Geochemistry of Evaporites in Lacustrine Basin, Dongying Depression, Bohai Bay Basin, China

Zhonghong Chen1*, Jiqing Zhang2, Li Wang2 and Ming Zha1
1School of Geosciences, China University of Petroleum, Qingdao, Shandong, China
2Petroleum Developing Center in Shengli Oilfield, SINOPEC, Dongying, Shandong, China

Abstract

The middle and lower parts of the Fourth Member of the Shahejie Formation in the Dongying Depression are representatives of evaporites and mud rocks in Eastern China. To investigate hydrocarbon potential, some deep wells, such as the Hk1 well and Fs2 well, were intensively cored and evaluated by geochemistry.

The total salinity of the formation waters is very high and is dominated by CaCl2, suggesting favorable seal development and preservation of petroleum. The content of Al, Mn, and Ba and the ratios of V/Fe and (Fe+Al)/(Ca+Mg) were low, whereas the content of Na, Ca, and Mg and the ratios of Sr/Ba, Fe/Mn, and Sr/Ca were high in the evaporites and mudstones of the Fourth Member compared with those in the mudstones of the Third Member. Sr content was remarkably high. The co-sedimentary system of evaporites and mudstones displays a heavier isotopic carbon. The values of δ13C for different group compositions and n-alkanes ranged from -28‰ to -21‰ and -18‰ to -32‰, the fractionation of δ13C between group compositions was no greater than -2‰, and δ13C of C15 and C16 significantly fluctuated. δ13C for high-maturity evaporite source rocks was significantly increased by thermal evolution and thermochemical sulfate reduction. High content of gammacerane and low Pr/Ph was exhibited in the evaporite system compared with that in the non-evaporite system. Different responses of biomarker parameters for the different sedimentary systems were exhibited, such as C19/(C19+C23) terpanes, C29/(C27+C28+C29) steranes, and C24/C23 and C22/C21 tricyclic terpanes. The reverse trend of the maturation parameters at depths corresponding to abundant carbonate minerals indicates that carbonate retards the thermal evolution of organic matter. Evaporites and mudstones can generate and expel hydrocarbons. The tested samples were mostly types I and II of organic matter, and their original generating capacity can reach 40 and 20 mg/g rock, respectively. The efficiency of hydrocarbon expulsion reached 60%, but the distribution of organic matter and its generative potential were highly variable. In general, the mudstones show greater generative potential than the evaporites.

Keywords: Mineral element; Carbon isotope; Evaporite; Hydrocarbon generation; Hydrocarbon expulsion; Dongying depression

Introduction

The formation, physical properties [1-4] and hydrocarbon potential of evaporites and mudstones has always been of interest to the petroleum industry [5,6]. Evaporite rocks are widely developed in potashiferous basins around the world and closely relate to the formation and distribution of the global giant oil and gas fields. Specifically in China evaporites are widespread in some Tertiary rifted depressions, covering an area of 250,000 km2. The formation, physical properties [1-4] and hydrocarbon potential of evaporites and mudstones has always been of interest to the petroleum industry [5,6]. Evaporite rocks are widely developed in potashiferous basins around the world and closely relate to the formation and distribution of the global giant oil and gas fields. Specifically in China evaporites are widespread in some Tertiary rifted depressions, covering an area of 250,000 km2. The Dongying Depression is a typical lacustrine basin containing evaporites and mudstones. To investigate hydrocarbon potential, some deep wells, such as the Hk1 well and Fs2 well, were intensively cored and evaluated by geochemistry.

The Dongying Depression is a typical lacustrine basin containing evaporites and mudstones in eastern China. Dark mudstone is mainly deposited in the upper part of Member 4 (Es3) and the lower and middle parts of Member 3 (Es2) of the Shahejie Formation in the Paleogene system; these parts are regarded as main hydrocarbon source rocks in the Depression [10-13]. However, exploration for many wells in the northern slope of the Depression has generated high-yield oil and gas from the evaporites and mudstones in the middle part of Member 4. This yield is a favorable prospect for oil and gas exploration and thus arouses considerable concern regarding evaporites and mudstones [9, 13]. However, the geochemistry of evaporites and mudstones has rarely been investigated for fewer wells drilled through deeply buried source rocks. Exploration has carried out few deep drillings, such as the Hao-1 Well and Fs2 well. On this basis, more intensive coring and calculating was conducted to study the geochemistry of evaporites and mudstones.

The study primarily aims to (1) better understand the response of mineral elements and related ratios in a saline lacustrine compared with those in a fresh lacustrine, (2) define the response of δ13C in a saline lacustrine, (3) improve understanding of biomarkers in a saline lacustrine, and (4) determine the capacity of evaporites to generate and expel hydrocarbon.

Geological Setting and Evaporite Distribution

The Dongying Depression is a typical negative rifted tectonic unit located in the southern part of the Bohai Bay Basin and characterized as northern faulted, southern overlapped, and opening-shaped. The depression developed from ancient Paleozoic bedrock with various tectonic movements, covering an area of 5700 km2.

A set of clastic materials is deposited in the Paleogene system and is dominated by mudstone intermixed with sandstone, carbonate, and gypsum-salt rocks. Under the impact of tectonic movement, the sediment has undergone a sedimentary process in three different stages: salted lake, deep lacustrine, and river delta. In the process, the main source rocks deposited in this depression are Es4, deposited at a hypersaline environment, and Es5, deposited at a fresh-brackish environment.

*Corresponding author: Chen Zhonghong, School of Geosciences, China University of Petroleum, Qingdao, 266580, Shandong, China, E-mail: hongczh@163.com

Received August 20, 2013; Accepted September 27, 2013; Published October 04, 2013

Citation: Chen ZH, Zhang JQ, Wang L, Zha M (2013) Geochemistry of Evaporites in Lacustrine Basin, Dongying Depression, Bohai Bay Basin, China. J Pet Environ Biotechnol 4: 156. doi:10.4172/2157-7463.1000156

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In the lower part of E₄, some brine-lake sediment, including gray shale, oil shale, salt rock, gypsum, and alluvial fan, is deposited in an evaporating environment under weak tectonic activity and extremely hot and dry climate (Figures 2 and 3). In the central part of the lake during this period can be found about 400 m to 600 m of thick gray mudstone with salt rock and gypsum layers. In mid-E₄ deposition, tectonic subsidence significantly increased, and a large rifted lake formed between high and continuous mountains. A large amount of lake water evaporated with result of high salinity. The highly saline center (>100 g/l) is distributed in the Minfeng sag of the northern basin, coinciding with the evaporite distribution (Figure 4). The salinity increased from the edge to the center of the salt lake, and some shales, marls, plasters, salts, and mudstones were deposited in sequence. In the alternation of oxygen environments with hypoxic environments, co-sediment of evaporites and mudstones was formed.

Experiment Methods

The contents of inorganic elements were measured by conventional ICP-AES and atomic absorption spectra in well drilling. Formation pressure and temperature were obtained by related testers in the drilling.

Sediment samples were crushed in a rotary mill, and aliquots were subjected to rock-eval pyrolysis to determine total organic carbon content, residual soluble hydrocarbon (S₁), pyrolysed hydrocarbon (S₂), production index, and hydrogen index (Iₜₚ), Tₚmax (temperature of maximum kerogen pyrolysate yield). Vitritine reflectance (Rₒ) was analyzed on randomly oriented grains by conventional microphotometric methods.

Chloroform is used as a solvent, and the samples are continuously extracted for 72 h at a constant temperature of 70°C in a Soxhlet extraction device. Ten to 15 milligrams of the samples were extracted, dissolved by petroleum ether, repeatedly washed, and moved in a flask, standing for 24 h (about 100 ml of petroleum ether in flask). The solution was finally filtered. The insoluble part was asphaltene, and the soluble part was a mixture of saturated hydrocarbons, aromatic hydrocarbons, and non-hydrocarbons.

A prepared filling column of silica-alumina and alumina was activated for 5 h at a constant temperature of 450°C, and silica was activated for 8 h at a constant temperature of 150°C. Filled column chromatography was conducted with the silica and alumina at a ratio of 3:1 (silica in the up of alumina). After the column was wet with petroleum ether, the sample filtrate was poured into the column and washed with petroleum ether, dichloromethane, and diethyl ether. Saturated hydrocarbons, aromatic hydrocarbons, and non-hydrocarbons were then obtained.

HP-Chemistation was used as a GC-MS analytical instrument (model: HP-5890 II type for GC, HP-5890A for MS; column: DB5-MS; column length: 60 m; internal diameter: 0.25 mm). The initial temperature was 100°C, the heating rate was 4°C/min, the final temperature was 320°C, and the final time was 20 min. The temperature of gasification and transmission pipeline was 310°C, the ion source...
temperature was 250°C, and the pre-column pressure was 170 kPa. The split ratio was 20:1, and the carrier gas was helium. Electron energy was 70 eV, the scan rate was 0.46 scans/s, and ionization was conducted by electronic impact.

Biomarkers were identified on the basis of mass fragment graphic responses and relative retention times, compared with a standard sample. Quantification was conducted by peak height responses in specific mass chromatograms, such as the m/z 191 and m/z 217 mass chromatograms.

The δ¹³C values for n-alkanes were obtained by a Finnigan Mat GC-IRMS with a delta-S mass spectrometer. In a typical analysis, 1 µl of n-alkane concentrate was injected into a DB-5 chemically bonded fused silica capillary column (30 m × 0.32 mm i.d.) with splitless mode. Helium was used as a carrier gas. The GC oven temperature was programmed from 60°C to 120°C at 30°C min⁻¹ and then to 310°C at 5°C min⁻¹. Injector temperature was 300°C. The measurement was conducted three or four times in most cases.

Results and Discussion

Inorganic elements in evaporites

In evaporate system the hydrocarbon generation is often associated with certain metals. For example, sulfate reducing agent can extract metal ions from salt water, and some metal ions can be selectively absorbed by the algae and higher aquatic plants around, which are released from decomposition of organic matter later. Amino acids generated from organic matter may improve the solubility of heavy metals. Therefore, the inorganic elements can reflect the sedimentary environment, water-rock interaction and hydrocarbon fluid flow.

The cores in the Paleogene system from the Hk1 Well were tested intensively and analyzed to investigate the inorganic elements of the evaporites and mudstone and to contrast the distribution rules for different sedimentary environments. The evaporites in Es₄ have low Al content (average=3.49%) but high Na, Ca, and Mg content (average=7.03%, 10.11%, 2.68%, respectively) compared with the mudstone in Es₃. Among the trace elements, Mn and Ba in the evaporites have significantly low content (average=697.9 × 10⁻⁶ and 473.6 × 10⁻⁶, respectively), whereas Sr content is remarkably high (average=1135.35 × 10⁻⁶). The ratios V/Ni and (Fe+Al)/(Ca+Mg) are low (average=1.99 and 0.60, respectively), whereas the ratios Sr/Ba, Fe/Mn, and Sr/Ca are high (average=3.99, 85.59, and 0.019, respectively). Particularly, the Sr/Ba ratio in the Es₄ evaporite system is five times that in the Es₃ non-evaporite system.

The chemical field of formation water also reflects the strong response of some major rock-forming elements in the evaporite system. Rock-forming elements sodium (Na), calcium (Ca), and magnesium (Mg) are the main cations in formation water, and their contents are well correlated with salinity. The salinity in the evaporite system is remarkably high, ranging from 1.63 g/l to 335.50 g/l and averaging 63.3 g/l.

The distribution of inorganic elements in the evaporites and
Figure 3: The lithologic log of the co-sedimentary system of evaporites and mudstone and their representative photos in the Es4 from Hk1 Well in Dongying depression.

1. mudstone; 2. sandy mudstone; 3. oil shale; 4. calcareous clay; 5. gypsum mudstone; 6. saline mudstone; 7. soft mudstone; 8. siltstone; 9. dolomitic siltstone; 10. fine sandstone; 11. argillaceous dolomite; 12. gypsum; 13. gypsum salt rock; 14. Salt rock
mudstone of Es4 shows distinct discreteness and fluctuation (Figure 5). The content of most inorganic elements is high in the mudstone but low in adjacent evaporites, except Na (Table 1). Such distribution is supposed to be associated with the special nature of the dissolution and adsorption of the inorganic element. When organic matter generates hydrocarbon, the organic acids generated can dissolve soluble inorganic salts in evaporites, and dissolved inorganic elements can easily migrate to adjacent mudstones and can accumulate related elements in clayed mudstones. The loss of elements in gypsum-salt for the clayed mudstones has a comparatively large surface area and strong adsorption. The evaporites and mudstone containing many catalytic transitional elements benefit the generation of oil and gas [14-16].

**Water type in the evaporate system**

The formation water chemical characteristics are reflection of different sedimentary and diagenetic environment, and ionic substances are the result of water-rock interaction and hydrocarbon accumulation. The ions Cl-, SO4^2-, HCO3^-, Na^+, K^+, Ca^2+ and Mg^2+ were known as the main ions in formation water in oil fields. In a closed hydrological system the composition of concentrated lake brines largely depends on labile lithologies that are leached and altered by inflow waters in the drainage basin surrounding a salt lake [17,18]. For example, flow across and through limestone aquifers produces inflow waters rich in Ca and HCO3, dolomite dissolution generates Mg-rich waters, igneous and metamorphic matrices tend to yield silica-rich Ca–Na–HCO3 waters, and pyritic shales and other sulfur-rich sediments, such as tuffaceous ashfalls, will contribute inflows rich in sulfate, whereas basic and ultrabasic rocks tend to produce alkaline Mg–HCO3 waters [19].

Due to the groundwater evaporation, concentration and salt precipitation chloride predominate in the groundwater with main NaCl salt. The long-term interaction between Water and rock make the greatly cation exchange effect for the absorbed Ca^2+ in the rock with Na^+ in groundwater, and the exchange reduced the dissolution of the Na^+, increase the dissolved Ca^2+, resulting in the formation of CaCl2 type water. Generally, CaCl2 type water originates from the deep earth, and reflects a reductive environment with water stagnated and salt highly concentrated. Here, the formation water is dominated by CaCl2, and some water shows NaHCO3. The Na2SO4 and MgCl2 types of formation water are rare. The development of CaCl2 type water in the evaporites suggested favorable seal development and petroleum preservation.

**Carbon isotope response to evaporites**

The value of the carbon isotopes of crude oil (δ13C) mainly depends on organic matter. High δ13C is displayed in many hypersaline environments, including some basins in China, such as Qaidam Basin [20], Qianjiang Depression in Jianghan Basin, [21] and Bohai Bay Basin [22]. The heavier carbon isotope is usually associated with the organic matter source mainly dominated by algae and aquatic bacteria [23,24].

High δ13C is also shown in the evaporites and mudstones. The δ13C of alkanes, aromatics, and non-hydrocarbons is greater than -27.6‰, -25.5‰, and -25.7‰, respectively, in the intervals of 3963 m to 3973 m and 4024 m to 4501 m of the F52 Well in the Dongying Depression (Table 2 and Figure 6). At the same time, the fractionation of δ13C between different group compositions is relatively small, and the difference is less than 2‰ mostly. The highest fractionation occurs between the saturated hydrocarbons and aromatic hydrocarbons, reflecting the characteristics of δ13C fractionation in salted lakes [25].
Figure 5: The distribution of the content of mineral elements in the Paleogene system from Hk1 Well in Dongying depression. (a, major minerals; b, trace minerals; c, element ratio).
### Table 1: The distribution of mineral elements and some ration parameters in the co-sedimentary system of saline and mud in the middle part of E4 from the well Hk1 in Dongying depression.

| Depth (m) | Lithology            | Na (%) | K (%) | Ca (%) | Mg (%) | Fe (%) | Al (%) | Mn (10⁶) | Ni (10⁶) | Zn (10⁶) | Sr (10⁶) | Co (10⁶) | Ba (10⁶) | V (10⁶) | Ca/Mg | V/Ni | Fe/Mn |
|-----------|----------------------|--------|-------|--------|--------|--------|--------|----------|----------|----------|----------|----------|----------|---------|-------|-------|-------|
| 3650      | Gray mudstone        | 3.23   | 1.39  | 6.38   | 2.38   | 2.4    | 6.67   | 496.38   | 28.37    | 92.75    | 829.9    | 12.91    | 396.2    | 86.04   | 2.08  | 2.68  | 3.03  |
| 3659      | Dark-gray mudstone   | 2.27   | 1.56  | 14.32  | 2.56   | 2.75   | 4.45   | 470.54   | 29.32    | 92.64    | 1399.7   | 12.14    | 324.1    | 62.37   | 4.32  | 5.59  | 2.11  |
| 3715      | Gypsum-salt          | 11.52  | 1.39  | 8.27   | 2.4    | 2.47   | 3.96   | 399.93   | 27.82    | 81.5     | 657.7    | 10.98    | 184.4    | 64.59   | 3.57  | 3.45  | 2.32  |
| 3718      | Gypsum-salt          | 32.26  | 0.81  | 5.25   | 1.3    | 1.47   | 0.84   | 293.49   | 11.85    | 41.45    | 652.6    | 5.05     | 277.3    | 4.04    | 1.18  | 1.02  | 50.09 |
| 3730      | Gray mudstone        | 4.65   | 3.53  | 10.21  | 1.83   | 6.37   | 4.11   | 298.16   | 30.45    | 204.7    | 735.3    | 14.94    | 281.5    | 5.58    | 1.97  | 213.6 |
| 3745      | Gypsum-salt          | 36.72  | 0.25  | 0.24   | 0.9    | 0.46   | 0.47   | 73.25    | 7.14     | 16.89    | 458.0    | 1.69     | 377.4    | 10.89   | 1.21  | 1.53  | 62.80 |
| 3772      | Gypsum-salt          | 24.61  | 0.22  | 9.02   | 0.34   | 0.39   | 0.33   | 28.2     | 5.88     | 11.76    | 770.2    | 1.84     | 283.9    | 6.93    | 2.71  | 1.18  | 138.3 |
| 3802      | Purple-red mudstone  | 2.22   | 1.39  | 14.11  | 2.01   | 2.54   | 5.08   | 547.92   | 32.09    | 89.59    | 1247.7   | 14.1     | 379.5    | 66.32   | 7.02  | 2.07  | 46.36 |
| 3856      | Purple-red mudstone  | 3.65   | 2.01  | 3.92   | 2.42   | 1.68   | 6.8    | 403.09   | 34.79    | 117.8    | 801.8    | 13.45    | 816.4    | 100.9   | 1.62  | 2.87  | 24.81 |
| 3862      | Gypsum-salt          | 27.08  | 0.28  | 3.4    | 0.58   | 0.46   | 0.53   | 68.91    | 7.09     | 16.63    | 351.5    | 1.94     | 254.4    | 8.91    | 1.38  | 5.86  | 126.75|
| 3884      | Gypsum-salt          | 34.98  | 0.2   | 3.17   | 0.29   | 0.33   | 0.42   | 28.3     | 4.67     | 11.25    | 368.8    | 2.45     | 262.4    | 7.23    | 1.41  | 10.9  | 116.6 |
| 3996      | Purple               | 5.21   | 1.79  | 4.33   | 1.87   | 3.09   | 3.59   | 422.06   | 31.69    | 107.5    | 1429.4   | 12.81    | 583.0    | 90      | 2.45  | 2.32  | 73.21 |
| 3902      | Purple               | 3.76   | 1.75  | 10.82  | 2.54   | 3.02   | 3.55   | 354.54   | 28.92    | 96.73    | 825.4    | 13.89    | 487.8    | 88.95   | 1.69  | 4.26  | 23.8  |
| 3914      | Purple               | 3.95   | 1.44  | 8.67   | 1.98   | 2.66   | 5.65   | 643.14   | 25.95    | 97.61    | 618.7    | 12.93    | 262.6    | 60.7    | 2.36  | 4.38  | 34.36 |

### Table 2: The characteristics of carbon isotope of the group composition of the organic matter in the middle part of Member-4 of Shahejie Formation from the Well Fs2 in the Dongying depression.

| Depth (km) | Lithology            | Value of δ¹³C/‰ |
|------------|----------------------|-----------------|
| 3.963      | Dark-gray salted mudstone | -27.6          |
| 3.968      | Dark-gray salted mudstone | -27.3          |
| 3.971      | Dark-gray salted mudstone | -27.4          |
| 3.973      | Dark-gray salted mudstone | -27.4          |
| 4.024      | Salted rock           | -25.0          |
| 4.295      | Laminated mudstone    | -24.2          |
| 4.301      | Laminated mudstone    | -21.0          |
| 4.498      | Dark-gray gypsum-salted mudstone | -22.4          |
| 4.499      | Dark-gray gypsum-salted mudstone | -25.2          |
| 4.500      | Dark-gray gypsum-salted mudstone | -22.1          |
| 4.501      | Dark-gray gypsum-salted mudstone | -23.9          |

### Figure 6: The characteristics of carbon isotope of n-alkanes from the source rocks of the middle part of Member-4 of Shahejie Formation from the Well Fs2 in Dongying depression.
The δ13C of the n-alkane series only slightly varies, except for a remarkably high δ13C of C_{17} and C_{18}. The δ13C of n-alkanes increase slightly with increasing carbon number, reflecting a characteristic of the δ13C from mature alkanes in hypersaline lacustrines.

Comparedly, the values of δ13C in the interval 3963 m to 3973 m are low, around -27.5‰, -25.5‰, and -25.5‰ for alkane hydrocarbon, aromatic hydrocarbon, and non-hydrocarbon, respectively. By contrast, the values of δ13C in the interval 4024 m to 4501 m are higher, distributed in the ranges -27.5‰ to -21‰, -24.3‰ to -21‰, and -25.6‰ to -22.6‰, respectively. These values reflect the effect of thermal evolution on the distribution of δ13C in similar sedimentary environments. The measured vitrinite reflectance (R) is 0.95% at a depth of 3972 m, whereas at 4500 m, R reaches 1.2%, indicating much higher thermal maturation.

The stability of various carbon bonds varies: 13C-13C > 13C-12C > 12C-12C. The generated hydrocarbon has lighter isotopic composition than its parent material because the 12C-12C bond is easier to break, resulting in a heavier isotope (δ13C) gathered in residual organic matter. This result explains the heavier carbon isotope distribution in the interval 4024 m to 4501 m.

The distribution of δ13C was not only controlled by maturity but also resulted from the reaction Thermochemical Sulfate Reduction (TSR). TSR is a function of the interaction between sulfate and hydrocarbons, and in the process, sulfate minerals are deoxidized into acid gases, such as H2S. In carbonate rocks, the TSR reaction is the widespread occurrence of geological phenomena and has been reported in some China basins [26,27]. A certain thickness of anhydrite, adequate hydrocarbons, and higher paleothermal energy are the prerequisites for TSR reaction. TSR reaction is a very complex chemical reaction that significantly changes carbon isotopes and enriches H2S [28].

In TSR reaction with the consumption of hydrocarbon, the 12C-12C bond is broken with priority, and more 13C is involved in the reaction. More 13C is retained in the residual hydrocarbons; thus, residual hydrocarbon after the reaction is relatively enriched in 13C, and the gained weight of δ13C can reach 2‰ to 4‰ in the Sichuan basin [26]. In the interval 4024 m to 4501 m of the Fs2 Well in Dongying Depression, conditions exist for TSR reaction with higher content of sulfate and high temperature. As the differentiation of the δ13C value in the two intervals (3963 m to 3973 m and 4024 m to 4501 m) reaches more than 5%, the TSR reaction contributes to δ13C distribution.

**Biomarker response in evaporite system**

**Source-related biomarker parameters:** The mature to highly mature hydrocarbon source rocks in the evaporite system tend to have a predominance of even carbon numbers, showing the "single-peak" distribution of n-alkanes (Figure 7). The peak carbon is located between nC_{14} and nC_{16}, and most of the peak carbons are nC_{15} and nC_{16}. The distribution of Pr/Ph is well differentiated by the two class of samples, and the obvious advantage of phytane (Pr/Ph<1.0) is demonstrated in the evaporite samples from E4 (Figure 8). Haven et al. [29] found that low Pr/Ph (<1) is typical of hypersaline environments, where the ratio appears to be controlled by the effects of variable salinity on halophilic bacteria [29]. Likewise, Schwark et al. [30] noted a general decrease in Pr/Ph during the deposition of the Permian Kupferschiefer sequence in Germany and interpreted it to indicate increasing paleosalinity [30]. Although the utility of Pr/Ph to accurately describe the redox state of paleoenvironments is limited by several factors, including variable source inputs, different rates of early generation, variations at higher maturity, and analytical uncertainty [31], the supporting relationship between Pr/Ph and the gammaceran index confirms the inferred salinity relationship.

In the series of tricyclic terpanes, gamma content is remarkably high, indicating of marked stratification in a salted water environment with halocline [32,33], because gammacerane indicates a stratified water column in marine and non-marine source-rock depositional environments [6], commonly resulting from hypersalinity at depth (Figure 9). The value 0.065 of γ/C_{30}H is a division for source rocks between the evaporite system and non-evaporite system. For the evaporite system, the value of γ/C_{30}H is generally higher than 0.065, but not for the non-evaporite system. This result indicates that increased water salinity during the deposition of source rocks increases gammacerane indices and reduces pristane/phytane, corresponding to the result of the investigation of the biomarkers of oils from lacustrine source rocks in Angola [32]. In the distribution of hopane series with a long chain, the characteristics of C_{27}hopane > C_{28}hopane > C_{29}hopane are shown, and the baseline shifts (Figure 9), fully reflecting the characteristics of biomarkers in a depositional environment with high salinity [5, 34-36]. At the same time, the differentiation of biomarker
responses for different source rocks in the evaporite system of \( E_4 \) and the non-evaporite system of \( E_3 \) in the Dongying Depression is exhibited by other indicators, such as \( C_{19}/(C_{19}+C_{23}) \) from terpanes, \( C_{29}/(C_{27}+C_{28}+C_{29}) \) from steranes, and \( C_{24}/C_{23} \) and \( C_{22}/C_{21} \) from tricyclic terpanes (Figure 10). The divided values of indicators \( C_{19}/(C_{19}+C_{23}) \) and \( C_{24}/C_{23} \) are 0.70, 0.27, and 0.30, respectively.

For the source rocks from the evaporite system, the values of \( C_{19}/(C_{19}+C_{23}) \) and \( C_{24}/C_{23} \) are generally lower than 0.70 and 0.27, respectively, whereas the values of \( C_{22}/C_{21} \) are generally higher than 0.30.

**Biomarker maturity parameters:** An analysis of many samples of the profile of the Paleogene System in the Dongying Depression reveals a distinct response of typical maturity-related biomarker parameters in
the evaporite system compared with that in the non-evaporite system. The values of $C_{29}\beta\beta/(\beta\beta+\alpha\alpha)$, $C_{29} 20S/(20S+20R)$, and $Ts/(Ts+Tm)$ began negative reversion at a depth of around 3.5 km to 3.6 km in the trend of their evolution, consistent with the sharp increase of the carbonate content in the interval (Figures 11 and 12).

Regional geological data show that at a shallower burial depth (<3.57 km), the main source rocks are calcareous shale and clayed mudstone and that at a great burial depth (>3.57 km), the gypsum and salt rock content increases significantly and the evaporite and mudstone form a co-sedimentary system. Take the HK1 Well as an example. Below 3.57 km, gypsum and salt rock developed, and in the symbiotic system of evaporite and mudstone, the content of elements, such as calcium and sodium, increased significantly than that in the upper layer; the ratios of Sr/Ba, Sr/Ca, and Fe/Mn showed a significant high-value response, indicating a typical high-salt environment [36]. In such a high-salt environment, the negative reversion of $C_{29}\beta\beta/(\beta\beta+\alpha\alpha)$, $C_{29} 20S/(20S+20R)$, and $Ts/(Ts+Tm)$ showed good relations with high levels of gypsum-salt, reflecting the effect of inhibition or retardation on thermal evolution by gypsum-salt minerals.

The retarding effect of carbonate minerals in the transforming process from organic matter to oil and gas is an important reason for the delay and backwardness of oil and gas generation in the source rocks of evaporite systems, and such retarding effect is mainly controlled by the absence of the catalysis of clay minerals [37]. The related thermal simulation experiment also showed that carbonate rocks have an anti-catalytic effect on hydrocarbon generation from kerogen and that increasing temperature significantly increases the anti-catalytic function [38]. Clay minerals in shale catalyze the transformation of kerogen, whereas calcites in gypsum-salt rock have a small or no role in the transformation of kerogen. In addition, organic matter in gypsum-salt rock is mainly found within cryptocrystalline or fine-grained carbonate minerals and is not easily released and thus may retard thermal evolution in evaporite systems [39].

Retardation was demonstrated by many immature to low-maturity oils discovered in several co-sedimentary systems of evaporite and mudstone. Examples include source rocks from the Lower Cretaceous in the South Florida Basin, the Guang-33 Well in the Jianghan Basin, the Yuecan-1 and Kaiqian-67 Wells in the Qaidam Basin, and the Baminhe area in the Dongying Depression. For instance, in the Biyang Depression in China, the threshold depth of oil generation from the dolomites was 300 m deeper than that from the mudstone in the Paleogene system, and the maturity indicated by biomarkers was significantly negatively correlated to the carbonate content in rocks.

Thermal retardation in the evaporite system is not only reflected by biomarker parameters but also displayed in the distribution of conventional thermal indicators, including vitrinite reflectance ($R_o$) and pyrolysis peak temperature ($T_{max}$). The negative departure of $R_o$ and $T_{max}$ in the overpressed interval (high overpressure, the pressure coefficient reaches 1.8) from 2400 m to 3400 m suggests the inhibition of thermal evolution in response to overpressure [36]. The greater decline of $R_o$ and $T_{max}$ in the hypersaline interval from 3400 m to 4000 m indicates similar retardation because of the presence of carbonate minerals, as with related biomarker maturity ratios [36].

**Hydrocarbon Generation and Expulsion from Evaporites**

**Group composition and organic matter type**

The capacity of hydrocarbon generation and expulsion from the co-sedimentary system of evaporites and mudstone (salted lake phase) is of widespread concern. Organic matter and high-quality source rocks are enriched in the depositional sequence of carbonate, sulfate, and chloride salt in salted lakes. In different sequences, the composition of organic matter and thermal evolution show a certain difference.

Evaporites in $E_s$ in the Dongying Depression are mainly composed of anhydrite and salt rock, and their paleosalinity is about 12% to 26% [40]. The high-salt environment may limit the growth of parasites and may be conducive to the formation of algal blooms and may thus increase the productivity of organic matter [13,41]. Moreover, the heat-conductive property of gypsum-salt is much higher than that of sand and shale; thus, source rocks in thick layers of gypsum rock can obtain heat earlier [9]. This early heating facilitates hydrocarbon generation.

To investigate the ability to generate and expel hydrocarbons from this co-sedimentary of evaporites and mudstone, a series of indicators was investigated from the Fs2 Well. The basic composition of the
organic matter is sapropel (the dominant component) and vitrinite (Table 3).

The sapropel and vitrinite content is 78% to 92% and 8.3% to 22.3%, respectively, and few exinite and inertinite exist. The index of the organic matter type is 58.9% to 85.4%, representing types I and II of kerogen. The group composition of the organic matter is mainly alkane (content: 74.49% to 80.98%) and aromatics and non-hydrocarbons in lower proportion (aromatics: 3.13% to 11.36%; non-hydrocarbons: 4.71% to 15.25%). The maturity parameters of vitrinite reflectance \( R_t \) and pyrolysis peak temperature \( T_{max} \) are greater than 0.93% and 400°C, respectively, indicating that the samples have passed the peak period of oil generation. However, the distribution of maturity parameters in the co-sedimentary system shows an irregular linear increase with burial depth. The increased thermal conductivity of the evaporites reduces the maturity parameters in the evaporites compared with those in the mudstone. For instance, \( R_t \) and \( T_{max} \) pyrolysis in the mudstone at 4300.92 m are 1.17% and 419°C, respectively, whereas in the gypsum-salted mudstone at 4498.8 m to 4499.8 m, the three samples of \( R_t \) are distributed from 1.13% to 1.14% and the \( T_{max} \) of two samples are 403 and 417°C, respectively.

**Capacity for hydrocarbon generation and expulsion**

A calculation model of hydrocarbon generation and expulsion was built based on geochemical information from pyrolysis [10]. A series of indicators was calculated to study the capacity for hydrocarbon generation and expulsion from the co-sedimentary system in the two intervals of the Fs1 Well.

Results show the stronger capacity of hydrocarbon generation and expulsion in the interval 3966 m to 3975 m (Figure 13): residual organic carbon content \( C_t \) ranged from 0.6% to 2.63% with an average of 1.54%; original organic carbon content \( C_w \) was distributed in the range 1.8% to 4.1%; the content of residual soluble hydrocarbons \( S_t \) and pyrolysis hydrocarbons \( S_w \) is distributed in the range 0.85 mg/g to 8.24 mg/g and 0.48 mg/g to 5.69 mg/g, averaging 4.31 and 2.83 mg/g, respectively; the remaining hydrocarbon generation potential \( S_t + S_w \) is distributed from 1.33 mg/g to 13.54 mg/g with an average of 7.14 mg/g; the soluble hydrocarbon index \( HI_t \), equivalent to \( S_t \times 100/C_w \), is distributed in the range 142 mg/g to 320 mg/g with an average of 255 mg/g, indicating that the organic matter entered the peak period of hydrocarbon generation. The conversion rate of pyrolysis \( K_t \), equivalent to \( S_t \times 100/\left( S_t + S_w \right) \), is high, generally greater than 55%, whereas the hydrogen index \( HI_t \), equivalent to \( S_t \times 100/C_w \), is significantly low (less than 200 mg/g). The original hydrocarbon generating potential \( S_t \) and the amount of hydrocarbon expelled \( S_w \) are distributed in 10 mg/g to 40 mg/g and 10 mg/g to 20 mg/g, respectively. The efficiency of hydrocarbon expulsion \( K_t \), equivalent to \( S_t \times 100/\left( S_t + S_w \right) \), is higher than 60% generally, indicating that the source rocks in the co-sedimentary system can generate and expel hydrocarbon.

In the 4498 m to 4504 m interval, source rocks enter the high-maturity stage, and the indicators are quite different, showing that the high thermal maturity reduced the residual capacity for hydrocarbon generation and expulsion: \( HI_t \) is no higher than 150 mg/g, and the \( HI_t \) of most of the samples is lower than 50 mg/g; \( KH_t \) is less than 50 mg/g; \( K_t \) is over 60% mostly; and \( K_t \) is more than 90%, whereas some samples even reach 95%. \( S_t \) and \( S_w \) are lower than 1.0 and 0.5 mg/g, respectively; \( S_t + S_w \) is no greater than 1.5 mg/g. The original potential of hydrocarbon generation and expulsion is also limited, lower than 15 mg/g, and that of most of the samples is below 6 mg/g.

**Discussion: Significance of the Evaporite**

The formation of evaporites

Regarding the formation of the evaporites here, Chen et al. [42] and Jian-hui et al. [43] suggested that evaporites form in a shallow lake, whereas mudstones form in a deep lake [42,43]. Laiyi et al. [44] considered that co-sediment is formed in alternating sedimentary environments from salted water to fresh water [44]. However, the current dominant view is that the co-sediment is formed in deep water, whereas a salted lake background is characterized by “mountains and deep basin” [45,46]. This point is based on the fact that the Paleogene Dongying Depression is surrounded by mountains, buried hills, and alluvial fans, which form a comparatively closed sedimentary system.

Salt lake in the western Qaidam basin in China is a typical inland salt lake, in which has formed extensive evaporite, and its formation mechanism can be referred. In the co-sedimentation of the evaporites and source rocks in age of Oligocene and Miocene in the western Qaidam basin, the prospective oil source rocks were just distributed stratigraphically and teconically in the locations where halite and gypsum were deposited. A genetic model was postulated for evaporites and source rocks, which was described as “big shallow water saline lake surrounding small deep water brine lake” [47]. Jin and Zha [9] believed that water depth controlled both on deposit ion of the evaporites and the accumulation of organic matter in the salt lake environment [47].

The significance of evaporites for the organic matter

At the beginning of the evaporite formation, due to the salinity increase or the mixture of different salinity water, the bottom water is at stagnation, thus causing the death of various organisms. But because the river continuously supplied biological and organic substances, lakes contained a large number of various biological residue and organic material. When the lake salinity increased, the various creatures will die and drown in the lake, and in consequence humus mud phase at

| Depth (m) | Lithology | Maturity parameters | Composition (%) | Organic matter type | Group (%) |
|----------|-----------|---------------------|----------------|---------------------|-----------|
|          |           | \( T_{max} (^\circ C) \) | \( R_t \) (%) | sapropel | vitrinite | Type index | type | alkanes | aromatics | Non-hydrocarbons |
| 3967.8   | Salted mudstone | 422 | 0.93 | 78.7 | 21.3 | 62.7 | II | 80.53 | 11.32 | 4.74 |
| 3970.7   | Salted mudstone | 418 | 1.00 | 89.7 | 10.3 | 81.9 | I | 79.90 | 11.08 | 5.15 |
| 3972.3   | Salted mudstone | 417 | 0.95 | 91.7 | 8.3 | 85.4 | I | 80.11 | 11.26 | 4.83 |
| 3972.5   | Salted mudstone | 419 | 0.98 | 91.3 | 8.7 | 83.8 | I | 75.65 | 10.21 | 4.71 |
| 4300.9   | Laminated mudstone | 419 | 1.17 | 85.7 | 14.3 | 74.9 | II | —— | —— | —— |
| 4498.2   | Gypsum-salted mudstone | 403 | 1.13 | 90.7 | 9.3 | 83.7 | I | 79.26 | 3.13 | 9.09 |
| 4499.2   | Gypsum-salted mudstone | 421 | 1.14 | 87.7 | 12.3 | 78.4 | I | 80.98 | 4.35 | 7.61 |
| 4499.8   | Gypsum-salted mudstone | 417 | 1.14 | 78.3 | 20.3 | 61.8 | II | 76.53 | 4.50 | 12.54 |
| 4501.2   | Gypsum-salted mudstone | 427 | 1.24 | 82.7 | 15.7 | 69.3 | II | 64.49 | 6.54 | 25.23 |

Table 3: The characteristics of the organic matter in the E1s from the Well Fs1 in the Dongying depression.
the bottom of the lake basin formed in a weak redox environment. The continuously conveying biological and organic substances from rivers provide material base for the hydrocarbon formation.

Generally, the formation and preservation of effect organic matter in salt lake is in an anaerobic environment (strong reducing environment). The high salinity in salt lake makes a halocline permanent hierarchy easy to form. The anoxic environment forms under bottom water characterized by high salinity, and the organic matter avoids the decomposition from oxidation, resulting in the formation of high quality source rocks [48]. This is reflected in the Chaerhan modern salt lakes in Qaidam basin: 1 m below the salt water is a permanent brine layers, in the depth of 3~4 m is strong reducing environment, where can form black mud.

Numerous continental salt basins in the eastern China show that at the second or third sedimentary cycle is the stage of source rocks formation. Many practice indicated that such a set of evaporite formation commonly was before the inevitable development of source rock. Peters et al. [32] pointed out that in barrier type of sedimentary basins; the content of organic matter was associated with the concentration of salt that deposited at the same time [49]. Tissot [50] observed that the organic matter content of rock salt is second only to clay (or shale) based on analysis on 108 pieces of rock salt [50].

Zhanhua depression is adjacent to the northeast of Dongying depression, thus the significance of evaporites for the organic matter in this depression can be compared. In the Zhanhua depression also develops the evaporite sediment combination up to 500 m on the upper E_r. Research shows the evaporite exists not only with source rock, but also with high quality one. In salt lake on saline lake, the salinity difference between surface and bottom layers could result in the salt transition layer and the division of upper and lower layers makes a long-rang delamination of salinity. Bottom water circulation is in an absolutely stagnant state and results in anoxia environment. This environment restrains the activity of benthon and makes a maximum preservation of sediment organic matters. Oil-source correlation shows that salt lake saline lake source rock is an important one in Zhanhua Depression and controls the formation and distribution of many pools.

The significance of evaporites for the thermal maturation and oil generation

The thermal conductivity of gypsum and cream salt layer is significantly higher than other rocks, and the heating rate is significantly lower than other rocks. Test of the rock from the Hk1 Well shows the average thermal conductivity of the gypsum halite layer dominated by salt rocks is 5.48 W/m.K, and the average heat production rate of gypsum and salt rock are only 0.07 μW/m m and 0.02μW/m m respectively (Table 4). Compared to the evolution of normal mature hydrocarbon source rock (mudstone and shale) the salt layer shows lower vitrinite reflectance ($R_o$), and correspondly the paleogeothermal is lower than normal source rocks (Figure 14). Therefore, the existence of the halite layer makes the lower strata heat easily lose, lowers the temperature of the deep under the salt, suppresses the diagenesis, causes the diagenetic evolution of salt formation relatively lags behind the diagenetic evolution in the region without salt or gypsum, and correspondly
The significance of evaporites for the oil reservoir and preservation

Salt-gypsum layer lead to the formation of overpressure sealing strip, which suppresses the compaction and is conducive to the preservation of primary pores. The inhibitory effect of abnormal high pressure on the thermal evolution of organic matter and hydrocarbon generation, make the release space of organic acids in the abnormal high pressure basin and the influence of organic acids on sandstone diagenesis greatly exceed the normal pressure basin. Organic acid and CO$_2$ will release in the process of thermal evolution of organic matter, reduce the pH value of pore water, further strengthen the dissolution of minerals, make the cement and clastic particles in sandstone dissolve, and promote the formation of secondary pore. The pressured seal formed by Salt-gypsum layers is beneficial to the growth of secondary porosity under salt and preserved. For example, in the Paleogene Ek (Kongdian Member) and Es$_4$ there develop two overpressure belts and two interval depths (3398-3754m and 4052-4275.5m) for the development of secondary porosity accordingly, which are associated with the development of two halite layers respectively (Figure 15).

Figure 15 indicated that the development of secondary porosity under salt-gypsum layers was affected by rock types, structure and the development of primary pore at the same time. When primary pore relatively developed, acid formation water can smoothly enter between the particles and dissolve rock components.

The concentration layers of salt-gypsum in the Paleogene Es$_4$ in Dongying depression forms a good regional cap rocks for the oil and gas below. This is indicated by the occurring of cracking gas and even industrial air flow in the Fs1, Fs2 and Fs3 Wells in the Minfeng sag in the north of Dong depression. The thick salt-gypsum layer becomes a good cover to prevent dissipation of oil and gas (Figure 16). In the salt-gypsum layer in the north-central zone of Dongying depression there develops abnormal high pressure, and the local pressure coefficient reaches 2.05. The existence of abnormal high pressure caused by the salt paste interval further enhances the sealing capability of the rocks.

Conclusions

This study showed the geochemical response in a co-sedimentary system by evaporites and mudstone in the Es$_4$ of the Dongying Depression in Eastern China.

The total salinity was generally high. The type water of CaCl$_2$ dominated. The Al, Mn, and Ba contents and the parameters of V/ Ni and (Fe+Al)/(Ca+Mg) showed low value responses. The Na, Ca, and Mg contents and the parameters of Sr/Ba, Fe/Mn, and Sr/Ca showed high values. Sr content was remarkably high compared with that in the non-evaporite system. The distribution of mineral elements had visible characteristics of discretion and fluctuation, and mineral elements, except for Na, had discrepancies in saline deposits and were enriched in adjacent mud.
A heavier isotope of carbon (high value of $\delta^{13}C$) was displayed, and the value of $\delta^{18}O$ for the high-maturity evaporite source rocks was significantly increased by thermal evolution and TSR reaction. High content of gammacerane and low value of Pr/Ph were exhibited. The indicators $C_{23}/(C_{19}+C_{23})$ from terpanes and $C_{24}/C_{23}$ from tricyclic terpanes were lower, whereas $C_{29}/(C_{27}+C_{28}+C_{29})$ from steranes were higher than those in the non-evaporite system in the Dongying Depression. A distinct negative reversion of biomarker maturity parameters $C_{29}^{\beta\beta}/(\beta\beta+\alpha\alpha)$, $C_{29}^{20S}/(20S+20R)$, and $Ts/(Ts+Tm)$ was demonstrated, indicating that thermal evolution was restrained and retarded for the high content of gypsum-salt minerals. The co-sedimentary system can generate and expel hydrocarbon, and high maturation can reduce the capacity for these actions.

**Acknowledgments**

This study was supported by the National Natural Science Foundation of China (Grant No. 41272140 and 40802026) and Shandong Province Natural Science Foundation (Grant No. ZR2011DM004).

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