Lithium isotope compositions of U.S. coals and source rocks: Potential tracer of hydrocarbons

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ABSTRACT

Kerogen in organic-rich rocks contains trace amounts of lithium (Li) that has been overlooked as a contributor to the global Li geochemical cycle. This study examined a variety of coals where kerogen is concentrated (>50% organic carbon) and hydrocarbon source rocks of different ages, depositional environments and thermal maturity to determine their range of Li isotopic compositions (δ7Li‰) and factors that influence their compositions.

Using Secondary Ion Mass Spectrometry (SIMS), we analyzed 22 coals and 4 hydrocarbon source rocks (Types I, II, III), to determine the δ7Li of kerogen in situ, without chemical isolation of phases that can alter their original isotopic compositions. The δ7Li values of the coals surveyed are distinctly isotopically light (<0‰) compared to most natural minerals and fluids. In immature coals, with a vitrinite reflectance in oil (VRo) of ≤0.5%, kerogen δ7Li values average –23.4 ± 1.1‰ and become heavier with increasing thermal grade to temperatures of gas generation (VRo ~1.3%). The linear correlation between δ7Li and VRo suggests that 6Li may be preferentially released to pore fluid from kerogen during thermal maturation. Notably, authigenic clays forming at diagenetic temperatures substitute Li from pore fluids into silicate layers, therefore, the Li isotopic composition of the clays may record fluid isotopic compositions influenced by organic-Li sources.

NanoSIMS isotopic maps of the Lower Bakken Shale, and SIMS measurements of the Green River Shale show isotopically light Li associated with C-dominated areas, and heavier δ7Li with Si-dominated areas of the hydrocarbon source rocks. We conclude that kerogen is a source of isotopically light Li that contributes to fluids during thermal maturation and hydrocarbon generation. Kerogen may be a significant contributor of Li to pore fluids and its distinctly light Li isotopic composition relative to other terrestrial waters and minerals could provide a tracer of organic inputs to the global geochemical cycle.

1. Introduction

Lithium isotopes are being increasingly utilized to aid in understanding a host of terrestrial and extraterrestrial processes. In terrestrial systems, there has been significant use of Li isotopes in low temperature crustal environments to investigate the geochemical processes of weathering and fluid dynamics (Penniston-Dorland et al., 2017; Tomascak et al., 2016). However, there is limited information regarding how Li from organic compounds may play a role in these processes. If there is a significant amount of Li bound in organic compounds, it is important to know its isotopic composition and how Li from organics would contribute to the global geochemical cycle if the Li were released from the organics (e.g. with increasing burial temperature). In this study, we measured the Li isotope compositions on a suite of 22 coals from across the United States, representing different ages, depositional environments and thermal maturities. In addition, 4 hydrocarbon source rocks containing Type I, II and/or III kerogen were analyzed using Secondary Ion Mass Spectrometry (SIMS) and NanoSIMS. The goal of this study is to evaluate the range of Li isotopic compositions in organic-rich rocks consisting of a variety of organic macerals which were exposed to a range of thermal alteration conditions.

1.1. Lithium resources

Lithium is an important natural resource that has increased in demand due to the rising use of Li ion batteries. The primary economic sources for Li are brines, salt lakes and pegmatite deposits (Kesler et al., 2012). Seawater contains only ~0.17 mg/L Li but it is extracted economically using manganese oxide membranes to concentrate Li (Hong et al., 2018). Lithium is more concentrated in oilfield brines with concentrations up to several 100 mg/L (Collins, 1975; Eccles and Berhane, 2011; Macpherson, 2015; Millot et al., 2011; Phan et al., 2011).
2000). Interestingly, Li concentrations in bitumen (extractable organic compounds) are reported to be higher than their reservoir rocks by up to 17 mg/L, suggesting an association of Li with extractable hydrocarbons (Hosterman, 1990). In hydrocarbon source rocks, Li concentrations are highest in authigenic clay minerals (primarily illite-smectite; I–S) and some studies suggest that Li may be linked to gas generation (Williams et al., 2013, 2015). Lithium substitutes in the octahedral sites of I–S and is also adsorbed in clay interlayers (Sposito et al., 1999). Bentonite from hydrocarbon reservoirs in the Baltic Basin, contain I–S with Li contents >600 ppm after removal of interlayer-Li (Williams et al., 2013), whereas average marine shales have a reported range of 50–75 ppm Li (Horstman, 1957; Pendias and Kabata-Pendias, 2000). In Marcellus Shale source rock, Phan et al. (2016) found or-
range of 50–75 ppm Li (Horstman, 1957; Pendias and Kabata-Pendias, 2000) with Li contents >600 ppm after removal of interlayer-Li. Bentonite from hydrocarbon reservoirs in the Baltic Basin, contain I–S with Li contents >600 ppm after removal of interlayer-Li (Williams et al., 2013), whereas average marine shales have a reported range of 50–75 ppm Li (Horstman, 1957; Pendias and Kabata-Pendias, 2000). In Marcellus Shale source rock, Phan et al. (2016) found organs and sulfide minerals to be the second largest (up to 20%) host for Li after the silicate fraction. For coals, the worldwide average reported Li-content is lower than for clays, with ranges from 6 to 44 ppm (Finkelman et al., 2018; Ketriss and Yudovich, 2009; Swanson, 1975). However, some coals from China have been identified as containing economically viable concentrations of Li (Qin et al., 2015). Coals within the Jungar Coalfield (Mongolia) have Li concentrations exceeding 500 ppm but in those deposits, Li is mostly concentrated in authigenic clays within the coal (Dai et al., 2012).

1.2. Geologic background

Kerogen is the insoluble organic material found in sedimentary rocks (Durand, 1980) that is derived from the remains of organisms and is the most abundant form of organic matter on Earth (Vandenbroucke and Largeau, 2007). Kerogen is composed of a variety of organic macerals, which are grouped based on their physical properties. Just as rocks are composed of multiple minerals that each might host Li, the macerals that comprise kerogen may each host Li in different structural arrangements. Because of the complexity of organic macerals in kerogen, a simple approach to assessing major compositional changes was developed by Van Krevelen (1950) comparing their major element ratios (H/C and O/C) to classify organic Types (e.g., I, II, III) (Fig. 1). Coals contain primarily Type III kerogen derived from woody remains of plants that contain high O/C and low H/C (Vandenbroucke and Largeau, 2007). Coals are dominated by vitrinite (>70% by volume; Orem and Finkelman, 2003), which is a group of organic macerals that can indicate the thermal maturity according to its reflectance in oil immersion microscopy (VRo; Mukhopadhyay, 1994). Type III kerogen is more prone to generate gas than liquid hydrocarbons. Types I and II kerogen are primarily derived from algae and phytoplankton, containing higher H/C and lower O/C and are more prone to generate liquid hydrocarbons (Vandenbroucke and Largeau, 2007).

1.3. Lithium isotope measurements of kerogen

Lithium is a valuable tracer of fluids because of its mobility during surface weathering and under hydrothermal conditions (Millot et al., 2010; Pistiner and Henderson, 2003). Lithium isotopes are commonly used for tracing lithologic sources that contribute Li to fluids (e.g., Misra and Froelich, 2012; Williams et al., 2013, 2015; Phan et al., 2016), as different sources have a range of Li isotope ratios (Fig. 2).

In this study, we measured Li isotopic compositions in kerogen using solid-state secondary ion mass spectrometry (SIMS), because it has been recognized that reagents (acids) used to isolate kerogen (or insoluble organic matter (IOM); Cronin et al., 1987) from sediments also react with organic functional groups in kerogen, thus altering their composition (Saxby, 1976). To demonstrate this, Williams and Bose (2018) extracted kerogen from the Lower Bakken shale using standard 5 N HF-1 N HCl digestion (Durand, 1980) and compared the Li/12C ratio of the acid treated IOM (Li/12C = 0.024 ± 0.004; n = 5) to unextracted kerogen in polished section (Li/12C = 0.244 ± 0.020; n = 6). This indicates an average loss of 90% Li from the acid treated kerogen. Furthermore, NanoSIMS in situ measurements were made (using a 300 nm spot size) of Li/12C on kerogen in an argon ion polished section of the same Bakken sample. This sample preparation makes it easier to avoid beam overlap with silicates, and the average Li/12C ratio on kerogen was even higher (0.414 ± 0.020; n = 11) confirming significant loss of organolithium in the acid treated IOM. To our knowledge, this study is the first survey of Li isotopes in kerogen that has not been isolated from silicates by acid treatments, which likely alters their Li isotopic ratios.

2. Methods

2.1. Samples

Powdered coal samples (<0.25 mm) were obtained from the Pennsylvania State University (PSU) coal repository and the National Institute of Standards and Technology (NIST). The PSU samples included 8 Department of Energy Coal Samples (DECS) and 7 Argonne National Laboratory Premium Coal Samples (APCS). Additionally, 3 NIST coal samples, 4 coal samples from mines in the Rocky Mountain region of the U.S., and 4 hydrocarbon source rocks were acquired to extend the range of age, thermal grade and kerogen Types studied. The coal samples come from most of the major sedimentary basins in the U.S., range in VRo from 0.25–5.19%, and in age from the Carboniferous to the Tertiary. Table 1 lists the samples studied and source rock data are shown in Table 2.

Additional bulk coal samples were studied in thick polished sections
for comparison to the powdered samples. These samples came from the Williston Basin (Montana), Powder River Basin (Wyoming) and San Juan River Basin (New Mexico). Two of the whole rock samples are from the Fruitland Fm. of the San Juan River Basin and were selected specifically because of their high (up to 39 ppm) Li content (Bregg et al., 1998), relative to average sub-bituminous coals (6–7 ppm). Coal from the Powder River Basin was selected because of its low ash content, while the coal from Williston Basin was obtained to determine the reproducibility of Li isotope compositions of samples taken from separate locations within the same deposit (DECS-25 and the Savage Lignite).

Source rock shales (Table 2) were acquired from the US Geological Survey (Denver, CO) and were previously studied for B isotope compositions (Williams and Hervig, 2004). These rocks were selected to determine whether the Li isotopic composition varied with organic Type; Type I (Green River Fm.), Type II (New Albany Fm.), Type I/II (Bakken Fm. Lower Bakken) and Type III (Wilcox Fm.). The Green River Fm. (Type 1) is the type locality for lacustrine deposition (Bradley, 1931). The kerogen found in these deposits comes primarily from algal sources, with total organic carbon (TOC) up to 15.8% (Vandenbroucke and Largeau, 2007). The Type IINewAlbanyFm. Blochermember from the Eastern Interior Basin contains TOC up to 9.4% (Lewan et al., 1995) and represents a typical Type II kerogen marine shale, where planktonic organisms are the primary input (Vandenbroucke and Largeau, 2007). The Lower Bakken Fm. (Type I/II) from the Williston Basin has a variable TOC up to 35% (Jin and Sonnenberg, 2012) and contains inputs of both terrestrial and planktonic origin where it was deposited in an offshore ramp depositional environment (Albert, 2014). Wilcox Fm. (Type III) from the Gulf of Mexico Sedimentary Basin has a primary input of higher plants and was deposited in a wide range of depositional environments (lagoonal, fluvial, and deltaic) (Nichols and Traverse, 1971). The Wilcox sample we analyzed is a lignite with > 50% TOC.

**Fig. 2.** Natural variations in $\delta^7\text{Li}$ in literature (Bottomley et al., 2003, 1999; Chan et al., 2002; Clergue et al., 2015; Eccles and Berhane, 2011; Lemarchand et al., 2010; Macpherson et al., 2014; Millot et al., 2011; Négrel et al., 2010; Penniston-Dorland et al., 2017; Phan et al., 2016; Tomascak et al., 2016). Coal kerogen measurements are from this study and contain lower $\delta^7\text{Li}$ values than all other Li-reservoirs.
Table 1
Coal data from the Penn. State Coal Repository and Lithium isotope data from this study.

<table>
<thead>
<tr>
<th>Sample</th>
<th>State</th>
<th>Coal Type</th>
<th>VRo %</th>
<th>Li/C ppm</th>
<th>Li/C - 100</th>
<th>n</th>
<th>Organic Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 PA</td>
<td>Carboniferous</td>
<td>13.18</td>
<td>30.1</td>
<td>88.08</td>
<td>4.84</td>
<td>4.72</td>
<td>1.60</td>
</tr>
<tr>
<td>2 WY</td>
<td>Tertiary</td>
<td>8.77</td>
<td>45.7</td>
<td>80.73</td>
<td>5.20</td>
<td>10.11</td>
<td>1.43</td>
</tr>
<tr>
<td>3 IL</td>
<td>Carboniferous</td>
<td>15.48</td>
<td>45.7</td>
<td>80.73</td>
<td>5.20</td>
<td>10.11</td>
<td>1.43</td>
</tr>
<tr>
<td>4 PA</td>
<td>Carboniferous</td>
<td>9.25</td>
<td>40.8</td>
<td>84.95</td>
<td>5.43</td>
<td>6.90</td>
<td>1.68</td>
</tr>
<tr>
<td>5 VY</td>
<td>Carboniferous</td>
<td>4.77</td>
<td>19.0</td>
<td>91.81</td>
<td>4.48</td>
<td>1.66</td>
<td>1.34</td>
</tr>
<tr>
<td>6 UT</td>
<td>Cretaceous</td>
<td>4.71</td>
<td>47.8</td>
<td>81.32</td>
<td>5.81</td>
<td>10.88</td>
<td>1.59</td>
</tr>
<tr>
<td>7 WV</td>
<td>Carboniferous</td>
<td>19.84</td>
<td>36.2</td>
<td>85.47</td>
<td>5.44</td>
<td>6.68</td>
<td>1.61</td>
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<tr>
<td>8 ND</td>
<td>Tertiary</td>
<td>9.72</td>
<td>42.9</td>
<td>74.05</td>
<td>4.90</td>
<td>19.13</td>
<td>1.17</td>
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Department of Energy Coal Samples

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<th>Sample</th>
<th>State</th>
<th>Coal Type</th>
<th>VRo %</th>
<th>Li/C ppm</th>
<th>Li/C - 100</th>
<th>n</th>
<th>Organic Type</th>
</tr>
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<tbody>
<tr>
<td>1 TX</td>
<td>Tertiary</td>
<td>15.81</td>
<td>55.5</td>
<td>76.13</td>
<td>5.54</td>
<td>51.78</td>
<td>1.50</td>
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<td>2 CO</td>
<td>Cretaceous</td>
<td>5.37</td>
<td>28.2</td>
<td>87.78</td>
<td>5.85</td>
<td>4.03</td>
<td>1.76</td>
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<tr>
<td>3 UT</td>
<td>Cretaceous</td>
<td>5.84</td>
<td>46.9</td>
<td>81.72</td>
<td>6.22</td>
<td>10.10</td>
<td>1.56</td>
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<tr>
<td>21 PA</td>
<td>Carboniferous</td>
<td>11.15</td>
<td>3.9</td>
<td>91.87</td>
<td>3.91</td>
<td>2.92</td>
<td>0.81</td>
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<td>22 PA</td>
<td>Carboniferous</td>
<td>23.27</td>
<td>37.8</td>
<td>88.64</td>
<td>5.75</td>
<td>3.03</td>
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<tr>
<td>25 MT</td>
<td>Tertiary</td>
<td>11.85</td>
<td>46.9</td>
<td>75.64</td>
<td>5.15</td>
<td>17.72</td>
<td>1.09</td>
</tr>
<tr>
<td>34 PA</td>
<td>Carboniferous</td>
<td>7.35</td>
<td>40.4</td>
<td>85.41</td>
<td>5.55</td>
<td>6.38</td>
<td>1.72</td>
</tr>
<tr>
<td>39 WY</td>
<td>Tertiary</td>
<td>8.03</td>
<td>46.4</td>
<td>74.75</td>
<td>5.43</td>
<td>18.87</td>
<td>1.01</td>
</tr>
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National Institute of Standards and Technology Samples

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<th>Sample</th>
<th>State</th>
<th>Coal Type</th>
<th>VRo %</th>
<th>Li/C ppm</th>
<th>Li/C - 100</th>
<th>n</th>
<th>Organic Type</th>
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<td>1632c</td>
<td>Carboniferous</td>
<td>7.16</td>
<td>nr</td>
<td>77.45</td>
<td>5.11</td>
<td>nr</td>
<td>1.50</td>
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<tr>
<td>2682b</td>
<td>Tertiary</td>
<td>6.32</td>
<td>48.5</td>
<td>68.43</td>
<td>4.88</td>
<td>16.24</td>
<td>1.37</td>
</tr>
<tr>
<td>2684b</td>
<td>Carboniferous</td>
<td>9.72</td>
<td>36.4</td>
<td>69.25</td>
<td>4.59</td>
<td>nr</td>
<td>1.45</td>
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</table>

Additional coal samples of this study

<table>
<thead>
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<th>Sample</th>
<th>State</th>
<th>Coal Type</th>
<th>VRo %</th>
<th>Li/C ppm</th>
<th>Li/C - 100</th>
<th>n</th>
<th>Organic Type</th>
</tr>
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<tr>
<td>San Juan</td>
<td>Cretaceous</td>
<td>16.77</td>
<td>40.05</td>
<td>69.90</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
</tr>
<tr>
<td>Four Corners</td>
<td>Cretaceous</td>
<td>15.78</td>
<td>38.12</td>
<td>70.30</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
</tr>
<tr>
<td>Savage</td>
<td>Tertiary</td>
<td>4.89</td>
<td>47.26</td>
<td>71.20</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
</tr>
<tr>
<td>Black Thunder</td>
<td>Tertiary</td>
<td>9.42</td>
<td>41.90</td>
<td>72.71</td>
<td>nr</td>
<td>nr</td>
<td>nr</td>
</tr>
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</table>

Table 2
Lithium isotope results for U.S. source rocks of different organic Types.

<table>
<thead>
<tr>
<th>Sample</th>
<th>State</th>
<th>Age</th>
<th>VRo %</th>
<th>δ7Li ‰ ± S.E.</th>
<th>n</th>
<th>Organic Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wilcox Fm. Lignite</td>
<td>TX</td>
<td>Lower Eocene</td>
<td>0.45</td>
<td>-19.8 ± 2.4</td>
<td>4</td>
<td>III</td>
</tr>
<tr>
<td>Bakken Fm. Lower Bakken Mbr</td>
<td>ND</td>
<td>Mississippian</td>
<td>0.50</td>
<td>-8.9 ± 1.9</td>
<td>8</td>
<td>II</td>
</tr>
<tr>
<td>Green River Fm.</td>
<td>WY</td>
<td>Eocene</td>
<td>0.65</td>
<td>-15.0 ± 3.6</td>
<td>4</td>
<td>I</td>
</tr>
<tr>
<td>New Albany Fm. Blocher Mbr</td>
<td>IL</td>
<td>Mississippian</td>
<td>0.95</td>
<td>-0.9 ± 2.2</td>
<td>8</td>
<td>II</td>
</tr>
</tbody>
</table>

Lithium isotope compositions determined by IMS-6f, where n = number of analyses on each sample; S.E. = 1σ standard error; nr = not reported; VRo - vitrinite reflectance values %; VRo values for Wilcox, Green River and New Albany shales are estimated from: ¹Mukhopadhyay, 1994; ²Canter et al., 2016; ³Pawlewicz and Finn, 2002; ⁴East et al., 2012; ⁵Lewan et al., 1995; ⁶Nuccio and Hatch, 1996.
2.4. Secondary ion mass spectrometry

Two different SIMS instruments, the Cameca Ametek Ion Mass Spectrometer (IMS-6f) and NanoSIMS 50 L (NanoSIMS) were used to obtain in situ Li isotopic ratios of samples. Both instruments have duoplasmatron sources which generate a plasma beam of O− ions. The primary beam is focused to a spot on the sample surface. This high energy ion bombardment sputters the sample surface ejecting ions, atoms, and electrons (Hervig, 1996). Positive secondary ions, such as 7Li+ and 6Li+, are transmitted through a series of lenses and apertures in the secondary column and separated by energy in an electrostatic analyzer. Ions are differentiated by their mass (m) to charge (z) ratio in the mass spectrometer and are detected using either an electron multiplier or faraday cup. For our analyses, all ions were detected using electron multipliers, calibrated using internal reference standards. Mass interferences must be eliminated by using energy filtering (for molecular interferences) or by adjusting the entrance and exit slits (or apertures) to the mass spectrometer to increase the mass resolving power (MRP = mass/Δmass). Primary interferences were 24Mg2+ and 11BH+ interfering with 12C+. The 14N2+ signal was easily resolved from 2Li, and sufficient MRP was used on both instruments to resolve m/z interferences.

To determine the instrumental mass fractionation (IMF) we used several standard reference materials (NIST SRM 610, NIST SRM 612), and illite from the Clay Minerals Society Source Clay Repository (IMt-1; www.clays.org). Each of these standards verified consistent values of IMF (mean daily S.E. ± 1.0‰, 1σ) and standard bracketing of unknowns, with one or more of these internal reference materials, was used to monitor instrumental drift during each analytical session (See Supplement material (SM) Fig. S1).

SIMS measurements are subject to matrix effects, where ions sputter at different rates (cps) from different matrices (Burnett et al., 2015; Hauri et al., 2006; Seah and Shard, 2018). We acknowledge that compositional differences between standards and unknowns (e.g., silicates vs. organics) may lead to matrix effects on the ionization process, which could shift the isotopic ratios measured (Hauri et al., 2006). For example, Bell et al. (2009) found that changes in Mg-content of olivine affected the δ7Li values by 1.3‰ per mole Mg substituted. There is not yet a kerogen standard with known Li isotopic composition verified by alternative analytical techniques. However, we suggest that δ7Li values within a kerogen Type class (I, II or III) will be comparable as their major element compositions are similar (e.g., Ishida et al., 2018). Furthermore, we found no correlation between δ7Li and the C-H-O contents of measured Type III kerogen in coals (SM; Figs. S6-S8). There is also no standard for calibration of Li contents in kerogen matrices but the 7Li/12C+ intensity ratio can be used to compare relative abundances among similar phases (e.g., Type III kerogen).

2.4.1. IMS-6f analytical protocol

The primary beam current and ion optics determine the beam diameter and therefore the analysis spot size. For the IMS-6f, the duoplasmatron source generates a primary beam of O− ions at ~12.5 kV and the sample is held at +9.0 kV for a total impact energy of 21.5 kV. The spot size can be as small as a few microns (~5 μm) in diameter, but in samples with low Li content, higher currents were necessary to provide enough Li+ secondary ions counts for statistical significance. Generally, a 5–20 nA primary current yielding a beam diameter of < 50 μm was used for spot analyses on all samples measured by IMS-6 f. Ions with an energy range ± 20 eV were allowed into the secondary magnet where the magnetic field can be adjusted to select the m/z of interest. For the IMS-6f analyses, each sample was analyzed on multiple spots as indicated in Tables 1 and 2 (n = 2 to 33). The IMF measured before and after analysis of unknowns was used to correct the raw isotope ratios.

2.4.2. NanoSIMS analytical protocol

A duoplasmatron source was also used for NanoSIMS measurements, generating a plasma beam of O− ions at ~−8 kV. The sample is held at +8 kV for a total impact energy of 16 kV. The NanoSIMS design uses a smaller primary beam of tens to hundreds pA of O−, to achieve greater spatial resolution than the IMS-6 f. The advantage of NanoSIMS is the ability to make ion images of the distribution of isotopes over a small area (typically 5–25 μm2) and at a greater spatial resolution. This allows spatial separation of measurements in the kerogen matrix avoiding silicate minerals (e.g., clays) in the kerogen matrix.

For the NanoSIMS analyses, ion maps were made in the imaging mode by rastering the beam over 25μm2 areas to measure the distribution of 7Li+, 12C+ and 30Si+ (cps). Images were taken at two different pixel resolutions (256 x 256 and 512 x 512) and varying spatial resolutions with beam diameter ranging from ~300 nm to 2 μm (3 pA – 500 pA primary current). All images were taken with a counting time of 1000 μs/pixel. NanoSIMS real time imaging was used in conjunction with the ion maps to pinpoint regions of interest (ROI) within the rastered area where the 12C+ or 30Si+ signals were high. Maps of 14N+, 24Mg+, 39K+, and 40Ca+ ions were also made to help identify the matrix being measured, e.g., determining if a spot with relatively high 12C+ counts was kerogen or a carbonate mineral, differentiating K-bearing clays vs. quartz, and to confirm that the 24Mg+ m/z was not interfering with the 12C+ m/z measurements. Using a primary beam current of 100–500 pA with a beam diameter of 1–3 μm we measured Li isotopic ratios in areas of high-12C+ versus high-30Si+ (cps). Using this protocol, we were able to separate analyses of kerogen and silicate phases and acquire their Li isotopic compositions at high spatial resolution.

3. Results

A histogram of all measurements of δ7Li by IMS-6f and NanoSIMS instruments for coals and source rocks is shown in Fig. 3. Multiple SIMS measurements on each sample were averaged and are summarized in Tables 1 and 2. The individual analyses and statistical evaluation can be found in Table S1. The δ7Li values measured are mostly < 0‰, except where analyses possibly overlapped silicates. We also characterized spot measurements according to their 12C+/30Si ratios and defined Si-rich areas where 12C+/30Si < 1.0, while C-rich areas ranged from 12C+/30Si = 1 to 12.000. For Si-rich areas (n = 28) the average δ7Li is
−7.3 ± 1.7‰, and for C-rich areas (n = 130) the average δ7Li is −23.6 ± 1.1‰ (Fig. S9). Each coal sample (Table 1) was measured in multiple spots (n = 2–33) and the mean and standard error of the mean for all analyses on each sample is plotted in Fig. 4. The variability in measurements from spot to spot within a single sample may be due to heterogeneity of the various organic macerals, beam overlapping multiple phases (kerogen and silicates), sample charging, or surface roughness of pressed powders. Nonetheless, the average analytical error on each spot analysis was 1.1‰ (1σ) which is far less than the observed range of values (SM; Table S1).

The source rock samples from the Green River Fm., New Albany Fm. and Wilcox Fm were measured by IMS-6f and the Lower Bakken shale sample was measured by both IMS-6f and NanoSIMS. Given the range of VRo values reported for these source rocks (Table 2), the more thermally mature New Albany shale has the highest δ7Li values, while the less mature source rocks have lower δ7Li values on average. Spot analyses in the Green River Fm. (Type I) showed a correlation (R² = 0.93) between 7Li/30Si ratios and 12C/30Si ratios (Fig. 5) with isotopically lighter values observed in C-rich spots.

### 4. Discussion

#### 4.1. Coal survey of δ7Li

Few studies have examined the organic vs. mineral hosts of Li in coal, although some bulk measurements of Li in coals have reported that Li is hosted by the silicates or “ash” portion (Finkelman, 1980; Dai et al., 2012). Our in situ study of Li in coals shows no correlation between δ7Li and the contents of elements commonly associated with ash (Si, Al, K) for measured APCS and DECS coals (SM; Figs. S3-S5). Swaine (1990) stated that Li is partly associated with organic matter, and Finkelman et al. (2018) later reported up to ~50% of Li in immature coals may be associated with organic material. Terrestrial plant matter that is the primary component of coals is one possible source of organolithium compounds. Gough et al. (1977) found the Li content of plants to be positively correlated with the Li content of the soils in which they grew. Li has a varying toxicity threshold among plants depending on the species, yet plants contain trace amounts of Li (0.01–31 ppm) on average (Pendias and Kabata-Pendias, 2000). Leguminous plants grown in Li rich soils (9–175 ppm) in New Zealand are reported to contain up to 143 ppm Li (Pendias and Kabata-Pendias, 2000). Beets can take up even higher amounts of Li (up to 5500 ppm) from Li-rich soils and have been proposed for agro-mining (Kavanagh et al., 2018). Even though Li is present in small concentrations in most plants, it can become concentrated during the formation of kerogen due to the breakdown of biomolecules and consolidation of plant material during burial and lithification. Littke and Leythauser (1993) found that a 1-m thick layer of peat generates ~20 cm of bituminous coal.
therefore 3 ppm Li in peat can produce a coal with 15 ppm Li.

The range of Li isotopes published for various terrestrial and extraterrestrial samples is summarized in Fig. 2 (Clergue et al., 2015; Lemarchand et al., 2010; Négrel et al., 2010; Penniston-Dorland et al., 2017; Tomascak et al., 2016). The coals measured in this study represent the lowest δ7Li values measured, typically <0‰ for the most organic-rich samples. This range of δ7Li values is similar to the range of δ11B values measured on these same coal samples (Williams and Hervig, 2004). The low δ7Li values in low rank coals suggest that immature organic macerals in the coal are enriched in 6Li compared to most rocks, minerals and waters evaluated in the global geochemical cycle (Fig. 2).

Consistent with our Li isotope results for kerogens, other studies of Li isotopes in organic materials have reported isotopically light δ7Li values. Lemarchand et al. (2010) found forest soil solutions as well as fir tree needles, branches and roots to δ7Li values lower (−5 to −17‰) than the parent granite (+0.3‰), soil (+0.7 to +3.3‰) and stream waters (+5 to +20‰) in a forested granitic catchment in France. Another study on peat from the Massif Central in France (Négrel et al., 2010)
2010) reported $\delta^7$Li values of −11 to 0‰.

Only two studies have reported $\delta^7$Li of coal constituents. Harkness et al. (2015) measured $\delta^7$Li values from −7 to +12.68% for aqueous leachates of the inorganic constituents of coals from the Powder River, Appalachian and Eastern Interior basins. He et al. (2019) used microwave-digested coal samples from the high Li, (> 500 ppm) Guanbanwusu mine (China) and reported $\delta^7$Li values from +6 to +8% on residues measured by ICP-MS. The anomalously high Li contents of this coal have been shown to be positively correlated with the ash content suggesting secondary Li enrichment from a hydrothermal source (Sun et al., 2012). Therefore, the somewhat heavier lithium isotope compositions reported for this Guanbanwusu coal are likely controlled by the silicate host minerals.

Our results show a clear positive correlation ($R^2 = 0.76$) between $\delta^7$Li of coals and thermal maturity as indicated by VRo (Fig. 4). Low rank coals (VRo < 0.5%) have the lowest $\delta^7$Li values (average = 23.4 ± 1.1‰) and the Li isotope composition becomes heavier (up to +5‰) with increasing rank up to VRo ~1.3% (bituminous), irrespective of age and depositional environment (Table 1). The observation that $\delta^7$Li values of kerogen in coals become higher as thermal maturity increases may be interpreted as either a preferential release of $^6$Li to pore fluids as temperature increases, or an uptake of $^3$Li from hydrothermal fluid during thermal maturation. However, no clear correlation was observed between Li/C and either VRo or $\delta^7$Li (SM; Fig. S2). Given that these samples each come from many different sedimentary basins of varying geological ages, they each have their own initial Li contents and diagenetic histories that would affect the Li/C ratios. Kerogen from different ages spanning 300 Myr may come from plants with varying Li contents. Considering that a positive trend is shown among all coals studied, it is most likely that the trend reflects a preferential loss of $^6$Li rather than incorporation of exogenous $^6$Li. While hydrothermal Li certainly influences some basins (e.g., Dai et al., 2012), if this were the case ubiquitously, we would expect higher Li/C ratios in hydrothermally influenced (more mature) samples, which is not observed.

The trend of increasing $\delta^7$Li with thermal maturity is consistent with a previous study showing $^6$Li enrichment in authigenic clays associated with the migration of gas into the Wattenberg gas field, Denver Basin (Williams et al., 2015). That study proposed that $^6$Li preferentially leaves organolithium compounds during gas generation from kerogen (VRo > 1.5%). Different kerogen Types generate hydrocarbons (oil and gas) over a range of burial temperatures. The ‘oil window’ spans a VRo range from 0.5 to 1.35‰ (Hunt, 1996; Makhpoapdhyay, 1994; Peters and Cassa, 2007; Pittion and Goudain, 1985; Taylor et al., 1998; Tissot et al., 1987; Tissot and Welte, 1984). On the other hand, Type III kerogen (coals) have an effective ‘oil window’ at VRo values of 0.65–2.0% with gas generation dominant above VRo 1.5% (Petersen, 2006). The trend in $\delta^7$Li values of coals (Fig. 4) clearly coincides with the temperatures of hydrocarbon generation from kerogen, supporting a preferential release of $^6$Li into hydrocarbon-related fluids.

Interestingly, we measured two over mature samples APCS-6 from UT (VRo 1.68%) and DECS-21 from PA (VRo 5.19%) that fell off the trendline demonstrated by other coals. These samples show low $\delta^7$Li values (~22 ± 0.8 and ~39 ± 4.0‰, respectively; Table 1). Clearly at very high temperatures, the coal remaining is largely residual C and ash. It is possible that at very high thermal maturity, most of the organolithium compounds have been degraded (released) during hydrocarbon generation, and that any remaining Li is hosted by silicates, or residual C.

### 4.2. Hydrocarbon source rocks

Oil and gas are generated from source rocks at temperatures between ~60–150 °C, generally at burial depths from 1 to > 4 km. The hydrocarbons can be expelled from source rocks and migrate along pathways through permeable rock into structural and stratigraphic traps, forming conventional hydrocarbon reservoirs. Black shale source rocks can also host oil and gas forming unconventional reservoirs. Therefore, a distinctly light Li isotopic composition of pore fluids, suggesting influx of hydrocarbon-sourced Li, may be useful in identifying productive intervals of a shale. We studied several well-known hydrocarbon source rocks to compare the Li isotopic compositions of their different kerogen Types.

It was recognized while measuring source rocks with the IMS-6f instrument, that the large (up to 50 μm) primary ion beam diameter on the IMS-6f and the predominance of silicates over kerogen in source rocks influenced analyses where kerogen and silicate analyses overlapped. This resulted in slightly more positive $\delta^7$Li values in the shale source rocks than coals (Fig. 3), which are dominated by carbonaceous material (ASTM Committee, 2019), thus decreasing overlap on minerals. Primary beam overlap on silicates was avoided using the NanoSIMS on the Bakken shale sample, so that Li isotope measurements could be compared between kerogen and mineral phases (Section 3.1). The Mississippian Lower Bakken shale was studied in detail because it is a Type I/II source rock with high total organic carbon (up to 35 wt%; Jin and Sonnenberg, 2012). NanoSIMS maps of the Bakken shale (Fig. 6) and IMS-6f measurements of Green River Fm. (Fig. 5) showed that the C-rich areas have isotopically lower $\delta^7$Li values than Si-rich areas of the source rocks. Like the coal measurements, immature kerogen in source rocks contains isotopically light Li (Table 2). The general trend for source rock kerogen similarly shows $\delta^7$Li increasing with thermal maturity over the temperature range of oil and gas generation.

### 4.3. Implications for Li geochemical cycles

The results presented here support the idea that Li derived from kerogen can be used to trace organic-related fluids because the Li isotopic composition of pore waters from kerogen should be lighter than most other natural waters (Fig. 2). Oilfield brines generally range from $\delta^7$Li +4 to +16‰ (Eccles and Berhane, 2011; Macpherson et al., 2014; Millot et al., 2011; Phan et al., 2016), except where evaporative processes enrich the brines in $^7$Li (Chan et al., 2002) (SM Table S2). These brines are lighter than seawater ($^7$Li ~ 31‰) (Chan and Edmond, 1988; Misra and Froelich, 2012), rivers (mean = +23‰), and lakes (+17 to +36). Hydrocarbon-related fluids contain elevated contents of many organically derived trace elements (e.g., Li, B, N) and many of these elements are substituted into authigenic clays that form under diagenetic conditions affecting coals and hydrocarbon sources (Williams et al., 2012, 2013, 2015). Given that up to 50% of Li in immature coals may be associated with kerogen (Finkelman et al., 2018) and the observation that more mature bituminous coal contains Li associated with the silicates (ash) (Finkelman et al., 2018), we hypothesize that the Li is released from dispersed organic matter and becomes concentrated in authigenic clays. Among coals surveyed by Finkelman et al. (2018), 6 immature coals contained on average only ~6 ppm Li while 14 bituminous (mature) coals contain on average 44 ppm Li. If Li accumulates in pore fluids during burial, then the pore fluids become enriched in the Li released from organolithium complexes in the source kerogen (Fig. 7).

The isotopic fractionation as Li is released from organic compounds in kerogen to pore fluid is unknown, but the isotopic fractionation of Li between water and authigenic clays (smectite-illite) is well-known (Wunder et al., 2007; Vigier et al., 2008; Williams et al., 2012). Therefore, where temperatures can be constrained, assuming equilibrium conditions, the Li isotopic composition of the fluid can be calculated. For example, the $\delta^7$Li of the high Si area in the Bakken shale (Fig. 6; Area 1; ROI-2) gave a value of ~5.5‰. High $^{13}$C in this area suggests that this may be illite (or possibly a detrital feldspar). Using the isotopic fractionation equation determined for illite-water fractionation:
1000 \ln \alpha = 6.15 - 0.07(1000/T (K))  \tag{2} 

(Williams et al., 2012) and estimating a temperature \( \sim 80 \, {^\circ}C \) based on the vitrinite reflectance of the Bakken immature kerogen, one can calculate a pore fluid \( \delta^7\text{Li} \) value of +14.5‰, which is in the range of other natural ground waters (Fig. 2). Realistically, the measured \( \delta^7\text{Li} \) measured (Fig. 6; Area 1; ROI-2) includes interlayer Li in addition to structurally bound Li in the clay, so this value may be slightly heavier than the water that equilibrated with the authigenic clay (Williams et al., 2015). Evaluating the isotopic fractionation of Li between kerogen and water will be useful in refining the potential use of Li isotopes for tracing hydrocarbon migration paths through organic-rich shales.

Another important implication for this work is its impact on interpretations of continental weathering based on \( \delta^7\text{Li} \) values. For example, the Li isotopic composition of seawater over the Cenozoic era has been interpreted to have been lighter in the past due to high rates of continental weathering based on the \( \delta^7\text{Li} \) of foraminifera (Misra and Froelich, 2012) and other biogenic carbonate (Sun et al., 2018). While the trend of decreasing \( \delta^7\text{Li} \) in foraminifera over the Cenozoic era is robust, the effects of diagenesis on the re-equilibration of carbonate \( \delta^7\text{Li} \) with diagenetically altered fluids, might warrant reconsideration in light of our data showing distinctly light Li isotopic compositions of kerogen. It is possible that increased nutrient supplies to the ocean, associated with high erosion rates, caused eutrophication and hypoxia, and these conditions led to preservation of sedimentary organic matter (Gallois, 1976; Liu and Wang, 2013; Loftus and Greensmith, 1988). Therefore, the low \( \delta^7\text{Li} \) values recorded by deeply buried forams, where diagenesis is likely, may be reflecting the isotopic composition of organic-derived Li on paleofluid compositions. Understanding the exchange of Li between organic matter, fluids and minerals in sedimentary basins contributes not only to our understanding of hydrocarbon generation and migration, but also to the evaluation of Li sources important to the global Li geochemical cycle.

5. Conclusions

Our survey of Li isotopic compositions in organic rich rocks of varying depositional environment, age, and thermal maturity leads to the following insights:

1. Immature kerogen measured in this study has a lighter Li isotopic composition than any other crustal material (mineral or fluid) yet reported.
2. With increasing thermal maturity, from immature to mature (up to VRo 1.3%), the \( \delta^7\text{Li} \) values of kerogen increase. We conclude that \( ^6\text{Li} \) is preferentially released from the organic matrix to pore fluids during thermal maturation.
3. The organic matrix in hydrocarbon source rocks has a lighter Li isotopic composition than the silicate matrix.
4. If Li released from kerogen is isotopically light relative to most pore fluids, then the uptake of Li from these fluids by authigenic clays will record the isotopically light Li related to hydrocarbon generation.
5. The recognition of isotopically light Li derived from organic sources could impact interpretations of continental weathering and should be further investigated for its significance in the global geochemical cycle.

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Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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