Interpretation of Nitrogen Isotope Profiles in Petroleum Systems: A Review

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Bulk sedimentary nitrogen isotope profiles are often used as proxies for depositional redox conditions, nitrogen cycling, and nutrient uptake in modern and ancient marine systems. The general preference in terms of analysis is that the sediments measured have undergone minimal thermal alteration, as post-depositional processes might have altered the initial $\delta^{15}N$ signal, thus complicating the interpretation of these records. Although not a traditional proxy for petroleum evaluation purposes, recently the use of nitrogen isotopes in petroleum systems has been investigated as potential proxies to reconstruct paleoenvironmental conditions such as redox, and for organic matter evaluation. In this paper we review the use of nitrogen isotope data in petroleum systems, their interpretations, and factors that may complicate their use as proxies. We review the evidence for nitrogen isotopic fractionation during diagenesis, catagenesis, and fluid migration as determined by lab experiments, and how these might impact interpretation of $\delta^{15}N$ data in petroleum systems. We also analyze the use and interpretation of $\delta^{15}N$ data from petroleum-producing reservoir units, including unconventional reservoirs and lacustrine basins. Lastly, we discuss potential applications for nitrogen isotopes in petroleum systems with regards to their use as both geochemical proxies and as tools to evaluate petroleum reservoirs.

Keywords: nitrogen isotopes, petroleum systems, diagenesis, redox, catagenesis, organic matter transformation

INTRODUCTION

Nitrogen isotopes are being used in an increasing number of recent sedimentary system studies as paleoenvironmental proxies. Nitrogen isotope ($\delta^{15}N$) measurements of sediments have been shown to record processes occurring in the water column, allowing them to serve as proxies for paleoredox conditions and nitrogen cycling (Altabet and Francois, 1994). As a result, sedimentary $\delta^{15}N$ data have been used to investigate the causes of mass extinctions, the extent of ocean anoxic events, and changes in oxygen minimum zones on glacial-interglacial time scales (e.g., Ganeshram et al., 1995; Calvert et al., 2001; Jenkyns et al., 2001; Quan et al., 2008; Ryabenko et al., 2012). Sedimentary $\delta^{15}N$ profiles have also been used to identify intervals and locations where the nitrogen cycle and nutrient regimes appear to have been significantly altered from that for the modern-day, including Precambrian conditions and Mediterranean sapropel deposits (e.g., Godfrey and Falkowski, 2009; Higgins et al., 2010; Ader et al., 2016). These evaluations of paleoredox conditions and changes in the nitrogen cycle can provide critical evidence for the occurrence of larger biogeochemical cycle changes or geological events,
such as emplacement of large igneous provinces, climate change, changes in export production, or the timing of the Great Oxygenation Event (e.g., Holloway and Dahlgren, 2002; Meyers et al., 2009; Algeo et al., 2014; Ader et al., 2016; Danzelle et al., 2020).

Nitrogen is an essential bionutrient for all living creatures, and is a critical component of many biological compounds, including proteins, enzymes, and nucleic acids. Nitrogen can also be found in a range of oxidation states ranging from -III in NH$_3$ and organic N compounds to + V in NO$_3^-$.

Nitrogen has two isotopes: $^{14}$N (99.63% abundance) and $^{15}$N (0.37% abundance); due to the much higher abundance of $^{14}$N, measured isotopic values are reported as $\delta^{15}$N in parts per thousand (per mil; %o):

$$\delta^{15}N = \left( \frac{^{15}N/^{14}N}_{\text{sample}} - 1 \right) \times 1000 \quad (1)$$

The reference standard is atmospheric nitrogen gas, set by definition to $\delta^{15}N = 0$%o. Measured samples that are depleted in $^{15}$N relative to the standard have negative $\delta^{15}$N values, while samples that are relatively enriched in $^{15}$N have positive $\delta^{15}$N values. Transformation of nitrogen between oxidation states and between organic and inorganic forms is predominantly mediated by organisms. Due to differences in bond strengths between $^{14}$N and $^{15}$N, isotopic fractionation occurs during these reactions, and therefore nitrogen isotopes can be used to provide information about the reactions involved and the larger nitrogen cycle. The

![Figure 1](image-url)
fractionation factor ($\alpha$) for any reaction can be calculated as the difference in $^{15}$N/$^{14}$N ratios between reactants ($R$) and products ($P$):

$$\alpha_{R \rightarrow P} = \frac{\left(\frac{^{15}N}{^{14}N}\right)_R - \left(\frac{^{15}N}{^{14}N}\right)_P}{\left(\frac{^{15}N}{^{14}N}\right)_P}$$

(2)

If no fractionation takes place, $\alpha = 1$; reactions that favor the light isotope have $\alpha > 1$, and reactions that favor the heavier isotope have $\alpha < 1$. Similarly, kinetic fractionation can be assessed using the kinetic isotope effect ($\epsilon$), which is the difference in rates of reaction for each isotope. For reactions taking place in natural systems, $\epsilon$ is essentially the difference between the $^{15}$N value of the reactant and that of the instantaneous product formed.

**Figure 1A** shows a schematic of isotope effects for nitrogen reactions that are important to the geologic aquatic nitrogen cycle, along with their associated fractionation factors ($\alpha$) and isotope effects ($\epsilon$) (Delwiche and Steyn, 1970; Wada et al., 1980; Sigman and Cassiotti, 2001; Talbot, 2001; Brunner et al., 2013). Measured fractionation factors for nitrogen processes may vary depending on the reaction pathway, culture or field measurement, and the specific species measured (Delwiche and Steyn, 1970; Wada et al., 1980 and references therein; Talbot, 2001 and references therein). If a reaction proceeds to completion, no fractionation is observed.

The interpretations of $\delta^{15}$N data with regards to paleoredox conditions and nitrogen cycling rely on the fractionation factors for different nitrogen reactions (Figure 1A), which are in turn set by environmental conditions, particularly oxygen concentrations. The details have been discussed in several papers (e.g., Delwiche and Steyn, 1970; Wada et al., 1980; Altabet and Francois, 1994; Talbot, 2001; Robinson et al., 2012), but in brief, the sedimentary $\delta^{15}$N values reflect the $\delta^{15}$N of sinking organic matter, which is in turn influenced by the nitrogen reactions and environmental conditions occurring in the water column. In general, higher sedimentary $\delta^{15}$N values are representative of low oxygen/suboxic water column conditions, because these conditions favor strongly fractionating denitrification and anammox reactions that result in organic matter enriched in $^{15}$N, which is recorded in the sediments as higher $\delta^{15}$N values (Altabet and Francois, 1994; Robinson et al., 2012 and references within). Lower $\delta^{15}$N values can represent either anoxic systems dominated by nitrogen fixation or ammonium assimilation, or oxic systems utilizing nitrate as the main source of nitrogen; the nitrogen reactions for both of these conditions have smaller fractionation factors, and therefore generate organic matter depleted in $^{15}$N. The conceptual model of how sedimentary $\delta^{15}$N values vary with oxygen levels is shown in Figure 1B.

Since sedimentary $\delta^{15}$N values represent the average of all nitrogen processes occurring in the water column at a particular location, nitrogen isotopes are a local proxy. Nitrogen isotope values in sedimentary systems also reflect the $\delta^{15}$N values of the initial nitrate or ammonium used to produce the organic matter, so variations in the $\delta^{15}$N record may reflect the input of different sources of nitrogen into the system (Altabet and Francois, 1994; Robinson et al., 2012 and references within). Therefore, correlating specific $\delta^{15}$N values to exact O$_2$ concentrations can be difficult, particularly for historical records.

One of the main concerns in the interpretation of sedimentary $\delta^{15}$N records is whether the sedimentary $\delta^{15}$N values measured accurately reflect the water column processes, or whether there is alteration of the record either during the initial organic matter sedimentation or through post-depositional processes. In brief, loss of nitrogen compounds could result in altering the original $\delta^{15}$N signal through removal of compounds that are either enriched or depleted in $^{15}$N (Robinson et al., 2012 and references within). This could result in measured $\delta^{15}$N values that do not correspond to the original environmental conditions and nitrogen reactions. Remineralization of nitrogen compounds is most likely to occur at locations at which the water column and sediments are predominantly oxic, sedimentation rates are particularly slow, or productivity is low, so use of nitrogen isotopes as proxies in these environments may not be straightforward. Isotopic alteration may also occur at the sediment-water interface, with some evidence for preferential loss of nitrogen compounds with lower $\delta^{15}$N values, resulting in a shift to higher $\delta^{15}$N values measured in the sediment (Freedenthal et al., 2001; Robinson et al., 2012 and references within). Other processes that may alter the initial $\delta^{15}$N signal include input of NH$_4^+$ or terrestrial nitrogen from outside sources into the sediments. These alterations to the bulk sedimentary $\delta^{15}$N signal may be identified by isolating and measuring specific fractions that are more resistant to alteration such as compounds derived from chlorophyll, or that have been physically protected, such as microfossil bound compounds. Bulk sedimentary $\delta^{15}$N measurements from areas where the organic matter is well-preserved are generally considered to be accurate proxies of water-column processes, so measurements from suboxic to anoxic environments or high sedimentation rate locations can be used for reconstruction of paleoenvironmental conditions with a high degree of confidence. In order to conform to the tenets of Fretwell’s Law (as cited by Kendall and Caldwell, 1998), which cautions against using stable isotope data as the sole source of information, other redox and environmental proxies, such as biomarkers, paleontological evidence, and element enrichment, should be used to provide additional support and context for interpretation of $\delta^{15}$N data (e.g., Lee et al., 2019; Adeboye et al., 2020).

Given the caveats associated with interpretation of $\delta^{15}$N profiles for modern and ancient sedimentary systems that have not undergone significant post-depositional alteration, the potential use of nitrogen isotopes as paleoenvironmental and nitrogen cycle proxies in petroleum systems deserves additional analysis. In this paper, we review the recent research on nitrogen isotopes in petroleum systems, including fractionation due to diageneis and catagenesis, and the influence of kerogen type, and highlight applications of $\delta^{15}$N data as paleoenvironmental proxies in petroleum systems.

**POST-DEPOSITIONAL NITROGEN FRACTIONATION**

The reliability of proxies to be used to decipher paleoenvironmental conditions relies on the ability to identify...
TABLE 1 | List of previously published $\delta^{15}$N values for a representative range of petroleum systems and petroleum-related samples.

| OM type | Geographic location | Formation | Range of $\delta^{15}$N values | Reporting reference(s) |
|---------|---------------------|-----------|-------------------------------|-------------------------|
| Bituminous sediment | Various | Various | +4.2 to +10.7‰ | Steihl and Lehmann (1980) |
| Coal | Antarctica | Beaver Lake | +3.0‰ | Bokhoven and Theeuwen (1986) |
| Coal | Australia | Various | +3.0 to +3.7‰ | Rigby and Batts (1986) |
| Coal | Canada | Various | −3.2 to 1.4‰ | Whiticar (1996) |
| Coal | China | Various | −2.9 to +3.6‰ | Ding et al. (2018) |
| Coal | China | Huaiyuan | +3.7 to +7.7‰ | Xie et al. (2021) and references therein |
| Coal | Various | Various | +3.5 to +6.3‰ | Steihl and Lehmann (1980) |
| Coal | Netherlands | State Mine Maurits | +2‰ | Bokhoven and Theeuwen (1986) |
| Coal | New Zealand | Stockton | +1.3‰ | Rigby and Batts (1986) |
| Coal after pyrolysis | Zwickau | Various | +3.9 to 5.1‰ | Steihl and Lehmann (1980) |
| Crude oil | China | Various | −5.79 to +18.11‰ | Chen et al. (2005) |
| Crude oil | Japan | Various | +2.3 to +5.1‰ | Wada et al. (1975) |
| Crude oil | United States | Various | +1 to +6.7‰ | Hoering and Moore (1958) |
| Neutral, low polarity oil fraction | China | Liaohe Basin | +10.6 to +12.2‰ | Oldenburg et al. (2007) |
| Neutral, high polarity oil fraction | China | Liaohe Basin | +9.0 to +11.3‰ | Oldenburg et al. (2007) |
| Basic oil fraction | China | Liaohe Basin | +4.1 to +5.8‰ | Oldenburg et al. (2007) |
| Kerogen (undifferentiated) | Australia | Various | −2.25 to +5.34‰ | Stiehl and Lehmann (2017) and references therein |
| Kerogen (undifferentiated) | Canada | Sulphur Mountain Formation | +2.92 to +7.42‰ | Stiehl and Lehmann (2017) and references therein |
| Kerogen (undifferentiated) | China | Various | −0.48 to +7.50‰ | Chen et al. (2005) |
| Kerogen (undifferentiated) | Mexico | Bandaraya | +2.4 to +4.2‰ | Peters et al. (1978) |
| Kerogen (undifferentiated) | South Africa | Witwatersrand Supergroup | −4.44 to −0.48‰ | Stiehl and Lehmann (2017) and references therein |
| Kerogen (undifferentiated) | Switzerland | Serpiano Oil Shale | −0.9‰ | Chicarelli et al. (1993) |
| Kerogen (undifferentiated) | Unite States | Various | −1.67 to +9.9‰ | Peters et al. (1978) and Stiehl and Lehmann (2017) and references therein; Williams et al. (1995) |
| Type I Kerogen | Russia | Shungite | +3.4‰ | Boudou et al. (2008) and references therein |
| Type I Kerogen | United States | Mahogany Shale, Green River Formation | +16.8‰ | Schimmelmann and Lis (2010) and references therein |
| Type II Kerogen | United States | New Albany Shale | −1.3 to +1.2‰ | Boudou et al. (2008) and references therein; Schimmelmann and Lis (2010) and references therein |
| Type II-S Kerogen | Jordan | Sononian Ghareb limestone | +9.0‰ | Boudou et al. (2008) and references therein; Schimmelmann and Lis (2010) and references therein |
| Type III Kerogen | Various | Various | +2.2 to +6.0‰ | Peters et al. (2000) and references therein |
| Sapropelic kerogen | Mexico | Laguna Mormon Algal mats | +1.87 to +6.18‰ | Peters et al. (1981) |
| Humic kerogen | United States | Staten Island Peaty Soil | +1.15 to +5.31‰ | Peters et al. (1981) |
| Mudstone | United States | Fordoche Field | +0.7 to +4.5‰ | Williams et al. (1995) |
| Natural gas | China | Yinggehai Basin | −9.0‰ to −1.8‰ | Zhu et al. (2000) |
| Natural gas | Germany | Mid-European Basin | +6.5 to +18.0‰ | Zhu et al. (2000) and references therein |
| Natural gas | Netherlands | Slosscheren | +12 to +18‰ | Bokhoven and Theeuwen (1986) |
| Natural gas | Poland | Zechstein Main Dolomite | −9.2 to +15.9‰ | Kotarba et al. (2020) and references therein |
| Natural gas | Russia | West Siberian Basin | −19.0 to −10.7‰ | Zhu et al. (2000) and references therein |
| Natural gas | United States | Various | −10.5 to +14.4‰ | Hoering and Moore (1958) and Zhu et al. (2000) and references therein |
| Oil shale | Australia | Various | −2.5 to +12.7‰ | Rigby and Batts (1986) |
| Organic extract | Poland | Various | −1.7 to +1.8‰ | Bauersachs et al. (2009) |
| Organic extract | Switzerland | Serpiano oil Shale | −3.97‰ | Chicarelli et al. (1993) |
| Organic matter | Germany | Wealden shale | +10.11 to +15.03‰ | Froidel et al. (2021) |
| Porphyrrins | Switzerland | Serpiano oil Shale | −3.04 to −3.36‰ | Chicarelli et al. (1993) |
| Shale | Oklahoma | Caney | +8.2 to +11.2‰ | Quan et al. (2013) |
| Shale | Oklahoma | Woodford | +1.5 to +9.5‰ | Quan et al. (2013) and Rivera et al. (2015) |
| Shale | Virginia | Chahtanooga | −0.7 to 3.0‰ | Tute et al. (2019) |

For analytical and sample details, please refer to the original articles. Please note that this is not an exhaustive list of all published $\delta^{15}$N values from petroleum-focused nitrogen isotope studies.

and constrain the relationship between the measured proxy and the parameter it represents. In the case of bulk sedimentary $\delta^{15}$N profiles in petroleum systems, this means trying to characterize any deviations from the initial $\delta^{15}$N value set by nitrogen reactions in response to paleoenvironmental conditions during deposition to those measured in the bulk sediment from core samples and outcrops. Since the organic matter has undergone significant transformation from fresh organic matter to
petroleum, it is likely that the nitrogen isotope values have also been altered by these same processes. Unfortunately, identification and quantification of these transformation processes and the nitrogen isotope fractionation factors associated with them are not easy to determine or constrain. The heterogeneity of organic matter, the difficulty in characterizing elemental and compositional changes during diagenesis and catagenesis, and the limitations of laboratory pyrolysis experiments are all issues that complicate diagenesis and catagenesis, and the limitations of laboratory characterization of elemental and compositional changes during those processes. Nitrogen isotope values of terrestrial organic material are often more variant and have a wider range of values than those from marine systems (Kendall, 1998 and references within). The influence of terrestrial input on the sedimentary organic matter can be traced through the C/N ratio, as terrestrial materials generally have a higher ratio than those of marine phytoplankton (Hedges and Mann, 1979). This is reflected in higher N$_{org}$/C$_{org}$ ratios for Type II kerogen compared to Type III kerogen at comparable maturities even after nitrogen is lost preferentially to carbon during diagenesis (Boudou et al., 2008). This difference in source is reflected in the measured $\delta^{15}$N values, with $\delta^{15}$N values for Type II kerogen generally similar to that measured in black shale samples (~2 to +2‰), and the $\delta^{15}$N values for Type III kerogen closely resembling those for coal (~+2 to +6‰) (Figure 2; Boudou et al., 2008). Similarly, the $\delta^{15}$N values x for coals and lacustrine and brackish-water oil shales are generally higher than for oil shales deposited under marine conditions, reflecting both organic matter source and the nitrogen utilization of the precursor organisms (Rigby and Batts, 1986). The $\delta^{15}$N values of coals, in particular, reflect the organic matter source and composition rather than maturity or diagenesis (Figure 2; Rigby and Batts, 1986; Ding et al., 2018; Xie et al., 2021). These distinctions between the marine and terrestrial organic matter sources indicates that if the nitrogen source or kerogen type is unknown, caution must be used when interpreting $\delta^{15}$N records in petroleum systems, as higher $\delta^{15}$N values could be a consequence of suboxic conditions characterized by predominant water column denitrification or reflect the presence of significant amounts of terrestrial input. While this is also true of all sedimentary systems, the transformation of the initial organic material during diagenesis and catagenesis may make identification of terrestrial input through C/N ratios more difficult.

**Diagenesis**

The loss of organic nitrogen during diagenesis and catagenesis is well documented (e.g., Tissot and Welte, 1984; Prokopenko et al., 2006 and references within; Vandenbrouke and Largeau, 2007); however, how these processes alter the $\delta^{15}$N signal in bulk sediment is not conclusive. Figure 3 summarizes four different possible scenarios for the relationship between the N/C ratio of kerogen and the $\delta^{15}$N$_{kerogen}$ in sediments. As stated earlier, analysis of modern-day core top and particulate organic
matter indicates that in general, sedimentary $\delta^{15}$N is an accurate recorder of water column processes, even when degradation has occurred (Peters et al., 1978; Altabet and Francois, 1994; Robinson et al., 2012 and references within). During diagenesis, organic nitrogen is converted to ammonium (NH$_4^+$) which can be trapped in the sediments, particularly in clay minerals. Detailed analysis of the nitrogen isotope values for the ammonium released provides conflicting information regarding fractionation during diagenesis. One scenario argues that since the NH$_4^+$ is generated from the organic nitrogen pool, the isotopic values are thought to be similar to the originating organic matter (Scholten 1991; Williams et al., 1995; Hoefs, 2004). Williams et al. (1995) compared the $\delta^{15}$N values of clay-fixed NH$_4^+$ from mudstones with the $\delta^{15}$N values of the associated kerogen and found no significant difference, leading them to conclude that the mudstones must be acting as a closed system (Figure 4). In this scenario, the $\delta^{15}$N profile of the remaining sedimentary organic matter would be unchanged despite the loss of N as illustrated by arrow 1 in Figure 3, and thus primarily reflect processes that set the $\delta^{15}$N values of the initial organic matter and not fractionation during diagenesis.

An alternative scenario is that nitrogen isotope fractionation does occur during the generation of NH$_4^+$, with the degree and direction of fractionation varying with environmental conditions during diagenesis. Incubation experiments under anoxic conditions show a decrease in the $\delta^{15}$N values of organic matter of approximately 3‰ due to microbial degradation (Lehmann et al., 2002). Similarly, analysis of $\delta^{15}$N-NH$_4^+$ in pore water in the modern-day Santa Barbara Basin shows an enrichment of 1–3‰ compared to the residual organic matter (Prokopenko et al., 2006). Anoxic degradation appears to result in either the loss of a pool of organic nitrogen that is $^{15}$N-enriched (Figure 3, arrow 2), or the addition of a $^{15}$N-depleted fraction to the organic matter pool (Figure 3, arrow 3). Generation of heavy NH$_4^+$ via organic matter degradation could be due to anoxic diagenesis of more labile, isotopically enriched marine organic matter over more refractory, more depleted terrestrial nitrogen (Prokopenko et al., 2006). Incorporation of organic compounds generated by bacteria could be the source of lighter N to the organic matter pool (Lehmann et al., 2002). Unlike the incubation experiments, the loss of heavy nitrogen via diagenesis did not significantly change the $\delta^{15}$N values of the sedimentary organic matter in the Santa Barbara Basin, likely due to sediment conditions favorable for good organic matter preservation (Figure 4; Prokopenko et al., 2006). It is probable that the
$\delta^{15}$N values of organic matter could decrease during diagenesis if sediments had lower organic matter concentrations, slower sedimentation rates, and more oxic conditions, or in settings where the input of lower $\delta^{15}$N compounds is significant. In contrast, oxic degradation increases the $\delta^{15}$N value of organic matter (Figure 3, arrow 4) as $^{15}$N-depleted nitrogen is preferentially lost as NH$_4^+$ during early diagenesis (Macko and Estep, 1984; Holmes et al., 1999; Freudenthal et al., 2001; Lehmann et al., 2002). The bulk sedimentary $\delta^{15}$N signal appears to reflect the $\delta^{15}$N values of organic matter and thus reflect water column processes in organic-rich areas with rapid accumulation rates. In contrast, sediments with low organic N concentrations and high NH$_4^+$ levels are more likely to have altered $\delta^{15}$N profiles (Robinson et al., 2012 and references within). In a petroleum-generating system characterized by high organic matter concentrations and suboxic to anoxic environmental conditions, the use of $\delta^{15}$N values as a redox proxy may be plausible. Plotting total organic carbon (TOC) concentrations vs total nitrogen (TN) concentrations for bulk sediment samples can provide information about excess inorganic N which may indicate whether significant diagenehas occurred and that the $\delta^{15}$N values for the bulk sediment may have been altered (Calvert, 2004).

**Catagenesis**

Even after diagenesis, there is still nitrogen present in kerogen that can be lost during catagenesis, potentially resulting in additional nitrogen isotope fractionation. While modern-day systems can be used to evaluate alteration of $\delta^{15}$N values during diagenesis, no such analogs exist for catagenesis. Changes in the nitrogen content and isotopic values due to catagenesis can only be evaluated through laboratory experiments or inferred through analysis of sediments from petroleum reservoirs. Though these analyses are imperfect due to the limitations of pyrolysis/thermal maturity experiments and the difficulty in isolating catagenic effects from other geological processes in petroleum-generating systems, all studies indicate that nitrogen is lost during catagenesis. Analysis of kerogen isolated from sediments after a series of thermal maturation experiments indicates that changes in nitrogen content with temperature were complex and varied with organic matter type, as algal-rich sediments showed a different pattern compared to more terrestrial peat-rich sediments (Peters et al., 1981). The amount of nitrogen lost during catagenesis also appeared to vary by kerogen maturity and on the method of evaluation. Analysis of nitrogen loss using pyrolysis results in low release of nitrogen from both Type I and Type II kerogen, generally less than 12% of the initial kerogen N concentration (Barth et al., 1996; Gillaizeau et al., 1997; Behar et al., 2000). In comparison, evaluation of C/N ratios and total N, organic N, and fixed-NH$_4^+$ concentrations of samples from both oil-producing and non-productive areas of the Santa Maria and San Joaquin Basins in California indicates that about half the nitrogen remaining after diagenesis is lost (Compton et al., 1992). Some of this difference may be due to a discrepancy in the inorganic N that is measured in the field versus what is measured in pyrolysis studies. The pyrolysis experiments found that the majority of the N released from kerogen was as N$_2$ gas, with little as NH$_3$ (Gillaizeau et al., 1997; Behar et al., 2000). In contrast, the field analyses generally measured NH$_4^+$ in the surrounding matrix, which could possibly have been brought in from outside the system, though C/N ratios document significant N loss (Compton et al., 1992). Field analyses in natural gas systems have measured N$_2$ levels, but it is unclear how much of the original kerogen-bound nitrogen was converted to gas (Zhu et al., 2000). Studies generally agree, however, that there is some amount of nitrogen in kerogen that is resistant to catagenic degradation, and that the majority of the nitrogen loss that does occur takes place at higher thermal maturities, with maximum NH$_4^+$ evolution in the peak oil generation stage and maximum N$_2$ release during gas generation (Compton, et al., 1992; Krooss et al., 1995; Littke et al., 1995; Barth et al., 1996; Gillaizeau et al., 1997; Behar et al., 2000). The degree and direction of nitrogen isotope fractionation that results from nitrogen loss during catagenesis is also highly variable (Figure 5). Some studies have found that increasing thermal maturity generally results in heavier $\delta^{15}$N values for the remaining nitrogen in kerogen, since the light N is preferentially removed (Figure 5; Stiehl and Lehmann, 1980; Peters et al., 1981). In contrast, other evidence suggests that the $\delta^{15}$N values for Type III kerogen and coals do not seem to change significantly with higher maturity levels even as N$_{org}$ is lost, though the data for Type I and Type II kerogen is limited and inconclusive (Figure 5; Boudou et al., 1984; Rigby and Batts, 1986; Whiticar, 1996; Ader et al., 1998; Ader et al., 2006; Boudou et al., 2008; Xie et al., 2021). In other experiments, $\delta^{15}$Nkerogen
values decrease with increasing metamorphic grade, while the silicate-bound nitrogen becomes correspondingly heavier (Figure 5; Stüeken et al., 2017 and references within).

**Fluid Migration**

An additional difficulty in interpreting bulk δ15N values in samples from petroleum systems is that these systems may experience nitrogen fractionation as a result of fluid interactions during migration, in addition to changes driven by diagenesis and catagenesis. Interaction with various fluids (e.g., during petroleum expulsion from source intervals, interaction with other non-petroleum fluids in reservoir intervals) can result in nitrogen exchange between kerogen and nitrogen compounds in fluids, which can alter the δ15N values of kerogen depending on the source and type of N compounds present in the fluids (Schimmelmann and Lis, 2010). Exchange within a basin may have the effect of homogenizing the δ15N values for different N pools and in different areas across the basin, eliminating any variations in δ15N values that may be diagnostic of prior geochemical processes or original paleoenvironmental conditions (Schimmelmann and Lis, 2010). Isotopic analysis of NH4+ or N2 generated by diagenesis and catagenesis may also be impacted by infiltration of those compounds from other sources or subsequent reaction in the fluid phase (e.g., Williams et al., 1995; Zhu et al., 2000; Xie et al., 2021). In particular, loss of light nitrogen either as NH3 (from the volatilisation of NH4+) or N2 gas would result in enrichment for the remaining nitrogen in fluids (Williams et al., 1995; Ador et al., 1998; Krooss et al., 2005). Trapping of this heavy fluid nitrogen by clays could lead to higher δ15N values for the bulk sediment in areas with significant concentrations of clay minerals (Charlesworth, 1986; Williams et al., 1995; Xie et al., 2021). The bulk sedimentary δ15N values of petroleum reservoir sediments may also be influenced by the enrichment of the δ15N values of organic nitrogen in petroleum due to migration (Williams et al., 1995 and references within).

**Impact on Interpretation of δ15N Data**

Based on the previous paragraphs, it is clear that there is no consensus regarding the impact of kerogen type, diagenesis, catagenesis, fluid interaction, and migration on bulk sedimentary δ15N values from petroleum systems. The heterogeneity of the organic matter, environmental conditions, and geological processes in petroleum systems make it complicated to interpret whether observed trends in bulk δ15N values reflect initial nitrogen reactions and paleoredox conditions or if these trends indicate alteration (Table 1). If some broad conclusions can be drawn, the reliability of bulk sedimentary δ15N records as redox proxies are higher if conditions are more conducive for organic matter preservation: high organic matter productivity, predominantly anoxic conditions, lower maturity, and minimal fluid interaction and migration.

Given all of the possible scenarios, it seems unlikely that the numerical value of bulk sedimentary δ15N from petroleum systems is the same as the initial numerical value set during sediment deposition; however, it is possible that the relative relationships of δ15N values within a sedimentary unit or location could be unchanged if the unit or location were subjected to similar geological processes over the interval in question. For example, this means that the presence of anoxic conditions could not be diagnosed based on a δ15N value near 0‰, but an interval of lower δ15N values within a profile of higher δ15N values could be interpreted as an anoxic interval, particularly if supported by other geochemical and stratigraphic proxies. This means that it would be difficult to make direct comparisons of δ15N values from sequences or locations that are too distant in either time or place, since they would likely be altered by different geological processes and therefore experience different influences with regards to isotopic fractionation. While there does seem to be some variation in sedimentary δ15N values from unaltered systems through time on a global scale (Algeo et al., 2014), the potential fractionation due to catagenesis and migration likely make it inadvisable to draw global-scale conclusions from δ15N measurements in petroleum systems.

**NITROGEN ISOTOPE MEASUREMENT**

Another potential complication in the comparison between δ15N values for different locations and time periods is that the δ15N values may also vary depending on the sediment fraction that is measured. The development of simpler and more sensitive isotope ratio mass spectrometer (IRMS) instrumentation in the last few decades means that the δ15N of various nitrogen fractions can be measured relatively easily and using smaller sample sizes than in the past. The most basic measurement is bulk sedimentary δ15N, as the sediment samples simply need to be powdered, dried, and wrapped in tin capsules prior to combustion and analysis in an elemental analyzer (EA)-IRMS system, an improvement over the off-line Kjeldahl digestion or thermal oxidation methods previously used. This bulk sediment measurement can be referred to as δ15Nbulk, δ15Nsed, or δ15NTN, and represents the δ15N of the total nitrogen in the sediments, a sum of both the organic material and any inorganic N trapped in mineral matrices. This has become the most common measurement used for the δ15N redox proxy and is generally thought to accurately represent the original δ15N of organic matter in the water column as any NH4+ generated from the remineralization of organic matter is retained in the surrounding mineral matrix and included in the bulk measurement (Altabet and Francois, 1994; Higgins et al., 2012; Robinson et al., 2012).

In some samples, low organic matter concentrations or high amounts of detrital material may mean that nitrogen concentrations in the bulk sediments may be too low or too diluted to meet the detection limit of the IRMS. If the sediment is carbonate-rich, acidification methods can be used to reduce the sediment mass and concentrate the nitrogen. Acidification can either be performed in centrifuge tubes, with the residual sediments rinsed with water to remove acid then dried before analysis (rinse method), or on a smaller scale directly in silver capsules with the acid removed by evaporation (capsule method) (Brodie et al., 2011). The measured values for δ15Nacidified are generally not the same as for δ15Nbulk and the rinse method and the capsule method often return different δ15N values. This
FIGURE 6 | (A): Ranges of measured $\delta^{15}$N values for shale samples from unconventional reservoirs as referred to in the text. Values shown are from Rigby and Batts 1986, Quan et al. 2013, Rivera et al. 2015, and Tutte et al. 2019; for analytical and sample details, please refer to the original articles. (B): Ranges of measured $\delta^{15}$N values for lacustrine nitrogen fractions as referred to in the text. Values shown are from Oldenburg et al. 2007; for analytical and sample details, please refer to the original article. (C): Ranges of measured $\delta^{15}$N values for porphyrin, kerogen, and organic extracts as referred to in the text. Values shown are from Chicarelli et al. 1993; for analytical and sample details, please refer to the original article.
indicates that some nitrogen is lost during the acidification process, possibly through loss of volatile nitrogen or water-soluble nitrogen, though the exact identity of the lost nitrogen fraction is still undetermined (Brodie et al., 2011). Therefore, the $\delta^{15}N_{\text{bulk}}$ and $\delta^{15}N_{\text{acidified}}$ fractions are not equivalent, and numerical comparison of these values should be done with caution.

Nitrogen isotope measurements of the isolated kerogen fraction can capture the organic N isotope signal without the interference of inorganic nitrogen compounds. This $\delta^{15}N_{\text{kerogen}}$ can be measured using solvent-extracted rock powder or on the kerogen obtained after removal of carbonate minerals by acidification and silicate minerals by HF (e.g., Higgins et al., 2012; Stüeken et al., 2017). Other nitrogen fractions can also be measured for $\delta^{15}N$, including silicate-bound, foram-bound, inorganic, porphyrins, polar N compounds, and neutral N compounds, if they can be isolated from the bulk sediment. As each of these fractions measures specific nitrogen fractions, comparison between the various $\delta^{15}N$ values may provide additional information regarding the partitioning of nitrogen and nitrogen isotopes between these pools (e.g., Oldenburg et al., 2007; Stüeken et al., 2017).

**INTERPRETATION OF $\delta^{15}N$ VALUES IN PETROLEUM SYSTEMS**

The use of sedimentary nitrogen isotopes in the evaluation of petroleum reservoirs has increased in recent years as the use of $\delta^{15}N_{\text{bulk}}$ as a redox proxy has become more established. While there is still much more work to be done on the theory and systematics of nitrogen isotope partitioning and fractionation in such systems, $\delta^{15}N$ values of various fractions can still be used to interpret depositional paleoenvironmental conditions, nitrogen cycling, and organic matter sources. Here we present some examples to illustrate how sedimentary $\delta^{15}N$ has been used in a range of petroleum systems to evaluate paleoredox conditions, lacustrine settings, and organic matter fractions.

**Unconventional Reservoirs**

Unconventional reservoirs are petroleum systems characterized by low porosity and permeability, which often results in a lack of a clear density separation for the hydrocarbons and formation fluids contained within the reservoir unit (J. Puckette, personal communication). A significant amount of research has been done over the last couple of decades into the systematics, depositional conditions, and reservoir characteristics of these organic-rich shale, mudstone, and carbonate deposits. Due to their low permeability and porosity, the $\delta^{15}N_{\text{bulk}}$ values from unconventional reservoirs are likely to retain the isotopic fingerprint of water column nitrogen sources and processes during deposition, as these systems would experience minimal migration of either the organic or inorganic nitrogen pools. Rigby and Batts (1986) measured the $\delta^{15}N$ of oil shale samples from Australia and noted that the $\delta^{15}N_{\text{bulk}}$ values reflected the type of organic material, nitrogen source, and paleoenvironmental conditions during deposition, similar to unaltered sediments (Figure 6A). To evaluate whether $\delta^{15}N$ profiles could be used as a paleoproxy for water column redox state, the $\delta^{15}N_{\text{bulk}}$ values for two productive units from Oklahoma with similar geological histories but different depositional redox conditions were compared. The $\delta^{15}N_{\text{bulk}}$ values from the Woodford Shale, deposited under anoxic conditions, were found to be lower than those from the overlying suboxic Caney Shale, supporting the interpretation that $\delta^{15}N_{\text{bulk}}$ profiles can be used as a paleoredox proxy even for oil-mature unconventional formations (Figure 6A; Quan et al., 2013). Measurement of the $\delta^{15}N_{\text{bulk}}$ values from Woodford Shale samples taken from the Anadarko Basin (Oklahoma) found that there were no trends in $\delta^{15}N$ values with thermal maturity, as inferred from vitrinite reflectance measurements (Figure 6A; Rivera et al., 2015). Correlation of the $\delta^{15}N_{\text{bulk}}$ values with measured Fe and Mo concentrations once again indicated that the primary control on the $\delta^{15}N_{\text{bulk}}$ values appeared be depositional redox conditions (Rivera et al., 2015). Nitrogen isotope analysis of acidified samples from the oil-producing Huron, Three Lick, and Cleveland Shale formations identified variations that corresponded to two distinct depositional environments (Figure 6A; Tuite et al., 2019). Since paleoredox conditions also affect the predominant nitrogen reactions, the identification of these different redox regimes provided information about changing productivity, nutrient cycling, and water column stratification during alternating icehouse/greenhouse conditions (Tuite et al., 2019).

The results from these unconventional petroleum reservoir studies indicate that the sedimentary $\delta^{15}N$ values accurately reflect organic matter sources and redox conditions during deposition, which can be used to characterize the unconventional reservoir and evaluate its formation, evolution, and potential hydrocarbon yield. In addition, the $\delta^{15}N$ profiles from unconventional reservoir formations can also provide valuable information regarding past biogeochemical cycles, environmental conditions, and climate regimes even though these sediments have undergone significant catagenic alteration (Quan et al., 2013; Rivera et al., 2015; Tuite et al., 2019). This potentially expands the number of locations and geological formations that can be used to address paleoenvironmental and climatic questions, provided that the cores have robust age models and have not been influenced by loss of either organic or inorganic nitrogen phases.

**Lacustrine Basins**

Nitrogen reactions and their isotopic signatures are more complex in lacustrine and other freshwater systems due to the influence of terrestrial nutrients and organic matter, the smaller size of the water bodies, and the impacts of seasonal cycles (Talbot 2001 and references within; Quan and Falkowski, 2009 and references within). This means that the use of sedimentary $\delta^{15}N$ profiles to interpret paleoenvironmental conditions in lakes is not as well constrained as in marine environments, and evaluation of thermally mature lacustrine formations even less so. As mentioned above, the $\delta^{15}N$ values and C/N ratios for Type III kerogen and coals do reflect the influence of terrestrial organic material and are distinct from the Type I and Type II kerogen values (Rigby and Batts, 1986; Boudou et al., 2008; Ding et al., 2018; Xie et al., 2021). Analysis of two different lacustrine...
source rocks from the Junggar Basin of China found that crude oils from brackish lacustrine basins were more enriched in pyrrolic-N containing compounds derived from biological sources than crude oils from alkaline lacustrine settings (Zhang et al., 2020). This observed enrichment was attributed to higher terrestrial plant contributions, particularly alkaloid and chlorophyll compounds, to the original organic matter deposits of the brackish lacustrine basin. While changes in N-compounds found in crude oils may also be a consequence of fractionation brought about by migration and thermal maturity, these were not considered to be the main processes responsible for differences in abundance and type of N-containing compounds seen in the end member oils derived from brackish vs. alkaline lacustrine settings (Zhang et al., 2020). Analysis of different organic nitrogen fractions extracted from a single crude oil sample from the Laohe Basin (China) indicated that the polar neutral nitrogen fraction (including pyrrolic-N compounds) was generally enriched in $^{15}\text{N}$ compared to basic nitrogen compounds by approximately 5‰ (Figure 6B; Oldenburg et al., 2007). While the $\delta^{15}\text{N}$ values of the crude oils from the Junggar Basin study were not measured, given the differences in organic matter sources and the enrichment of specific pyrrolic-N compounds, it would not be unexpected for the $\delta^{15}\text{N}_{\text{oil}}$ values to be significantly different in the brackish basins than the alkaline settings. Oldenburg et al. (2007) attributed the higher $\delta^{15}\text{N}$ values for pyrrolic-N compounds to be a result of increasing thermal maturity; however, Zhang et al. (2020) did not find higher concentrations of pyrrolic-N compounds in more mature samples. Resolution of this discrepancy by future studies may provide important information on the relative influence of organic matter source versus thermal maturity in lacustrine and freshwater systems. Given the complexity of processes that could potentially have a significant influence on the $\delta^{15}\text{N}$ signals in lacustrine systems, it is likely that the processes that impact lacustrine systems will vary depending on geologic setting and history of the particular paleolakes being studied.

**Porphyrians**

Porphyrians have traditionally been used as biomarker proxies for thermal maturity and to illustrate different depositional environmental conditions (Mawson et al., 2004). Since porphyrian compounds are direct breakdown products of chloropigments, the $\delta^{15}\text{N}$ values of porphyrians are considered to be a direct representation of the isotopic composition of those original chloropigments and thus of the original primary producers (e.g., Hayes et al., 1987; Boreham et al., 1989; Boreham et al., 1990; Chicarelli et al., 1993; Kashiyama et al., 2008). Porphyrians have been isolated in petroleum systems in samples with a wide range of maturities and retain the $\delta^{15}\text{N}$ signal of the original organic matter, so they could be a way to evaluate water column N reactions during deposition that would not be altered by post-depositional isotopic fractionation processes. Comparison of two different porphyrian fractions, kerogen, and total organic extract isolated from Serpiano oil shale sediments show that porphyrians fractions (average $\sim-3.1$% and $\sim-3.3$‰) are depleted in $^{15}\text{N}$ relative to the kerogen (average $\sim-0.9$%) and slightly enriched relative to the total extract (average $\sim-4.0$%) (Figure 6C; Chicarelli et al., 1993). While the depletion in the porphyrian fractions relative to the kerogen may indicate the influence of an additional nitrogen source, the low $\delta^{15}\text{N}$ values for all three fractions reflects the influence of predominant N$_2$ fixation to the original organic matter pool (Chicarelli et al., 1993). This indicates that $\delta^{15}\text{N}_{\text{porphyrin}}$ values can serve to determine whether isotopic fractionation has altered $\delta^{15}\text{N}_{\text{kerogen}}$ and/or $\delta^{15}\text{N}_{\text{bulk}}$ values for petroleum systems during catagenesis or migration. While the isolation and measurement of $\delta^{15}\text{N}$ for porphyrians is not trivial, it may be necessary if post-depositional fractionation processes are suspected and alteration of the $\delta^{15}\text{N}_{\text{bulk}}$ signal may have occurred.

**CONCLUSION AND FUTURE WORK**

In summary, nitrogen isotopes can be a powerful tool in petroleum systems, particularly for evaluating unconventional reservoirs, palaeoenvironmental conditions, and organic matter sources. Uncertainties regarding how diagenesis, catagenesis, hydrothermal fluids, and hydrocarbon migration processes may have fractionated the initial organic nitrogen deposits can result in significant caveats in terms of interpretation of the $\delta^{15}\text{N}$ values, and these may limit the use of $\delta^{15}\text{N}$ profiles as a redox and nitrogen cycle proxy for more complex petroleum systems. The more that is known about non-nitrogen parameters in the petroleum system being analyzed, such as fluid interactions, mineralogy, kerogen type, and porosity and permeability, the more constrained the interpretation of the $\delta^{15}\text{N}$ measurements can be, and the greater the potential to conclusively identify and characterize palaeoenvironmental regimes and biogeochemical cycles. In addition to additional research to better constrain post-depositional nitrogen processes and fractionation, more $\delta^{15}\text{N}$ data from more locations, wider thermal maturity ranges, and different depositional environments is necessary to link measured $\delta^{15}\text{N}$ values to specific palaeoenvironmental conditions and processes. While caution is advised against placing too much emphasis on specific $\delta^{15}\text{N}$ values in terms of interpretation, overall trends and excursions present in $\delta^{15}\text{N}$ profiles from petroleum systems can still provide critical information about reservoir characteristics, particularly depositional paleoredox conditions and organic matter sources.

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All authors listed have made a substantial, direct, and intellectual contribution to the work and approved it for publication.

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Nitrogen Isotopes in Petroleum Systems

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