The Geochemical Characteristics of Coals from the Junggar Basin in Northwest China and the Relation of the Configuration of Pristane with Maturity in Highly Mature and Over-Mature Samples

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Abstract — Coal samples analyzed from Junggar Basin are characterized by high Total Organic Carbon (TOC) contents (63.7-82.2%) in all samples, except the highly over-mature samples Ha-01 and Ha-02, and high S2 (60.5-83.0 mg HC/g rock) and Hydrogen Index (HI) values (77-118 mg HC/g TOC), indicating the coal rocks have excellent source rock potential. The Tmax values (418-531°C), C29 sterane 20S/(20S+ 20R) ratios (0.20-0.54) and C31 homohopane 22S/(22S+22R) ratios (0.26-0.61) indicate an immature to highly over-mature stage of organic matter, which is supported by the organic geochemical maturation parameters. In addition, n-alkane carbon numbers range from C11 to C33 with maxima at n-C19 or n-C23, exhibiting unimodal distribution patterns except Ai-13 and Ha-01, and low Pr/Ph ratios (0.29-0.80) in Ai-01, Ai-13, Ha-01 and Ha-02, indicating a suboxic reductive environment. In Wu-8, Wei-09 and Shinan-20, the Pr/Ph ratios are clearly greater than 1.0 (2.43-3.23), indicating oxic depositional conditions.

Furthermore, pristane isomers were identified in extracts from these coal samples. Using coals of different maturity, which correspond to vitrinite reflectances (%R0) of 0.36-2.99%, the Pristane Isomerization Ratio (PIR) (PIR = [6(R)10(R)+ 6(S)10(S)]/6(R)10(S)) ranges from 0.42 to 0.97 for the coal extracts. The value of the PIR, which is a molecular maturity parameter, is evaluated by analyzing a series of samples using known values of the molecular maturity parameter based on the sterane and hopane isomerization indices, and the Methyl Phenanthrene Index (MPI). Changes to the PIR in highly mature Ai-13 (R0 > 1.5%) and over-mature Ha-01 and Ha-02 (R0 > 2.0%) are clear; thus, a linear correlation is evident between the PIR and R0. The results suggest that the PIR is an appropriate indicator of maturity for the highly mature and over-mature coal samples in the Junggar Basin.

Résumé — Les caractéristiques géochimiques des charbons du Bassin de Junggar au Nord-Ouest de la Chine et la relation entre la configuration du pristane avec la maturité dans des échantillons hauteur matures ou post-matures — Des échantillons de charbon analysés du Bassin de Junggar se caractérisent par une forte teneur en carbone organique total (TOC) (63,7-82,2 %) dans tous les échantillons,
INTRODUCTION

Molecular maturity parameters are useful geochemical parameters employed widely since the recognition of systematic changes in biomarker composition associated with an increase in depth and thermal maturation in natural burial sequences [1-3]. The sterane and terpane isomerization ratios (C_{29} steranes 20S/20S+20R, C_{31} homohopanes 22S/(22S+22R), T_m/T_{max}) and Methyl Phenanthrene Index (MPI) have been broadly applied to assess thermal maturity in crude oils and their source rocks [1, 4, 5]. Most of these maturity parameters are mainly based on the relative abundances of two stereoisomers with a relative increase in the more thermally stable, non-biological isomer compared with the isomer that exhibits the original biological inherited sterane skeleton. These parameters include the absolute concentrations of steranes and homohopanes, which decrease gradually with increasing maturity, the ratio of tricyclic terpanes/tricyclic terpanes + homohopanes, and the ratio of T_m/T_{max}, which increases rapidly at high maturities [1]. However, it was proposed that the ratios of tricyclic terpanes/tricyclic terpanes + homohopanes and T_m/T_{max} could be used to investigate maturities (vitrinite reflectance values, %R_o) up to 1.3-1.4% [4]. Normally, maturity parameters, which are based on biomarkers, cannot be applied to highly mature oils and source rocks (with corresponding vitrinite reflectances (R_o) > 1.5%) or to over-mature oils and source rocks (R_o > 2.0%). This is due to the fact that either the biomarkers are at their equilibrium values or the compounds employed in the parameters are thermally destroyed at the highly mature and over-mature stages. Therefore, the maturity of samples cannot be assessed using these parameters. Various high maturity parameters have been described and employed with great success in the last two decades (aromatic hydrocarbons, Methyl Diamantane Index (MDI)) [1, 2, 6-9]. However, most maturity parameters lack sensitivity in the highly mature area; thus, samples with similar maturity parameters cannot be distinguished. In addition, several classical aromatic hydrocarbon-based parameters show a reversal during higher levels of thermal maturity. One example is the Methyl Phenanthrene Index (MPI-1) [6], which may not be adequate when comparing oils with high maturity condensates. The MDI can only be used to measure the maturity of source rocks at highly mature stages in the vitrinite reflectance range of 0.9-2.0% R_o. As a result, the MDI cannot be used as a maturity parameter for over-mature gas source rocks at a R_o greater than 2.0% [2].

Acyclic isoprenoids are widely utilized biomarkers in organic geochemistry. Studies of the lipid components of archaea continue to broaden the known range of acyclic isoprenoid skeletons in organisms, and many biomarkers of acyclic isoprenoids have been identified in sediments, crude oils and coals. In particular, regular head-to-tail linked isoprenoids are compounds which appear to be uniquely synthesized by archaea [10]. The isoprenoid hydrocarbons pristane (2,6,10,14-tetramethylpentadecane) and phytane (2,6,10,14-tetramethylhexadecane) are typical acyclic isoprenoids and have been widely assumed to be diageneric...
products of the phytol side chain of chlorophyll [4, 11], although alternative sources of precursors have been suggested. The ratio of these two compounds is usually interpreted to be an indicator of the oxicity of the environment of deposition [12]. In addition, pristane is thought to derive from the degradation of α-tocopherol in anoxic sediments and contains 2 chiral centers (at C-6 and C-10) and can, therefore, exist as 3 possible isomers [13]. In this case, the 6R, 10S isomer (Fig. 1a,b) is separable from the unresolved enantiomers (Fig. 1c,d).

Previous studies have shown that with an increase in burial depth and increase in temperature (maturation), isomerization may occur at chiral centers. For pristane, there is a loss of stereospecificity, resulting in a mixture of the three isomers (Fig. 1a-c) at a ratio of 2:1:1 [14, 15]. In a number of cases, the structures of the geolipids were related to the enzymatically controlled stereospecificity of the presumed preceding natural products. For example, the pristane in the relatively immature Green River and Messel shales is primarily the 6(R), 10(S) (meso) compound [14].

The detailed stereochemistry of acyclic isoprenoid alkanes was studied in a suite of coal samples, ranging from lignite, sub-bituminous coal and bituminous coal to anthracite, in the Junggar Basin using GC-MS. To ensure that all lignocarboid products of the phytothyl side chain of chlorophyll [4, 11], although alternative sources of precursors have been suggested. The ratio of these two compounds is usually interpreted to be an indicator of the oxicity of the environment of deposition [12]. In addition, pristane is thought to derive from the degradation of α-tocopherol in anoxic sediments and contains 2 chiral centers (at C-6 and C-10) and can, therefore, exist as 3 possible isomers [13]. In this case, the 6R, 10S isomer (Fig. 1a,b) is separable from the unresolved enantiomers (Fig. 1c,d).

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The detailed stereochemistry of acyclic isoprenoid alkanes was studied in a suite of coal samples, ranging from lignite, sub-bituminous coal and bituminous coal to anthracite, in the Junggar Basin using GC-MS. To ensure that all of the changes observed would occur as far as possible from maturational differences, seven samples were chosen from different areas and were sorted to have different maturities. The samples reflect a wide range of burial history with a maximum vitrinite reflectance maturity assessment of 0.36-2.99% Rm. The aim of this study is to illustrate the suitability of the PIR as a parameter for high-maturity shales.

1 GEOLOGICAL SETTING

The Junggar Basin, located in the northern part of Xinjiang province, Northwest China, is a typical late Paleozoic, Mesozoic and Cenozoic superimposed basin (Mesozoic-Cenozoic terrestrial sedimentary basin) with an area of approximately 1.3 x 10^5 km^2 and also famous for its energy resources including oil, gas, coal and shale [16, 17]. The basin underwent pre-Carboniferous folding and basement metamorphism [18, 19]. From the Devonian to the Carboniferous, the basin is a platform basin in the Xingmeng Ocean surrounded by deep troughs [18, 20]. Later during the Late Carboniferous, the basin transformed from an open marine basin into a closed continental basin [18, 20]. After that, from the Permian to the Quaternary, the basin mainly experienced three tectonic stages: a foreland basin (Permian), an intracontinental depression (Triassic-Paleogene) and a rejuvenated foreland basin (Neogene-Quaternary) [21].

The northwestern margin of the basin lies at the junction of the western Junggar Hercynian Fold Belt [22]. The coal resources are rich in the southern margin of the Junggar Basin, fulfilling the strategic goal of coal exploration. Several coal fields have been discovered primarily in the fault zone area (Fig. 2), which includes 100-meter-thick high-quality coal seams and oil shales. The coal rocks in the study area include the middle Jurassic system (J2) and lower Carboniferous system (C1) [17, 23], and the middle Jurassic Xishanyao Formation (J2x) is the main coal-bearing stratum, with many workable coal seams in the Xishanyao Formation [23].

2 MATERIALS AND METHODS

The samples used in this study were collected from the seven coal mines within the Junggar Basin (Fig. 2, Tab. 1) including more immature coal (Wu-8, Rm = 0.36%), early mature coal (Wei-9, Rm = 0.60%), late mature coal (Shinan-20, Rm = 0.80%), late to past-oil-window coal (Ai-01, Rm = 1.25%), past mature coal (Ai-13, Rm = 1.53%), and over-mature coals (Ha-01, Rm = 2.32% and Ha-02, Rm = 2.99%). The samples are rich in vitrinite (70.0-98.6%), as is common for humic coal (Tab. 1).

About twenty grams of coal sample were broken into pieces; the broken samples were ultrasonically extracted (3x) with dichloromethane/methanol (93:7, v/v) for 30 min at 35°C, with centrifugation after each extraction. The supernatants were combined and carefully evaporated to dryness under a steady stream of N2 at 35°C. The fractions obtained were analyzed using the full chromatography, with a mixture of pre-activated silica gel/alumina (4:1, v/v) as a stationary phase and with n-hexane, dichloromethane and methanol as eluents, respectively. The fractions obtained were analyzed using the full scan and Selected Ion Monitoring (SIM) modes.

GC-MS analysis was performed using a Hewlett-Packard 6890 Gas Chromatograph (GC) coupled to a Hewlett-Packard 5973 MSD. The GC was fitted with a HP-5 capillary column.
Helium was used as the carrier gas at 0.9 mL min^-1. The following oven temperature program was applied: the oven temperature was raised from 80 to 280°C (held for 30 min) at 3°C/min for the full scan of samples (Fig. 3). In addition, an initial temperature of 80°C for 1 min, an increase in temperature to 100°C at a rate of 3°C min^-1, an increase in temperature to 130°C at a rate of 0.3°C min^-1, an increase in temperature to 280°C at a rate of 3°C min^-1 and holding at this temperature for 30 min for the separation of pristane was carried out (Fig. 4). The injection port was set at 280°C. The interface, ion source and quadrupole temperature were maintained at 280°C, 230°C and 150°C, respectively. The ionization was carried out in an electron impact mode at 70 eV, and the data were acquired using both the full scan and SIM modes. All enantiomeric separations were confirmed with GC-MS fragmentation comparisons with the NIST library. Regular steranes were examined with m/z 217 fragmentograms. Terpanes, primarily consisting of homohopanes, were examined with m/z 191 fragmentograms. Acyclic isoprenoid alkanes and methyl phenanthrene were examined using m/z 85, 123, 169, 183 and 192 mass chromatograms (Fig. 4). Nor-pristane, pristane and phytane were examined and quantified on m/z 254, 268 and 282 fragmentograms, respectively. Assignments for these compounds are given in Figure 5. Identification of the isomerization of acyclic isoprenoids in these samples is either based on comparisons with the results of Peters et al. [4] or on the examinations of the mass spectra of those compounds.

### TABLE 1
Vitrinite reflectance characteristics of the coal samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Age</th>
<th>Relative maturity</th>
<th>Maceral composition (%)</th>
<th>( R_o ) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Vitrinite</td>
<td>Liptinite</td>
</tr>
<tr>
<td>Wu-8</td>
<td>J2X</td>
<td>Immature</td>
<td>79.0</td>
<td>0.43</td>
</tr>
<tr>
<td>Wei-09</td>
<td>J2X</td>
<td>Early mature</td>
<td>83.2</td>
<td>0.60</td>
</tr>
<tr>
<td>Shinan-20</td>
<td>J2X</td>
<td>Late mature</td>
<td>82.1</td>
<td>0.98</td>
</tr>
<tr>
<td>Ai-01</td>
<td>J2X</td>
<td>Late to past mature</td>
<td>98.6</td>
<td>1.25</td>
</tr>
<tr>
<td>Ai-13</td>
<td>J2X</td>
<td>Past mature</td>
<td>97.2</td>
<td>1.53</td>
</tr>
<tr>
<td>Ha-01</td>
<td>C1</td>
<td>Over-mature</td>
<td>86.3</td>
<td>2.40</td>
</tr>
<tr>
<td>Ha-02</td>
<td>C1</td>
<td>Over-mature</td>
<td>70.0</td>
<td>2.00</td>
</tr>
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(100 m × 0.32 mm × 0.25 μm, Agilent, USA). Helium was used as the carrier gas at 0.9 mL min^-1. The following oven temperature program was applied: the oven temperature was raised from 80 to 280°C (held for 30 min) at 3°C/min for the full scan of samples (Fig. 3). In addition, an initial temperature of 80°C for 1 min, an increase in temperature to 100°C at a rate of 3°C min^-1, an increase in temperature to 130°C at a rate of 0.3°C min^-1, an increase in temperature to 280°C at a rate of 3°C min^-1 and holding at this temperature for 30 min for the separation of pristane was carried out (Fig. 4). The injection port was set at 280°C. The interface, ion source and quadrupole temperature were maintained at 280°C, 230°C and 150°C, respectively. The ionization was carried out in an electron impact mode at 70 eV, and the data were acquired using both the full scan and SIM modes. All enantiomeric separations were confirmed with GC-MS fragmentation comparisons with the NIST library. Regular steranes were examined with m/z 217 fragmentograms. Terpanes, primarily consisting of homohopanes, were examined with m/z 191 fragmentograms. Acyclic isoprenoid alkanes and methyl phenanthrene were examined using m/z 85, 123, 169, 183 and 192 mass chromatograms (Fig. 4). Nor-pristane, pristane and phytane were examined and quantified on m/z 254, 268 and 282 fragmentograms, respectively. Assignments for these compounds are given in Figure 5. Identification of the isomerization of acyclic isoprenoids in these samples is either based on comparisons with the results of Peters et al. [4] or on the examinations of the mass spectra of those compounds.

Figure 2
Locations of the five wells in the Junggar Basin from which coal samples were collected for this study.

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</tr>
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<td>Shinan-20</td>
<td>J2X</td>
<td>Late mature</td>
<td>82.1</td>
<td>0.98</td>
</tr>
<tr>
<td>Ai-01</td>
<td>J2X</td>
<td>Late to past mature</td>
<td>98.6</td>
<td>1.25</td>
</tr>
<tr>
<td>Ai-13</td>
<td>J2X</td>
<td>Past mature</td>
<td>97.2</td>
<td>1.53</td>
</tr>
<tr>
<td>Ha-01</td>
<td>C1</td>
<td>Over-mature</td>
<td>86.3</td>
<td>2.40</td>
</tr>
<tr>
<td>Ha-02</td>
<td>C1</td>
<td>Over-mature</td>
<td>70.0</td>
<td>2.00</td>
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(100 m × 0.32 mm × 0.25 μm, Agilent, USA). Helium was used as the carrier gas at 0.9 mL min^-1. The following oven temperature program was applied: the oven temperature was raised from 80 to 280°C (held for 30 min) at 3°C/min for the full scan of samples (Fig. 3). In addition, an initial temperature of 80°C for 1 min, an increase in temperature to 100°C at a rate of 3°C min^-1, an increase in temperature to 130°C at a rate of 0.3°C min^-1, an increase in temperature to 280°C at a rate of 3°C min^-1 and holding at this temperature for 30 min for the separation of pristane was carried out (Fig. 4). The injection port was set at 280°C. The interface, ion source and quadrupole temperature were maintained at 280°C, 230°C and 150°C, respectively. The ionization was carried out in an electron impact mode at 70 eV, and the data were acquired using both the full scan and SIM modes. All enantiomeric separations were confirmed with GC-MS fragmentation comparisons with the NIST library. Regular steranes were examined with m/z 217 fragmentograms. Terpanes, primarily consisting of homohopanes, were examined with m/z 191 fragmentograms. Acyclic isoprenoid alkanes and methyl phenanthrene were examined using m/z 85, 123, 169, 183 and 192 mass chromatograms (Fig. 4). Nor-pristane, pristane and phytane were examined and quantified on m/z 254, 268 and 282 fragmentograms, respectively. Assignments for these compounds are given in Figure 5. Identification of the isomerization of acyclic isoprenoids in these samples is either based on comparisons with the results of Peters et al. [4] or on the examinations of the mass spectra of those compounds.

Figure 2
Locations of the five wells in the Junggar Basin from which coal samples were collected for this study.
Component ratios for the isoprenoid alkanes were calculated from GC peak areas using the appropriate standards.

Vitrinite is a coal maceral derived from the lignin and cellulose of plant cell walls and fillings. Vitrinite reflectance ($R_o$) is a measurement of the percentage of reflected light at high (500°C) magnification in oil immersion. Vitrinite reflectance increases with the increasing thermal maturity of the sample. The mounted sample plugs were polished and analyzed for $R_o$ using a Leitz MVP III microscopic photometer system [11, 24].

Total Organic Carbon (TOC) contents for the samples were measured with a LECO CS-334 elemental analyzer. Prior to analysis, the samples were dried, powdered and leached with 12% HCl to remove the inorganic carbon and washed with distilled water to neutral pH, followed by drying at low temperature. All TOC values were reported in percent relative to the dry weight of the sample.

3 RESULTS

3.1 Vitrinite Reflectance and Rock-Eval Pyrolysis Generation Characteristics

The vitrinite reflectance ($R_o$) and Rock-Eval and TOC data are summarized in Tables 1 and 2. The vitrinite reflectance
and Rock-Eval pyrolysis are used to determine the hydrocarbon potential, thermal maturity of the organic matter and its ability to generate oil and/or gas [11]. The pyrolysis gives rise to the parameters $S_1$, $S_2$, Potential Yield (PY), the Hydrogen Index (HI), the Oxygen Index (OI) and $T_{\text{max}}$. The vitrinite reflectance ($R_o$) ranges between 0.36 and 2.99 with a big change interval (Tab. 1). The $T_{\text{max}}$ values range from 418 to 531°C (Tab. 2). These results are consistent with the low maturity to high over-maturity of the organic matter in the study area. The TOC in the samples are high, from 63.70 to 82.20%; the average value is 74.31% (Tab. 2). The average $S_1$, $S_2$ and PY values are 2.83, 71.72 and 73.83 mg HC/g coal rock for samples, with the exception of samples Ha-01 and Ha-02, which were 0.63, 4.00 and 4.63 mg HC/g coal rocks, respectively (Tab. 2). The HI values are from 77 to 118 mg HC/g TOC in Wu-8, Wei-09 and Shinan-20, the Ph content is higher than the Pr content, and the Pr/Ph ratios range from 2.43 to 3.23. The Pr/Ph ratios are clearly greater than 1.0, indicating oxic depositional conditions. However, in Ha-01, Ha-02, the Py content is higher than the Pr content, and the Pr/Ph ratios range from 0.73 to 0.80 (Fig. 3, Tab. 3). This shows a suboxic reductive environment [4]. In Ai-01, the ratio of Pr/Ph is 0.28. This is thought to be due to reduction environments [4]. The difference in Pr/Ph values here may also be caused by the impact of maturity to a certain degree [25].

### 3.3 Separation of Pristane using GC-MS

The C$_{19}$ alkane 2,6,10,14-tetramethylpentadecane has been clearly identified in a series of coals from the Junggar Basin. The process of determining the relative configuration of pristane in geological samples has been discussed [15], in which the meso (I, II) isomer is separated from the two enantiomers (III, IV). The samples (Wu-8 and Wei-09) showed a preference in favor of the meso isomer (Fig. 6). Although the absolute configuration has not been examined, all of the samples appear to show some degree of isomerization. The relative abundance of the III and IV enantiomers (SS + RR) in all of the samples shows a trend of increasing isomerization (Fig. 6), yet Ha-02 shows a nearly complete isomerization. With previous geochemical investigations, stable hydrocarbons increase in relative abundance over less stable ones with an increase in thermal stress. Therefore, we assume that the acyclic isoprenoids can be used in maturity assessment for highly mature coals and oils.

### 4 DISCUSSION

#### 4.1 Depositional Environment

The $n$-alkanes are in the C$_{11}$ to C$_{33}$ range, with peaks of $n$-C$_{19}$ or $n$-C$_{23}$ (Fig. 3); except for Ai-13 and Ha-01, the coal rocks studied show a unimodal distribution. The OEP is significantly greater than 1.0 for Wu-8, Wei-09 and Shinan-20 (Tab. 3), indicating that odd-numbered $n$-alkanes are dominant in these samples. However, it is nearly 1.0 for Ai-01, Ai-13, Ha-01 and Ha-02, indicating that odd-numbered $n$-alkanes are not dominant in these samples. In Wu-8, Wei-09 and Shinan-20, the Pr content is higher than the Ph content, and the Pr/Ph ratios range from 2.43 to 3.23. The Pr/Ph ratios are clearly greater than 1.0, indicating oxic depositional conditions. However, in Ai-13, Ha-01 and Ha-02, the Ph content is higher than the Pr content, and the Pr/Ph ratios range from 0.73 to 0.80 (Fig. 3, Tab. 3). This shows a suboxic reductive environment [4]. In Ai-01, the ratio of Pr/Ph is 0.28. This is thought to be due to reduction environments [4]. The difference in Pr/Ph values here may also be caused by the impact of maturity to a certain degree [25].
microorganisms or by microbial reworking of plant
n-alkanes [26]. The pristane is formed from the chlorophyll
phytvl side chain by an oxidative pathway, while phytane is
generated through various reductive pathways; the ratio of
pristane to phytane (Pr/Ph) has been proposed as an indicator
of the oxicity of the depositional environment [28, 29].
As shown in Table 3, the ratio of Pr/Ph is within the range
of 0.29 to 0.80 for Ai-01, Ai-13, Ha-01 and Ha-02. These
values are thought to be due to a reduction to weak oxida-
tion-reduction environment, which is possibly favorable
for reducing conditions in the lacustrine sediments. How-
ever, in Wu-8, Wei-09 and Shinan-20, a Pr/Ph > 3.0 typi-
Fig. 5
Full scan mass spectra of nor-pristane, pristane and phytane isomers.

4.2 Maturity of the Organic Matter

The kerogen maturity was evaluated on the basis of Rock-
Eval $T_{\text{max}}$. The range of $T_{\text{max}}$ values varies between 418
and 531°C (Tab. 2). This range indicates that the thermal
maturity level of the organic matter belongs to the immature
to highly over-mature [4]. The ratios of C$_{29}$ steranes 20S/
(20S+20R), C$_{29}$ steranes $\beta\beta/(\beta\beta+\alpha\alpha)$ and C$_{31}$ homohopanes
22S/(22S+22R) have been widely used as maturity indicators [30, 31]. In our studied samples, the ratios of C29 steranes 20S/(20S+20R), C29 steranes bb/(aa+bb) and C31 homohopanes 22S/(22S+22R) are in the range of 0.20-0.54, 0.21-0.44 and 0.26-0.61, respectively (Tab. 3). This range indicates that the thermal maturity level of the organic matter also belongs to the immature to highly over-mature [4]. For the samples Ai-01, Ai-13, Ha-01 and Ha-02, these two parameters of C29 steranes 20S/(20S+20R) and C29 steranes bb/(aa+bb) with values of > 0.4 and C31 homohopane 22S/(22S+22R) ratios > 0.5 reached equilibrium values indicating the thermal maturity level for coal sources [31]. However, for Wu-8, Wei-09 and Shinan-20, these parameters show values of less than 0.4 and 0.5, respectively. This suggests an immature to early mature range [4].

4.3 Pristane Isomerization Ratio (PIR) as a Highly Over-Mature Parameter

The Pristane Isomerization Ratios (PIR) = [6(R)10(R)+6(S)10(S)]/6(R)10(S) obtained from the different coal samples are presented in Table 3 with the values of the molecular maturity parameters measured by 20R-20S isomerization of sterane, 22R-22S isomerization of 17α(H)21β(H)- homohopane, Odd/Even Predominance (OEP) and the MPI. Figure 7 and Table 3 show that the

### TABLE 2
Results of Rock-Eval/TOC analysis and calculated parameters

<table>
<thead>
<tr>
<th>Sample</th>
<th>Tmax (°C)</th>
<th>S1 (mgHC/g)</th>
<th>S2 (mgHC/g)</th>
<th>PY (S1+S2)</th>
<th>TOC (%)</th>
<th>HI (mgHC/gTOC)</th>
<th>OI (mgCO2/g TOC)</th>
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<tr>
<td>Wu-8</td>
<td>418</td>
<td>2.15</td>
<td>75.2</td>
<td>77.35</td>
<td>63.70</td>
<td>118</td>
<td>78</td>
</tr>
<tr>
<td>Wei-09</td>
<td>435</td>
<td>1.95</td>
<td>60.50</td>
<td>62.45</td>
<td>73.30</td>
<td>82</td>
<td>11</td>
</tr>
<tr>
<td>Shinan-20</td>
<td>431</td>
<td>1.25</td>
<td>72.90</td>
<td>74.15</td>
<td>69.70</td>
<td>104</td>
<td>14</td>
</tr>
<tr>
<td>Ai-01</td>
<td>463</td>
<td>5.30</td>
<td>83.00</td>
<td>88.30</td>
<td>77.2</td>
<td>107</td>
<td>7</td>
</tr>
<tr>
<td>Ai-13</td>
<td>479</td>
<td>3.50</td>
<td>63.50</td>
<td>67.00</td>
<td>82.20</td>
<td>77</td>
<td>6</td>
</tr>
<tr>
<td>Ha-01</td>
<td>531</td>
<td>0.70</td>
<td>6.25</td>
<td>6.95</td>
<td>79.40</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>Ha-02</td>
<td>530</td>
<td>0.55</td>
<td>1.75</td>
<td>2.30</td>
<td>74.70</td>
<td>2</td>
<td>2</td>
</tr>
</tbody>
</table>

TOC = Total Organic Carbon; S1 = free hydrocarbons; S2 = pyrolyzable hydrocarbons; PY = Potential Yield; HI = Hydrogen Index; OI = Oxygen Index; Tmax = temperature of maximum S2.

### TABLE 3
Geochemical parameters of the coals studied in the Junggar Basin

<table>
<thead>
<tr>
<th>Sample</th>
<th>OEP</th>
<th>Isoprenoids</th>
<th>C29 xα 20S/(20R+20S)</th>
<th>C29 ββ/(α+ββ)</th>
<th>MPI-1</th>
<th>MPI-2</th>
<th>C31 αβ 22S/(22R+22S)</th>
<th>PIR</th>
<th>Ro (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wu-8</td>
<td>2.45</td>
<td>Pr/Ph</td>
<td>0.20</td>
<td>0.21</td>
<td>0.25</td>
<td>0.38</td>
<td>0.26</td>
<td>0.42</td>
<td>0.36</td>
</tr>
<tr>
<td>Wei-09</td>
<td>1.90</td>
<td>Pr/Ph</td>
<td>0.30</td>
<td>0.24</td>
<td>0.42</td>
<td>0.50</td>
<td>0.39</td>
<td>0.67</td>
<td>0.60</td>
</tr>
<tr>
<td>Shinan-20</td>
<td>1.19</td>
<td>Pr/Ph</td>
<td>0.45</td>
<td>0.33</td>
<td>0.57</td>
<td>0.64</td>
<td>0.45</td>
<td>0.74</td>
<td>0.80</td>
</tr>
<tr>
<td>Ai-01</td>
<td>1.12</td>
<td>Pr/Ph</td>
<td>0.51</td>
<td>0.43</td>
<td>1.13</td>
<td>1.30</td>
<td>0.58</td>
<td>0.81</td>
<td>1.25</td>
</tr>
<tr>
<td>Ai-13</td>
<td>1.09</td>
<td>Pr/Ph</td>
<td>0.54</td>
<td>0.38</td>
<td>1.76</td>
<td>2.11</td>
<td>0.60</td>
<td>0.84</td>
<td>1.53</td>
</tr>
<tr>
<td>Ha-01</td>
<td>1.01</td>
<td>Pr/Ph</td>
<td>0.51</td>
<td>0.44</td>
<td>0.73</td>
<td>0.85</td>
<td>0.61</td>
<td>0.90</td>
<td>2.32</td>
</tr>
<tr>
<td>Ha-02</td>
<td>1.07</td>
<td>Pr/Ph</td>
<td>0.53</td>
<td>0.41</td>
<td>0.31</td>
<td>0.49</td>
<td>0.59</td>
<td>0.97</td>
<td>2.99</td>
</tr>
</tbody>
</table>

OEP = (C23+6C25+C27)/4(C24+C26); Pr/Ph = pristane/phytane; MPI-1 = 1.5×(MP+3MP)/(P+1MP+9MP); MPI-2 = (3×2MP)/(P+1MP+9MP); PIR = 2.6,10,14-tetramethylpentadecane [6(R)10(R)+6(S)10(S)]/6(R)10(S); C29 xα 20S/(20R+20S) = C29-5α(H),14α(H),17α(H),24-ethylcholestane 20S/(20S+20R); C29 ββ/(α+ββ) = C29-5α(H),14α(H),17α(H),24-ethylcholestane ββ/(α+ββ); C31 αβ 22S/(22R+22S) = C31 17α(H),21β(H)-30-homohopane 22S/(22R+22S).
PIR in all the coal samples gradually increases; thermal maturity is represented by the $R_o$ values ranging from 0.36% to 2.99%. A comparison of the PIR with traditional maturity parameters (OEP, $C_{31}$ homohopanes 22S/(22S+22R), $C_{29}$ steranes 20S/(20S+20R), $C_{29}$ steranes $bb/aa$) was made to assess whether the PIR is directly related to maturity. The PIR increases from 0.42 to 0.97, which is similar to an increase in the ratio of $C_{31}$ homohopanes 22S/(22S+22R) from 0.26 to 0.61 (Tab. 3). Changes are relatively small within the OEP and MPI indices, which are part of the overly mature sections ($R_o > 2.0\%$); however, no linear correlations between OEP and $R_o$ or MPI and $R_o$ have been reported (Fig. 7). In addition, the MPI shows a reversal trend at higher levels of thermal maturity (Fig. 7). All of these phenomena suggest the possibility that the OEP, saturate biomarkers $C_{31}$ homohopane and sterane parameters and aromatic biomarkers MPI may have a limited application in highly mature samples. However, the PIR continues to increase with higher vitrinite reflectance values. Therefore, the PIR may be considered as a novel high-maturity index that can evaluate the maturation of coal samples in the Junggar Basin of China, and possibly other highly mature petroleum and coal samples.

Figure 6
Example of the variations in the degree of isomerization with $R_o$ in seven coal sample sequences in the Junggar Basin.

Figure 7
Comparison between a) MPI, b) OEP, c) $C_{29}$ sterane and $C_{31}$ homohopane and d) PIR ratios versus $R_o\%$.
The stereoisomer distributions indicate a loss of stereospecificity of the phytol-derived 6(R), 10(S) pristane, with an increase in geological maturation. For example, the least mature geological sample Wu-8 contains only 31% of the 6(R), 10(R) and/or 6(S), 10(S) isomers and 69% of the 6(R), 10(S) isomer. However, the over-mature sample Ha-02 contains 49% of the 6(R), 10(R) and/or 6(S), 10(S) isomers and 51% of the 6(R), 10(S) isomer (Tab. 3). These results suggest that isomerization located at the chiral centers in acyclic isoprenoid alkanes is enhanced with thermal maturation. In the over-mature samples Ha-01 and Ha-02 (R_o > 2.0), the isomerization differences of pristane isomers are still obvious, and the relative abundances of the 6(R)10(R) + 6(S)10(S) isomers show significant changes with an increase in R_o. These results suggest that the PIR is an appropriate indicator of maturity for the highly mature and over-mature coal samples in the Junggar Basin.

This case study shows that the PIR increases with increasing vitrinite reflectance (%R_o). This suggests an isomerization of the RS isomer into the RR and/or SS isomers. The ratio reached a maximum value of 0.97 in the samples examined. In a further study, it would be interesting to find a possible correlation between the vitrinite reflectance values (%R_o) and/or the maturity and the values of the [6(R)10(R) + 6(S)10(S)]/6(R)10(S) ratio.

In summary, the PIR may be of potential interest in that it can be used efficiently in measuring the maturity of highly mature coals and oils and may also be used beyond the oil window stage. However, the current work did not investigate the dependence of the PIR on kerogen type. It is necessary to further investigate and collect data of the PIR from other samples at other locations in order to improve upon this study.

CONCLUSIONS

High TOC contents, with high S_2 and HI values, indicate that Wu-8, Wei-09, Shinan-20, Ai-01 and Ai-13 have excellent source rock potential. For Ai-01, Ai-13, Ha-01 and Ha-02, the Pr/Ph ratios were between 0.5 and 1.0, indicating weak oxic-anoxic environments, which are possibly favorable for reducing conditions in the lacustrine sediments and microbial activity. However, for Wu-8, Wei-09 and Shinan-20, the Pr/Ph > 3.0 typifies terrigenous organic matter input under oxic conditions. The Pr/n-C_17 and Ph/n-C_18 values are smaller than 1.0 in Ai-01 and Ha-02 and this suggests that the microbial activity was less intense. In other coal samples, the Ph/n-C_18 values are smaller than 1.0; however, the Pr/n-C_17 values are greater than 1.0; this indicates that there was microbial activity, but not too strong. The T_max values (418-531°C), C_29 sterane 20S/(20S+20R) ratios (0.20-0.54) and C_31 homohopane 22S/(22S+22R) ratios (0.26-0.61) indicate an immature to highly over-mature stage of organic matter.

In addition, pristane enantiomers were completely separated and identified using GC-MS in extracts from the seven coal rocks of the Junggar Basin in China. The PIR (pristane isomerization ratio) ranges from 0.42 to 0.97 with an increase in thermal maturity evaluated from the vitrinite reflectance (%R_o) from 0.36% to 2.99%.

Pristane is an important molecular marker which can be used for petroleum geochemical investigations. The PIR could be used in combination with sterane and hopane maturity parameters, as well as a MPI, to measure hydrocarbon maturity from the immature (R_o < 0.6%), mature (R_o, 0.6-1.3%) and past mature (R_o, 1.3-2.0%) to over-mature stages (R_o > 2.0%).

An increase in the pristane [6(R)10(R) + 6(S)10(S)]/6(R)10(S) maturity parameter resulted from the relatively greater rate of generation of the 6(R)10(R) and 6(S)10(S) isomers, combined with the earlier decline in concentration (by isomerization) of the 6(R)10(S) component. The isomerization at the chiral centers in pristane is obvious as thermal maturation increases, and the PIR shows a significant increase at high maturity, due to the greater relative thermal stability of the 6(R)10(R) and 6(S)10(S) isomers.

The PIR can be used to measure the maturity of source rocks even at highly mature stages in the elevated vitrinite reflectance scale of maturity (%R_o) of 2.0-3.0%.

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