Carbon and sulfur isotopic signatures of ancient life and environment at the microbial scale: Neoarchean shales and carbonates

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ABSTRACT

An approach to coordinated, spatially resolved, in situ carbon isotope analysis of organic matter and carbonate minerals, and sulfur three- and four-isotope analysis of pyrite with an unprecedented combination of spatial resolution, precision, and accuracy is described. Organic matter and pyrite from eleven rock samples of Neoarchean drill core express nearly the entire range of δ13C, δ34S, Δ33S, and Δ36S known from the geologic record, commonly in correlation with morphology, mineralogy, and elemental composition. A new analytical approach (including a set of organic calibration standards) to account for a strong correlation between H/C and instrumental bias in SIMS δ13C measurement of organic matter is identified. Small (2–3 μm) organic domains in carbonate matrices are analyzed with sub-permil accuracy and precision. Separate 20- to 50-μm domains of kerogen in a single ~0.5 cm3 sample of the ~2.7 Ga Tumbiana Formation have δ13C = −52.3 ± 0.1‰ and −34.4 ± 0.1‰, likely preserving distinct signatures of methanotrophy and photoautotrophy. Pyrobitumen in the ~2.6 Ga Jeerinah Formation and the ~2.5 Ga Mount McRae Shale is systematically 13C-enriched relative to co-occurring kerogen, and associations with uraniferous mineral grains suggest radiolytic alteration. A large range in sulfur isotopic compositions (including higher Δ33S and more extreme spatial gradients in Δ35S and Δ36S than any previously reported) are observed in correlation with morphology and associated mineralogy. Changing systematics of δ34S, Δ33S, and Δ36S, previously investigated at the millimeter to centimeter scale using bulk analysis, are shown to occur at the micrometer scale of individual pyrite grains. These results support the emerging view that the dampened signature of mass-independent sulfur isotope fractionation (S-MIF) associated with the Mesoarchean continued into the early Neoarchean, and that the connections between methane and sulfur metabolism affected the production and preservation of S-MIF during the first half of the planet’s history.

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INTRODUCTION

The search for evidence of life and habitable environments in ancient sedimentary rocks is facilitated by the fact that elements essential to all known organisms (e.g., C, H, O, N, S) have multiple stable isotopes whose relative abundances change in characteristic ways when processed biologically. Isotopic fractionation results from both equilibrium and kinetic processes and can generate measurable differences in the compositions of metabolic substrates and products. Preservation of these metabolic substrates and products over million- and billion-year time scales, however, commonly includes their further transformation into relatively recalcitrant, complex mixtures of organic matter (kerogen and bitumen) or minerals (e.g., carbonates, quartz, oxides, sulfates, sulfides, etc.) through secondary (early diagenetic) processes that can impart additional, abiotic fractionations. After these secondary transformations, isotopic compositions of sedimentary organic matter and biominerals may be further altered by tertiary (late diagenetic) processes including metamorphism and metasomatism. Detection of stable isotopic signatures of life and environment, then, requires deconvolution of these post-metabolic, secondary, and tertiary alteration processes. The likelihood of biosignature alteration or destruction increases with increasing age, and it follows that any records of life and environment preserved in the most ancient rocks on Earth (and beyond) are likely to be sparse and ambiguous.

At the regional, outcrop, or drill-core scale, the extent to which biologically driven isotopic fractionation processes may be recorded in sediments can be assessed by geologic mapping, visual inspection of hand sample lithology, detailed petrographic analysis of selected samples, and bulk geochemistry. So-called ‘bulk’ or ‘whole rock’ stable isotope analysis provides unique insights into biosignature formation and alteration processes by revealing spatial heterogeneity and compositional patterns masked by bulk techniques. In rocks older than ~2.4 Ga, deviations in the relative abundances of the rare $^{33}$S and $^{36}$S relative to the more common isotopes of sulfur ($^{32}$S and $^{34}$S) from those expected due to purely mass-dependent processes ($\Delta^{33}$S and $\Delta^{36}$S) are thought to have been produced during UV disassociation of SO$_2$ and possible only in an anoxic atmosphere (Farquhar et al., 2000). The preservation of such ‘mass-independent’ or ‘non-mass-dependent’ fractionation of sulfur isotopes (S-MIF) further requires that sulfur left the atmosphere as multiple, discreet molecular species (H$_2$SO$_4$ and S$_8$), and escaped homogenization due to a low concentration of sulfate in the oceans (Pavlov & Kasting, 2002; Ono et al., 2003; Ono, 2008). Measurement of S-MIF has thus been used to constrain the timing of atmospheric oxygenation (Farquhar & Wing, 2003; Bekker et al., 2004; Papineau et al., 2007; Guo et al., 2009; Williford et al., 2011). Recent empirical and theoretical analysis of small deviations in biological fractionation effects between the individual sulfur isotopes (Johnston et al., 2005; Johnston, 2011), as well as correlations between anomalously low bulk $\delta^{13}$C$_{org}$ values and small deviations in the ratio of $\Delta^{33}$S to $\Delta^{36}$S typical for Archean rocks (Thomazo et al., 2009a,b, 2013; Zerkle et al., 2012), suggests that information about atmosphere–biosphere interactions is uniquely accessible by coordinated carbon and sulfur four-isotope analysis (Ono et al., 2006).

In this study, we build upon insights into the behavior of the Neoarchean (2.8–2.5 Ga) biosphere and atmosphere gained from coordinated bulk stable carbon and sulfur four-isotope analysis by developing and applying the techniques to detect these signals in situ, at the scale of the micro-organisms and microenvironments within which they were originally produced and preserved. Here, we report new methods for, and data from, carbon isotope analysis of kerogen, pyrobitumen, calcite, and dolomite, and sulfur three- and four-isotope analysis by developing and applying the techniques to detect these signals in situ, at the scale of the micro-organisms and microenvironments within which they were originally produced and preserved. Here, we report new methods for, and data from, carbon isotope analysis of kerogen, pyrobitumen, calcite, and dolomite, and sulfur three- and four-isotope analysis of pyrite using secondary-ion mass spectrometry (SIMS). These measurements are made in petrographic context provided by light and scanning-electron microscopy (SEM), as well as elemental analysis by energy-dispersive x-ray spectroscopy (EDS). Although this approach is resource intensive compared to traditional bulk measurements, and is therefore applied to a relatively small number of rock samples, the information density per sample is extraordinarily high and inaccessible by other means. The approach we outline here for spatially resolved, in situ stable carbon and sulfur isotope analysis provides unique insights into biosignature formation and alteration processes by revealing spatial heterogeneity and compositional patterns masked by bulk techniques.

\footnote{We report isotopic compositions in standard ‘delta notation’, for example $\delta^{13}$C$_{org} = \left( \frac{^{13}C}{^{12}C} \right)_{org} \left( \frac{^{12}C}{^{12}C} \right)_{standard} - 1 \times 1000.$}

\footnote{$\Delta^{33}$S = $\delta^{33}$S – 1000 * $\left( \frac{1 + \delta^{32}$S_{000}}{1 + \delta^{36}$S_{000}} \right)^{0.515} - 1$;
$\Delta^{36}$S = $\delta^{36}$S – 1000 * $\left( \frac{1 + \delta^{32}$S_{000}}{1 + \delta^{36}$S_{000}} \right)^{1.9} - 1$}
**Archean organic geochemistry**

Lipid biomarkers, including polycyclic terpenoids such as hopanes and steranes (the diagenetic products of bacterial and eukaryotic membrane lipid components, respectively), extracted from shales and carbonates of Neoarchean strata in the Pilbara Craton of Western Australia (WRL-1, RHDH2, and SV-1 cores), and sediments from the Transvaal Supergroup of the Kaapvaal Craton of South Africa (GKF and GKP cores) were proposed to record some of the oldest molecular evidence of life on Earth, including the oldest evidence for eukaryotes, and to indicate oxygenic photosynthesis hundreds of millions of years prior to initial atmospheric oxygenation (Brocks et al., 1999, 2003a,b,c; Eigenbrode, 2004; Eigenbrode et al., 2008; Waldbauer et al., 2009).

These reports described the risks of false positives in the search for syngenic Archean hydrocarbons, and the authors took great care to evaluate the possibilities for contamination that would have confounded their interpretations. Recently, as part of a wide-ranging collaborative effort to test whether the Archean biomarkers are indigenous and syngeneric using unprecedented contamination control measures during drilling and later sample handling, new drill cores intersecting sediments of the Fortescue and Hamersley Groups in the Pilbara have been obtained by the Agouron Institute Drilling Project. Analyses of the interior portions of these new core samples yielded hopane and sterane biomarker concentrations comparable to procedural blanks and negative controls, whereas analyses of exterior portions (and re-analyses of the formerly studied and correlative samples) yielded concentrations an order of magnitude higher, consistent with contamination (French et al., 2015). This work demonstrates that alternative analytical approaches are needed to understand the biogeochemical records preserved in Archean organic matter.

**SIMS analysis of Archean organic matter**

The syngenericity of Neoarchean biomarkers was called into question earlier by a study that used spatially resolved, in situ $\delta^{13}C$ measurements of kerogen and pyrobitumen in samples of the Roy Hill Member of the Jeerinah Formation taken from the WRL-1, DDH 186, and FVG-1 drill cores. The authors argued on the basis of a 10–20$^{\circ}$ error in $\delta^{13}C$ between the extracted hydrocarbons and less mobile, and relatively $^{13}C$-depleted organic phases such as kerogen and pyrobitumen that the hydrocarbons must postdate peak metamorphism at ~2.2 Gyr ago (Rasmussen et al., 2008). The measurements by Rasmussen et al. (2008) were made using a CAMECA NanoSIMS 50 (commonly called ‘NanoSIMS’ and distinct from the instrument used to collect data in this study) that rastered a beam of Cs$^+$ ions across a 25-$\mu$m$^2$ area on the surface of a sample. NanoSIMS excels at 50- to 100-nm scale elemental mapping (e.g. Oehler et al., 2009).

However, a number of factors such as quasi-simultaneous arrival (Slodzian et al., 2004), electron multiplier aging, and extreme sensitivity to surface relief due to the short working distance (Fletcher et al., 2008) are impediments to achieving the sub-permil accuracy and precision for natural abundance light-stable isotope analysis at micrometer-scale spatial resolution for which the large-radius multicollector CAMECA IMS 1280 SIMS (commonly called ‘1280’, or simply, ‘SIMS’) instrument was designed (Valley & Kita, 2009). For example, a relation between instrumental bias and H/C was observed during $\delta^{13}C$ measurements of organic matter using a large-radius IMS 1270 (Sangdaly et al., 2005, 2007), and the extent to which this affects NanoSIMS measurements is unknown or has yet to be reported. We refer the reader to Orphan & House (2009) for further detailed discussion of the relative merits of various SIMS techniques for geobiological investigations.

‘Orientation effects’, or measurable relations between instrumental bias and crystal orientation, can also adversely affect accuracy of SIMS measurements. In such cases, relatively high ‘spot-to-spot’ precision as measured on an individual standard grain in a single orientation may belie significant ‘grain-to-grain’ differences in bias that arise from different crystal orientations and thus degrade true accuracy. For example, orientation effects in sphalerite and galena lead to degraded precision and accuracy in $\delta^{34}S$ analyses of these minerals relative to analyses of pyrite made with the same instrumental settings, but for which there is no known orientation effect (Kozdon et al., 2010). An orientation effect of several per mil has been observed in $\delta^{13}C$ measurements of individual graphite crystals in the WiseSIMS laboratory (T. Ushikubo, S. R. Dunn & J. W. Valley, unpubl. data). To our knowledge, the impact of orientation effects on NanoSIMS measurements of natural abundance stable isotope ratios remains unstudied. Here, we report our efforts using a large-radius IMS 1280 to develop a multiproxy approach for the detection of signs of life and planetary evolution in ancient geologic samples, an approach that in our view provides an optimal combination of spatial resolution, precision, and accuracy.

**Geologic setting**

The samples used in this study come from the Mount Bruce Supergroup, a 2.77–2.43 Ga package of sedimentary and volcanic rocks deposited in the Hamersley Basin on an Archean granite greenstone terrain within the Pilbara Craton of Western Australia (Trendall & Blockley, 1970; Blake et al., 2004). Figs S1 and S2 in the Supplementary Information (SI) show a geologic map and composite stratigraphy, respectively.

These rocks are remarkably well preserved for their age, having been subjected only to prehnite-pumpellyte facies burial metamorphism at the basin scale and hydrothermal
alteration locally (Smith et al., 1982; White et al., 2014b). Samples were selected to document in greater detail the range of organic and isotope geochemical heterogeneity observed in several earlier studies of Neoarchean drill core from the Pilbara Craton, Western Australia (Ono et al., 2003; Eigenbrode & Freeman, 2006; Kaufman et al., 2007; Rasmussen et al., 2008; Czaja et al., 2010). Eleven samples ranging in age from ~2.5 to ~2.7 Ga were subsampled from the ABDP-9, WRL-1, RHDH2a, and SV-1 drill cores: Samples from the WRL-1, SV-1, and ABDP-9 drill cores are proximal to the depocenter of the Hamersley Basin, whereas samples from the RHDH2a core, drilled in the Ripon Hills area, likely represent the basin margin (Yamaguchi, 2002; Yamaguchi et al., 2005; Eigenbrode & Freeman, 2006; Czaja et al., 2010). Stratigraphic correlation and depositional age models are facilitated by impact spherule layers observed ~2.5 m below the Jeerinah/Marra Mamba contact near the depocenter in the main outcrop area (Hassler et al., 2005) and ~60 m above the Jeerinah/Carawine contact near the margin in the Oakover River area (Rasmussen et al., 2005). An important implication of this correlation is that the Carawine Dolomite in the Oakover River area represents carbonate deposition in a shelf setting that is time-equivalent to deeper water iron formation deposition in the Marra Mamba Fm located to the southwest (Eigenbrode, 2004; Eigenbrode & Freeman, 2006). On this basis, rocks from the RHDH2a and WRL-1 cores are considered to have been deposited along an increasing depth gradient across the basin. In that light, near-zero bulk $\delta^{56}$Fe values near the center of the basin were previously interpreted to represent a dominance of hydrothermal and clastic Fe, whereas correlated low $\delta^{56}$Fe and total Fe values on the margin suggest active dissimilatory iron reduction (Yamaguchi et al., 2005; Czaja et al., 2010).

SAMPLES AND METHODS

The following represents a summary of methods developed and applied during this study. The reader is encouraged to consult the detailed reporting of our methodology available in the supplementary material.

Descriptions of samples and standards

Samples

Sample selection for this study was guided by previously reported stratigraphic and bulk geochemistry data from the SV-1, RHDH2a, and WRL-1 cores (Eigenbrode & Freeman, 2006; Czaja et al., 2010) and from the ABDP-9 core (Kaufman et al., 2007), and we refer the reader to those publications for further contextual information about the specific sample depths discussed below. Locations of drill cores and their general stratigraphic contexts are shown in Figs S1 and S2.

Samples chosen for this study from the SV-1 core (55.3 and 126.4 m) come from the ~2.7 Ga Tumbiana Fm, a unit in the lower part of the Fortescue Group represented in deeper parts of the basin by shales (as is the case with these samples), and in shallower parts by (commonly stromatolitic) carbonates. One sample from the lower portion of the RHDH2a core (285.3 m) comes from the pyritic black shale of the Roy Hill Member of the ~2.6 Ga Jeerinah Fm, deposited in relatively deep water and located near the top of the Fortescue Group. A single sample from ~30 m higher in the RHDH2a core (252.2 m) represents shallower (shelf facies) deposition in the Carawine Dolomite (Eigenbrode & Freeman, 2006). Bulk inorganic carbon isotope values reported from this unit fall around $0 \pm 0.8\%$ (Shields & Veizer, 2002) and are interpreted to represent original Archean carbonate. Two samples of Wittenoom Formation (289.45 and 520.6 m), one sample from the Marra Mamba Formation (659.55 m) and one sample of the Jeerinah Fm (679.9 m), were taken from the WRL-1 core. Three samples of pyritic black shale in the Mount McRae Shale were taken from the ABDP-9 core: above (111.22 m), within (148.37 m), and below (161.02 m) the interval within which multiple geochemical proxies (including $\Delta^{34}$S) were interpreted to record a transient episode, or ‘whiff’ of surface oxygenation (Anbar et al., 2007; Kaufman et al., 2007). Samples from the WRL-1, RHDH2a, and SV-1 cores were chosen to represent the wide range of published, bulk $\delta^{13}$C values interpreted as evidence for Neoarchean aerobiosis (Eigenbrode, 2004; Eigenbrode & Freeman, 2006). Figure S1 shows the locations of drill cores sampled for this study, and Fig. S2 shows the relative position of the sampled units within composite Neoarchean stratigraphy of Western Australia. Reflected light and backscattered electron mosaic images of each sample with locations of individual SIMS targets are shown in S1 (Figs S10–S18).

Existing SIMS standards

To optimize SIMS precision and accuracy, analyses of samples of unknown isotopic composition are bracketed by analyses of standard materials that, ideally, are mounted together with samples and meet the following criteria: (i) bulk isotopic composition is calibrated by ‘conventional’ analysis (e.g., by gas-source isotope ratio mass spectrometry), (ii) ‘spot-to-spot’ and ‘grain-to-grain’ isotopic composition is homogenous at the micrometer scale based on repeated SIMS analysis, and (iii) chemical composition and crystal structure (if applicable) is as similar as possible to the samples of interest (Valley & Kita, 2009). For sulfur isotope analyses in pyrite and carbon isotope analyses in calcite and dolomite, these criteria had been demonstrated in previous studies for working standards UWPy-1, Balmat pyrite with $\delta^{34}$S = 16.04 ± 0.18\% (all uncertainties are reported as ±2SD unless otherwise indicated),
\[ \Delta^{32}S = -0.003 \pm 0.009_{\text{ppm}} \text{ and } \Delta^{36}S = -0.21 \pm 0.24_{\text{ppm}} \] (Ushikubo et al., 2014), UW-3 calcite, which is from a granulite facies diopside-bearing calcite marble with

\[ \delta^{13}C = -0.91 \pm 0.08_{\text{ppm}} \] (Kozdon et al., 2009), and UW6220 dolomite with \[ \delta^{13}C = 0.79 \pm 0.71_{\text{ppm}} \] (Ferry et al., 2010).

### New standards for SIMS analysis of organic matter

For \( \delta^{13}C \) analysis of organic matter by SIMS, establishing a suite of working standards was an important part of this study. Because systematic bias between H/C and \( \delta^{13}C \) for SIMS analysis of organic matter had been previously observed in at least one other laboratory (Sangely et al., 2005; Fig. S3, this study), a suite of materials representing the wide range of H/C variability in solid organic materials was assembled. To represent the low H/C end-member, a shungite (SH95S1a; Buseck et al., 1997) and the oldest known coal, known as the Michigamme Anthracite (ca. 1840 Ma) UWMA-1 (Tyler et al., 1957; Pufahl et al., 2010) were chosen. Graphite was analyzed during our first session, but we do not recommend its use as a standard for organic carbon isotope analysis of amorphous organic materials including kerogen and pyrobitumen because it is a crystalline material, and we observe an orientation effect of several per mil. To represent the high H/C end-member, samples of Baltic and Dominican amber were used. Finally, a suite of coals was assembled from the research collection at Pennsylvania State University (PSU) to represent intermediate H/C values. These coals are identified as follows: DESC-21 (Lykens Valley #2 Seam, PA, USA), PSOC-1468 (Buck Mtn. Seam, PA, USA), PSOC-1534 (Waterfall Seam, AK, USA), UWLA-1 (LCNN Anthracite; Pappano, 2001), UWJA-1 (Jeddo Anthracite; Pappano, 2001), and UWHA-1 (Harmony Anthracite; Pappano, 2001).

### Petrography and elemental analysis

Petrography was carried out on a Hitachi S-3400 variable pressure SEM in secondary electron (SE) mode to investigate morphology and backscattered electron (BSE) mode to investigate relative chemical heterogeneity based on grayscale of the BSE image (e.g. Figs S10–S18). Images were typically acquired with an accelerating voltage of 15 kV and a working distance of 5 mm (for best spatial resolution) or 10 mm (for EDS). Elemental composition of targets and associated minerals was measured in spot or mapping modes with an EDS detector (Thermo Scientific, Waltham, MA, USA) interfaced to the Hitachi S-3400.

### Secondary-ion mass spectrometry

All SIMS measurements were performed using the CAMECA IMS 1280 large-radius multicollector ion microprobe in the WiscSIMS Lab., Department of Geoscience at the University of Wisconsin–Madison. Data in this study were gathered during eight separate analytical sessions. In general, we followed techniques described previously (Kita et al., 2009; Valley & Kita, 2009; Williford et al., 2011; Ushikubo et al., 2014), and specific analytical conditions for the different applications discussed here are given in Table 2. In all cases, sample mounts were allowed to degas for 24–48 h in the 10–7 torr vacuum of the SIMS airlock (pressure in analytical chamber is 10–9 torr). A primary beam of \( ^{133}\text{Cs}^- \) was focused at the surface of the sample; beam diameter and intensity varied depending upon the technique, but total impact energy was always 20 keV (with a secondary-ion accelerating voltage of 10 keV). An electron gun oriented normal to the sample surface and gold coating provided charge compensation. Secondary ions of the major and minor isotope(s) were collected using Faraday cup detectors (with choice of resistor depending upon abundance, below), and relevant hydride ions were measured either simultaneously (e.g., \( ^{13}\text{CH}^- \)) or by DSP2X scanning immediately after each analysis (e.g., \( ^{12}\text{CH}^-, ^{32}\text{SH}^- \)) with either electron multipliers (for low intensity beam conditions) or Faraday cup detectors (for high-intensity beam conditions). An electron multiplier was used to collect \( ^{13}\text{C}^- \) during small (1–3 µm) spot analyses. Each analysis consisted of intervals for (i) pre-sputtering (to penetrate gold coating and any minor surface contamination), (ii) centering secondary ions in the field aperture, and (iii) isotope ratio measurement.

Correction for ‘bias’, the sum of mass fractionation effects internal to the SIMS instrument, was achieved using an alpha correction as follows:

\[ \alpha = (1000 + \delta^{13}\text{C}_{\text{raw}})/(1000 + \delta^{13}\text{C}_{\text{true}}) \] (1)

where \( \alpha \) is bias, \( \delta^{13}\text{C}_{\text{raw}} \) is the uncorrected value for a standard analysis, and \( \delta^{13}\text{C}_{\text{true}} \) is the value for the standard as determined by conventional, gas-source isotope ratio mass spectrometry (for sulfur isotope analysis, \( \delta^{34}\text{S} \) is substituted for \( \delta^{13}\text{C} \)). Each set of ~10 unknown sample analyses was bracketed by eight analyses of a working standard (UWLA-1 for organic carbon, UW-3 for calcite carbon, UW6220 for dolomite carbon, and UWPy-1 for pyrite sulfur). Bias and external precision are defined for each bracket as the average \( \alpha \), and two standard deviations (2SD, expressed in permil) of the eight standard analyses, respectively. Internal precision is defined as two standard errors (2SE) of the individual isotope ratios calculated from multiple measurement cycles within each analysis. Average count rate for the major ion (e.g. \( ^{12}\text{C}^- \) or \( ^{32}\text{S}^- \)) was also calculated using the multiple measurement cycles in each analysis, and this ion count rate for each sample analysis is reported as a percentage relative to the average for the eight bracketing standard analyses as \( ^{12}\text{C}_{\text{rel}} \) or \( ^{32}\text{S}_{\text{rel}} \). Our experience with the analytical protocols and samples of this study suggests
that the intentional acquisition or interpretation of data at major ion count rates less than ~10% that of the standards should be avoided, as many possible confounding factors associated with SIMS analysis in mixed phases at low count rates can significantly degrade accuracy beyond the values reported here (see Table 1 and supplementary data).

For organic carbon analyses, bias correction was supplemented by an additional correction for H/C. To achieve this additional correction, multiple calibration standards with a wide range of H/C values (described above) were analyzed by gas-source isotope ratio mass spectrometry. Carbon isotopic and elemental compositions of standards, and bulk carbon isotope analysis methods are reported in the supplementary information. At the beginning of each SIMS organic carbon isotope analysis session, six to ten of these standards were analyzed multiple times, bracketed by analyses of the primary working standard UWLA-1. A linear regression was calculated using differential bias (α*, or the ratio of the bias for a given organic standard to the bias for eight bracketing analyses of UWLA-1) and 13CH−/12CH− (a proxy for H/C, measured simultaneously during each individual SIMS δ13C analysis) as the dependent and independent variables, respectively. Differential bias is defined as the ratio of bias for each calibration standard to bias for UWLA-1 during the calibration routine, and this method permits calculation of α* for each sample analysis using 13CH−/12CH−. For each δ13C analysis of sample organic matter, total bias correction (including H/C correction), was achieved using the following equation:

\[
\delta^{13}C = \frac{1000 + \delta^{13}C_{Raw}}{(\alpha \times \alpha^*)} - 1000.
\]  

Relative bias regression data and 95% confidence interval uncertainty calculations are reported in supplementary data. A quadratic propagation of uncertainty including external precision of bracketing standards, internal precision of individual analyses (affected in some cases by natural inhomogeneity at the scale of an individual analytical pit), and relative bias correction is also reported in supplementary data tables, as uc. For sulfur isotope analysis and carbon isotope analysis of carbonates, bias correction was achieved as above, but without inclusion of the α* term (and substitution of δ34S for δ13C).

RESULTS AND DISCUSSION

Data for SIMS δ13C analyses of organic matter, calcite, and dolomite, as well as δ34S and Δ34S of pyrite are plotted against estimated age in Fig. 1 (Δ34S data, acquired from only five of ten samples, are omitted). To maximize the clarity of this large and complex data set, we have combined the presentation of results with our interpretations. A key consideration from the outset of this study has been to present our data in a way that maximizes their relevance as the interpretive context inevitably evolves and facilitates target relocation for future analysis. To that end, we report every SIMS analysis in order of acquisition, for samples and standards in a supplementary data file. Reflected light and backscattered electron images of each SIMS sample mount are reported in a separate supplementary file, with all targets in each sample labeled (Figs S10–S18). Higher magnification SEM images of each analyzed target (acquired after analysis whenever possible to show analytical pits) are also provided in this supplementary file (Fig. S19). These post-analysis images allow the isotopic data reported here to be interpreted in the context of analyte morphology and associated mineralogy. The supplementary data file also includes additional bulk geochemical data for standards and samples, as well as bias vs. H/C calibration data and uncertainty analysis.

Organic carbon isotope analysis

**New organic carbon isotope standards for SIMS**

In session 1, beginning on December 14, 2010, ten organic materials whose bulk δ13C had been measured

<table>
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<tr>
<th>Start date</th>
<th>No.</th>
<th>Purpose</th>
<th>Spot size (μm)</th>
<th>Standard</th>
<th>2SD (‰)</th>
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</thead>
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<td>6 μm</td>
<td>UWLA-1</td>
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<td>δ13C organic matter</td>
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<td>UWPy-1</td>
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<tr>
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<td>UWPy-1</td>
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<tr>
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<td>10 μm</td>
<td>UWPy-1</td>
<td>0.09</td>
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<tr>
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<td>δ13C calcite</td>
<td>8 μm</td>
<td>UWC-3</td>
<td>0.87</td>
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previously by elemental analyzer continuous flow isotope ratio mass spectrometry (EA-CF-irMS) were analyzed by WiscSIMS to identify the material with the greatest spot-to-spot and grain-to-grain micrometer-scale carbon isotopic homogeneity. This was employed as a primary working standard for organic carbon isotope analysis with two FC detectors, and to determine whether H/C and bias are correlated on the WiscSIMS instrument as had been previously observed in another laboratory (Sangley et al., 2005). That study used Fourier transform infrared spectroscopy to analyze the aliphaticity (as a proxy for H/C) of each organic target after SIMS analysis. The alternative technique we report here has the significant advantage that H/C proxy data are acquired simultaneously with $^{13}$C/$^{12}$C ratios. Multiple grains of each of the organic carbon standards were analyzed in this session, with the exception of the two ambers, added later as high H/C end-members.

Graphite UWGr-1 was prepared as a powder from a graphite rod used for carbon arc coating with the intention to randomize crystal orientations and thus minimize orientation effects. Small ‘clumps’ of graphite powder that had been epoxy-mounted together with potential standards were analyzed three to four times each, and these showed high apparent heterogeneity ($\pm 4\%$, 2SD). We have considered whether the unexpectedly high heterogeneity we observed in graphite powder was due to variable inclusion of epoxy (that would have penetrated along grain boundaries during sample preparation) during individual analyses. Three analyses of UWGr-1 have $\delta^{13}$C$_{Raw}$ from $-58$ to $-59.5\%$, and the other seven analyses have $\delta^{13}$C$_{Raw}$ between $-62$ and $-63\%$, similar to epoxy in this standard mount that has $\delta^{13}$C$_{Raw} = -63.6 \pm 0.4\%$ ($n = 4$). However, values of $^{13}$CH/$^{12}$C for UWGr-1 range from 0.008 to 0.017, whereas values for epoxy are much higher and range from 0.138 to 0.140. We observe no correlations between $^{13}$CH/$^{12}$C or $^{13}$C count rate and $\delta^{13}$C$_{Raw}$ for UWGr-1: We thus attribute the three anomalous analyses to true heterogeneity in the graphite.

---

Fig. 1 Data from SIMS in situ carbon isotope analysis of organic matter, dolomite, and calcite, and SIMS sulfur three-isotope analysis of pyrite in ten Neoarchean samples ranging in age from ~2.7 to ~2.5 Ga. Samples were previously analyzed for bulk carbon and iron isotopes by Eigenbrode & Freeman (2006) and Czaja et al. (2010), respectively, and estimated age is assigned based on the model by Czaja et al. (2010). SIMS carbon isotope analyses of calcite and dolomite that had $^{12}$C count rates higher than the average for organic-free standards likely sampled organic inclusions and are labeled ‘above range’ (small, gray squares and diamonds). Analyses with count rates at or slightly below standard values likely sampled organic-free domains and are labeled ‘within range’ (small, black squares and diamonds). Linear correlations between $^{12}$C count rate and $\delta^{13}$C were used to calculate $\delta^{13}$C of calcite (large squares) and dolomite (large diamonds) for individual samples (see text).
Because of this heterogeneity and an orientation effect observed previously, we rejected graphite as a standard for SIMS δ¹³C analysis of amorphous organic matter in this study.

The number of grains analyzed, total number of spots, and 2SD heterogeneity for each material observed during this session are shown in Table 2. UWLA-1, prepared from LCNN anthracite, was chosen as the primary working standard due to its high spot-to-spot and grain-to-grain homogeneity (average 0.34‰ 2SD). This material enables external precision with SIMS organic carbon isotope analysis with a 6-µm spot that is comparable to bulk carbon isotope analysis with EA-CF-IRMS. To further constrain the bulk δ¹³C of this material, three individual grains of UWLA-1 were analyzed separately at Wisconsin using the sealed quartz tube method and a dual-inlet gas-source mass spectrometer (Kitchen & Valley, 1995). The results of these three analyses were remarkably consistent at −23.56‰, varying by <0.01‰. Typical precision for this method with multiple analyses of USGS24 graphite is ±0.15‰ (2SD); thus, we report the bulk δ¹³C of UWLA-1 to be −23.56 ± 0.15‰ (2SD).

During the second session, beginning April 19, 2011, seven of the ten potential standards analyzed during the first session were reanalyzed, and the two ambers were analyzed for the first time. Eighteen analyses on three grains of Baltic amber had a 2SD heterogeneity of 0.68‰ and eighteen analyses on three grains of Dominican amber had a 2SD heterogeneity of 1.2‰. Analyses of δ¹³C and ¹³CH/¹³C were used to generate the first ¹³CH/¹³C vs. bias working curve during this session. We observed a strong correlation, as shown in Fig. S3. This correlation demonstrates an effect of H/C on bias of up to 4‰ (expressed in per mil as (α − 1) *1000), indicating that monitoring this effect is particularly important when sub-permil accuracy is required. The validity of this approach is supported by correlation between ¹³CH/¹³C and H/C of the six standard materials for which we have bulk H and C abundance data. As is the case with any SIMS method, accuracy is optimized using frequent measurements of a running standard and calibration against standards that are as chemically similar to the sample as possible. The novel H/C correction method reported here, incorporating simultaneous collection of ¹²C/¹³C and H/C of the six standard materials for which we have bulk H and C abundance data. As is the case with any SIMS method, accuracy is optimized using frequent measurements of a running standard and calibration against standards that are as chemically similar to the sample as possible. The novel H/C correction method reported here, incorporating simultaneous collection of ¹²C/¹³C and H/C of the six standard materials for which we have bulk H and C abundance data.

Table 2

<table>
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<th>Material</th>
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Systematic offsets and ion count rates

Data from all analytical sessions for $\delta^{13}C$ of Neoarchean organic matter are shown in Fig. 2. The $\delta^{13}C$ data in Fig. 2 are plotted against $^{12}C_{\text{rel}}$, or the $^{12}C^{-}$ count rate of an individual analysis relative to the average count rate of the eight corresponding bracketing standard analyses (of UWLA-1). For the data set as a whole, we do not observe any systematic offset between bulk $\delta^{13}C$ and our SIMS measurements in this study. For example, SIMS $\delta^{13}C$ data for ABDP-9 148.37 are higher, and for RHDH2a 285.30 are lower than the respective bulk values for these samples. Nor do we observe a systematic change in SIMS $\delta^{13}C$ with decreasing $^{12}C_{\text{rel}}$ count rate. For $^{12}C_{\text{rel}}$ count rates between 15% and 93% of bracketing standards, data for ABDP-9 148.37 range within ~2% of the average for the sample. Internal precision, or two standard errors (2SE) of the 20 measurement cycles comprising an analysis, is also shown for each analysis in Fig. 2. Internal precision is controlled by counting statistics and signal-to-noise ratio as well as any natural isotopic heterogeneity encountered with depth in a single analysis pit. The decrease in internal precision with $^{12}C_{\text{rel}}$ is apparent in Fig. 2. Although precision is poor at low count rates, acceptable accuracy may be possible to achieve in aggregate. Nine analyses in a sample of the Wittenoom Formation (WRL-1 289.45) have $^{12}C_{\text{rel}}$ count rates lower than 15%, with an average of 6%. When the analysis with the lowest count rate (2%; n.b. $\delta^{13}C_{\text{raw}}$ for this analysis is 20% higher than any of the others in

Fig. 3 Secondary electron images showing a ~0.03-mm² region of the Carawine Dolomite (RHDH2a 252.2 m) before (A) and after (B) SIMS $\delta^{13}C$ analysis. Enlargements (B–H) show analytical pits (pit edges indicated with white arrows) from five analyses of varying proportions of pyrobitumen and calcite matrix and one analysis of apparently organic-free calcite (H). Individual analysis numbers are indicated at the top right, and $\delta^{13}C_{\text{org}}$ (VPDB) and ion count rate relative to UWLA-1 (anthracite) standard ($^{12}C_{\text{org}}$) are indicated at bottom right of each panel. Due in part to low ionization efficiency of inorganic relative to organic carbon, pure calcite yields only 2% of the $^{12}C$ counts yielded by pure organic matter, and values of $\delta^{13}C$ for these analyses are within uncertainty at organic:inorganic carbon ratios as low as 30%. Scale bars are 50 $\mu$m in (A) and (B), and 10 $\mu$m in (C–H).
the sample) is rejected, the remaining eight analyses have $\delta^{13}C = -30.0 \pm 4.5$‰, only $1$‰ different (lower) than the bulk value for this sample.

**SIMS $\delta^{13}C$ analysis of organic matter in carbonate matrices**

Kerogen in a sample of the Carawine Dolomite (RHDH2a 252.2, Fig. S14) occurs in stylolites associated with dolomite and pyrite (Fig. S7, SI). Pyrobitumen in this sample (Figs 3, S7, S14) is concentrated along cracks in a prominent calcite vein, and is sometimes associated with quartz. This pyrobitumen has $\delta^{13}C$ that is $\sim 3$‰ lower ($-52.6 \pm 0.8$‰; targets 4 and 5, Figs S14, S19), as well as a lower H/C ($^{13}CH/^{13}C = 0.65$) than the average for kerogen in this sample ($-49.8 \pm 1.4$‰; $^{13}CH/^{13}C = 0.78$; targets 6–10, Figs S14, S19). By contrast with pyrobitumen in the Mt. McRae Shale samples (ABDP-9; discussed below), the difference in both $\delta^{13}C$ and H/C, as well as the modes of occurrence of pyrobitumen and kerogen in the Carawine Dolomite sample, is consistent with post-depositional introduction of pyrobitumen together with vein-fill calcite, whereas the kerogen now concentrated in stylolites is more likely to be syndepositional. Rasmussen (2005) interpreted pyrobitumen nodules and streaks as evidence for petroleum generation and migration in the Jeerinah Fm and suggested that the pyrobitumen was locally sourced as a result of burial maturation $< 2.45$ Ga. Peak metamorphism and metasomatic alteration of the Mount Bruce Supergroup coincided with a generally northward migration of hydrothermal fluids as a result of the collision of the Pilbara and Yilgarn cratons during the Capricorn orogeny $< 2.2$ Ga (Rasmussen et al., 2005; White et al., 2014a). It is possible that (pyro)bitumen was redistributed or emplaced in the Carawine Dolomite as a result of this event. A similar inference of large-scale migration has been made for vein pyrobitumen with higher $\delta^{13}C$ than the in situ kerogen in the Jeerinah Fm on the basis of NanoSIMS analyses (Rasmussen et al., 2008).

Five analyses of varying amounts of pyrobitumen in a calcite matrix made during session 2 indicate that WiscSIMS organic carbon isotope analysis of organic domains in carbonate using the $6 \mu$m, two FC techniques at count rates as low as $30$% (i.e., $30$% organic matter in carbonate) of the average for the working standard are possible with near permil accuracy (Fig. 3). Four of these five analyses of pyrobitumen in calcite (with $^{12}C_{rel}$ from 0.92 to 0.29) are tightly distributed along a linear trend of $\delta^{13}C$ vs. $^{12}C_{rel}$ indicating a positive shift of only $0.4$‰ from the highest to lowest count rate. A single analysis with $^{12}C_{rel} = 0.31$ indicates a positive shift of $1.3$‰ from the analysis with the highest count rate.

One analysis of an organic carbon-free domain of calcite adjacent to pyrobitumen has a count rate only $2$% that of the working standard carbonate relative to organic carbon with this SIMS technique. Multiple factors are likely responsible for the low relative C/C₀ production in carbonate relative to organic carbon. Calcite and dolomite are $\sim 12$ and $13$% C by mass, compared to typical sedimentary organic matter that is $60$ to $> 90$% C by mass. Carbonate carbon also has a lower ionization efficiency than organic carbon (see further discussion below). A mixing model (not shown) assuming a $\sim 40$‰ difference in inorganic and organic $\delta^{13}C$ and a $2$% contribution from calcite suggests that the threshold for sub-permil accuracy would be $^{12}C_{rel} = 0.5$. Work continues to understand the constraints to accuracy in SIMS $\delta^{13}C$ analysis of small organic domains in carbonate, and the possible applications are numerous.

**Fig. 4** Secondary electron images (A, C) and energy-dispersive x-ray spectroscopic elemental maps (B, D) small (3 μm) spot $\delta^{13}C$ analyses (VPDB ± 2SD external precision) of typical kerogen (ker) in a shale sample from the ~2.6 Ga Marra Mamba Iron Formation (WRL-1 659.55 m). Kerogen in this sample is associated with dolomite (Dol), siderite (Sid), quartz (Qtz), chlorite (Clc), pyrite (Py). Pressure solution features and micrometer-scale veining are apparent at intersections between quartz and siderite in (A). Scale bars are 10 μm.
Spatial resolution and heterogeneity

A total of 121 organic targets in nine Neoarchean samples ranging in age from ~2.7 Ga to ~2.5 Ga were analyzed in this study. Most targets were analyzed using a 6-μm-diameter beam and Faraday cup detectors for both 12C\(^{13}\)C and 13C\(^{13}\)C. To investigate the limits of spatial resolution for the instrument, we analyzed six targets each in samples of the Marra Mamba Iron Fm (WRL-1 659.55 m) and the Jeerinah Fm (WRL-1 679.90 m) with a 3-μm-diameter beam, using an electron multiplier for 13C\(^{13}\)C (Fig. 4). Average SIMS \(\delta^{13}C\) values in these two samples are 3.1 and 0.4\(^{13}ppm\) lower than bulk \(\delta^{13}C\) values, respectively, and we have no reason to suspect a decrease in accuracy with the smaller beam condition (although external precision using this condition is not as good as the larger, 6 μm diameter condition due to fewer secondary ion counts). We started this session using a 1-μm-diameter beam to analyze 1 μm or smaller targets; however, the ~1 μm uncertainty in stage motion (due to backlash) combined with our inability to resolve 1-μm features in the optical system of the IMS 1280 made targeting difficult. To compensate for this difficulty, we increased the size of the beam to 3 μm, which can be imaged during analysis, and generated 13C\(^{13}\)C- ion images (Fig. S6) prior to analysis to facilitate accurate aiming. Software (digital zoom) and hardware (optical registering at 0.1-μm scale), and optical (365-nm sample illumination, Kita et al., 2015) improvements in the WiscSIMS instrument made after completion of this study have improved targeting, and we frequently achieved 1 μm targeting accuracy in later efforts, although analysis spots should always be verified by SEM. Figure 4 shows representative targets and analytical pits for small spot 13C analysis in sample WRL-1 659.55 m.

The analytical techniques presented here are designed to evaluate micrometer-scale variability, including correlations between isotopic composition and morphology, mineralogy, and elemental compositions, and in this study we report several such correlations that are invisible to bulk techniques. However, given (i) a sufficient number of SIMS analyses, (ii) sufficient accuracy of SIMS analyses, and (iii) sufficient sampling of characteristic phases in a particular sample, then the average SIMS value should be similar to the bulk value for that sample. In this study, bulk \(\delta^{13}C\) values and averaged SIMS \(\delta^{13}C\) values are remarkably consistent. The difference between the averaged SIMS value and the bulk value was calculated for each sample, and the average of these calculated differences is 0.02%\(^{13}\)C.

It is important to distinguish apparently random heterogeneity observed in SIMS data (that could simply indicate poor precision and/or accuracy) from systematic variability that can be correlated to textures or compositions (and is a signature of a natural process operating in the sedimentary or diagenetic system). Five of the nine samples analyzed in this study had a range in SIMS \(\delta^{13}C\) of less than 5%\(^{13}ppm\), and the average range is 7.4%\(^{13}ppm\) (Table 3). A sample of the ~2.7 Ga Tumbiana Fm exhibits the most dispersion observed in this study, and the dispersion is demonstrably real by systematic correlation to textures. A single kerogenous domain associated with filamentous TiO\(_2\) has \(\delta^{13}C = -52.3 \pm 0.1\) for \(n = 2\), consistent with the lowest values reported in earlier studies of Tumbiana Fm drill core (Eigenbrode & Freeman, 2006; Thomazo et al., 2009a), suggesting a metabolically recycled, methanotrophic carbon source. More typical kerogen observed in this sample, associated with clay minerals and pyrite, has \(\delta^{13}C = -32.8 \pm 3.6\) for \(n = 4\); e.g., Fig. 5). This kerogen component is isotopically similar to a relatively high %\(^{13}C\) organic component in the Tumbiana Fm identified by earlier bulk measurements (Eigenbrode & Freeman, 2006; Thomazo et al., 2009a), and likely derives from the bio-

### Table 3

Average and range of SIMS \(\delta^{13}C\) measurements of organic matter and/or carbonate minerals in eleven Neoarchean rock samples. Values of \(\delta^{13}C\) for dolomite (\(\delta^{13}C_{\text{calc}}\text{calc})\) and calcite (\(\delta^{13}C_{\text{calc}}\text{calc})\) were estimated for samples that contained varying degrees of inter- and intracrystalline organic matter in analyzed carbonate domains that likely drives linear relations between \(\delta^{13}C\) and \(^{12}C\) count rate relative to organic-free bracketing standards by assuming carbonate \(\delta^{13}C\) homogeneity and using an interpolated value at \(^{12}C_{\text{calc}} = 1.0\). These values were then used to calculate the average offset between organic carbon, calcite (\(\Delta^{13}C_{\text{calc-org}}\)) and/or dolomite (\(\Delta^{13}C_{\text{calc-org}}\)) for each sample

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<th>Depth (m)</th>
<th>Unit</th>
<th>Est. Age (Ma)</th>
<th>(\delta^{13}C_{\text{org}}) (‰, VPD8)</th>
<th>(\delta^{13}C_{\text{calc}}\text{calc})) Max</th>
<th>Min</th>
<th>Avg</th>
<th>(\Delta^{13}C_{\text{calc-org}}) (‰)</th>
<th>(\Delta^{13}C_{\text{calc-org}}) (‰) Max</th>
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Fig. 5 Two types of kerogen situated ~6 mm apart in a single sample of pyritic shale from the ~2.7 Ga Tumbiana Formation (SV-1 55.3 m; Fig. S16) indicating the preservation of multiple metabolic signatures. Type A kerogen is associated with filamentous TiO₂ and has low Δ²⁷C. Type B kerogen is associated with clay minerals and pyrite (Py), and has higher Δ²⁷C. Values of Δ²⁷C for these two different kerogen types bound the bulk rock value for this sample as shown at the bottom. Scale bars are 10 μm.

Fig. 6 Secondary electron (A, D), backscattered electron (B), and energy-dispersive x-ray spectroscopic elemental map (C) of pyrobitumen (OM) surrounding a uraniferous (U) mineral grain (A, C) and typical kerogen (B, D) in a sample of the Mount McRae Shale (ABDP-9 111.22 m). Analytical pits from SIMS Δ²⁷C analysis of organic matter (OM), calcite (Cal) and dolomite (Dol) are shown, and external precision for these analyses is indicated at lower right in panel (D). Pyrobitumen in this sample is associated with detrital quartz (Qtz) and chlorite (Chl), and is ~8‰ higher in Δ²⁷C than co-occurring kerogen in this sample, the latter suggesting either a different source or alteration mechanism for these two organic components in the sample. Calcite in this sample is likely secondary, but the offset between dolomite and kerogen Δ²⁷C provides an estimate of the apparent metabolic fractionation: ~34‰. Scale bars are 10 μm.
two classes of organic matter in close spatial association that are (i) indigenous and syngenetic, and (ii) distinct in morphology, associated mineralogy, and carbon isotopic composition represent a robust biosignature (Lepot et al., 2013; Williford et al., 2013).

The bulk δ13C value for this sample of the Tumbiana Fm is −49.7‰, suggesting that the low-δ13C phase we observed, or another similar phase is relatively abundant but may typically occur in nanometer-scale domains that we did not sample (and would be too diffuse for accurate SIMS analysis). This highlights an important difference between spatially resolved SIMS analysis and bulk analysis: although the former potentially yields far higher ‘information density’ by increasing spatial resolution, a selection bias introduced by analysis parameters (SIMS beam diameter, spatial resolution of light, and electron microscopy techniques, etc.) may cause important phases to be overlooked. Complementary molecular analyses, such as H/C and N/C mapping with NanoSIMS (Wacey et al., 2010), micro-Raman, or X-ray absorption spectroscopy (Lepot et al., 2009), may help to track distinct micro-reservoirs of organic matter and constrain their importance in the studied rocks. Bulk and spatially resolved analyses are complementary, and an integrated approach will maximize the information yield for a sample.

Sample ABDP-9 148.37b contains pyrobitumen nodules surrounding central sulfide grains, and these domains have slightly higher δ13C (−33.2 ± 0.4‰) than, but similar H/C (13CH/13C = 0.069) to the average for kerogen in this sample (−35.0 ± 0.8‰; 13CH/13C = 0.066). A similar pyrobitumen nodule with a central uranium-rich grain (Fig. 6A,C) in ABDP-9 111.22 was analyzed three times, and it has significantly higher δ13C (−26.9 ± 0.8‰) than the average for kerogen in the sample (−35.0 ± 0.8‰), but H/C (13CH/13C) in pyrobitumen is also similar (0.068) to kerogen (0.066). This δ13C enrichment in 13C observed in the Mount McRae pyrobitumen is larger than can be explained by typical thermal maturation processes (Des Marais, 2001). Similar associations in the Jeerinah Fm have been previously reported as evidence for Archean petroleum generation (Rasmussen, 2005), and it is possible that the 13C enrichment observed here resulted from radiolytic alteration of pyrobitumen, as has been proposed for the uraniumiferous Cigar Lake (Landais et al., 1993) and Athabasca (Sangély et al., 2007) deposits. Alternatively, the U-associated pyrobitumen observed in ABDP-9 111.22 may originate from another source rock.

Pyrobitumen nodules such as those described above do not occur in the oldest sample of the Mount McRae shale analyzed in this study (ABDP-9 161.02). Organic matter in this sample occurs in cores of ‘nodular’ associations of siderite and pyrite (Fig. 7C–D) and has low δ13C (−40.4 ± 1.6‰). A single large pyrobitumen nodule surrounding a small (~3 μm) central grain of chalcopyrite, and associated with chlorite and pyrite, was observed in a sample of the Jeerinah Formation (RHDH2a 285.3#1;
Carbonate carbon isotope analysis

As discussed previously, the conversion of inorganic carbon to C\(^{13}\) is a multistep process and inefficient (e.g., ~0.9 MHz \(^{12}\)C/nA/wt.% for calcite standards) relative to production of C\(^{13}\) from organic carbon (~4.7 MHz \(^{12}\)C/nA/wt.% for anthracite standards) during SIMS. This affords the unique ability to accurately analyze the isotopic composition of small organic carbon domains in carbonate matrices. However, the high relative efficiency of organic carbon ionization presents a challenge for accurate carbon isotope analysis of sedimentary carbonates that have small organic inclusions. In this study, we analyzed 80 targets of calcite and dolomite in seven samples (ABDP-9 111.22 and 148.37; RHDH2a 252.2 and 285.3; WRL-1 289.45 and 520.6; SV-1 55.3) and four siderite targets in one sample (ABDP-9 161.02). In general, achieving accurate individual SIMS analyses of carbonate was impossible due to the presence of organic matter in the analytical region.

We used \(^{12}\)C\(_{rel}\) of each carbonate analysis to assess whether organic inclusions had been analyzed: \(^{12}\)C\(_{rel}\) > 1.0 typically indicated the presence of organic matter. We note that further complexity in \(^{12}\)C count rates can be introduced by the presence of non-carbon bearing (e.g., quartz) inclusions in carbonate, but we suggest based upon our petrographic observations (Fig. S19) that the extent to which this affects our conclusions is minimal, as inclusions of carbon-free minerals are readily detected with SEM and EDS. In some samples, a linear correlation between \(^{13}\)C and \(^{12}\)C\(_{rel}\) was observed, with higher count rates yielding lower \(^{13}\)C. In these cases, it was possible to estimate a \(^{13}\)C value for organic-free carbonate minerals (similar to a bulk value) using the \(^{13}\)C value calculated for \(^{12}\)C\(_{rel}\) = 100% (an example is shown in Fig. S9). These \(^{13}\)C\(_{calc}\) calc and \(^{13}\)C\(_{cal}\) calc values are shown in Table 3. Further, it was then possible to calculate \(^{13}\)C\(_{dol-org}\) and \(^{13}\)C\(_{cal-org}\) using average SIMS \(^{13}\)C of organic matter in the samples.

Primary vs. secondary signals

With the exception of the millimeter-scale calcite vein in the Carawine Dolomite (RHDH2a 252.2) with \(^{13}\)C\(_{cal}\) calc = 0.5\(^{\text{‰}}\), all estimates for \(^{13}\)C of carbonate minerals are <0\(^{\text{‰}}\). Estimates for dolomite range from –9 to –1.4\(^{\text{‰}}\) and for calcite from –18.5 to –7.1\(^{\text{‰}}\). The strongly negative \(^{13}\)C estimated for most calcites analyzed in this study suggests that they precipitated after deposition in the presence of remineralized organic carbon. It is possible, however, that calcite in sample RHDH2a 252.2 represents a primary phase formed in surface waters and that dolomite formed in sediments in the presence of organic matter. The relative recalcitrance of this early diagenetic dolomite would have led to its survival of the dissolution event that formed the stylolites in this sample.

Assessing metabolic and diagenetic signatures using paired organic and inorganic SIMS carbon isotope analysis

Given the difficulty of accurate SIMS \(^{13}\)C analysis of sedimentary carbonate minerals introduced by small organic inclusions (discussed previously), we caution that the data presented here for inorganic carbon are less robust than those for organic carbon and pyrite sulfur. In some cases, however, our SIMS data align well with earlier conventional \(^{13}\)C\(_{calc}\) data from the same samples, validating the approach we outline here. With these caveats in mind, and because this approach may be useful in future investigations, we present the following as a case study for this technique.

Although the minimum \(^{13}\)C of organic matter is quite low (<-40\(^{\text{‰}}\)) in the majority of samples, RHDH2a 252.2 (Carawine Dolomite) is the only sample for which paired, \(in\;situ\) inorganic and organic carbon isotope data from both calcite and dolomite suggest a large (>40\(^{\text{‰}}\)) \(^{13}\)C\(_{calc-org}\) offset inconsistent with carbon fixation via the Calvin–Benson–Bassham (CBB) reductive pentose phosphate pathways used by photo- and chemo-autotrophs. This is broadly consistent with earlier evidence from bulk \(^{13}\)C data (Eigenbrode, 2004; Eigenbrode & Freeman, 2006), strengthens the case for a significant paleoecological role of methanotrophy in the Neoarchean (Hayes, 1983, 1994; Eigenbrode & Freeman, 2006) and provides strong, new evidence for the presence of this metabolism in shallow shelf environments (likely in sediments).

In ABDP-9 111.22 and 148.37, \(^{13}\)C\(_{dol-org}\) is 30.1\(^{\text{‰}}\) and 33.2\(^{\text{‰}}\), respectively (Table 3). This is self-consistent and agrees with previously reported, paired bulk inorganic and organic \(^{13}\)C data for ABDP-9 111.22 that showed \(^{13}\)C\(_{calc-org}\) = 30.1\(^{\text{‰}}\) (Kaufman et al., 2007). However, \(^{13}\)C\(_{calc-org}\) is smaller in these samples, 24.4\(^{\text{‰}}\) and 23.6\(^{\text{‰}}\), respectively (Table 3). An extrapolation of the equilibrium carbon isotope fractionation expression for coexisting metamorphic calcite and dolomite to 20°C indicates that dolomite is \(^{13}\)C-enriched by 2.4\(^{\text{‰}}\) relative to calcite (Sheppard & Schwarzb, 1970). If calcite is a primary carbonate phase in the upper Mount McRae Shale as previously reported by Kaufman et al. (2007), and represented a reli-
In three separate analytical sessions during this study, 266 pyrite targets were analyzed for sulfur three-isotopic composition with a ~10-μm-diameter beam. After minimal repolishing to remove analytical pits from sulfur three-isotope analysis, 58 additional pyrite targets were analyzed for sulfur four-isotopic composition using a ~20-μm-diameter beam. Values of $\delta^{34}$S and $\Delta^{34}$S measured in this study of 10 samples of <1 cm$^2$ each encompass nearly the entire range reported in earlier compilations of bulk sulfur three-isotope data from Neoarchean samples (Williford et al., 2011; Canfield & Farquhar, 2009; Fig. 8). This suggests that significant unrecognized variability exists in samples that have previously been analyzed only by bulk techniques, consistent with other SIMS studies of Archean and Proterozoic sulfides (Philippot et al., 2007, 2012; Bontognali et al., 2012; Farquhar et al., 2013; Johnson et al., 2013; Fischer et al., 2014; Zhelezinskaia et al., 2014).

**Morphologically correlated sulfur isotopic heterogeneity in the Jeerinah Fm**

Of forty pyrite targets analyzed in sample WRL-1 679.9 of the Jeerinah Fm, 32 define a relatively tight field with $\delta^{34}$S ranging from $-0.9_{\text{vo}}$ to $4.1_{\text{vo}}$ and $\Delta^{34}$S from $-10_{\text{vo}}$ to $0.7_{\text{vo}}$ (Fig. 9C). These pyrites occur in agglomerations of euhedral to subhedral crystals whose relatively narrow range of sulfur isotopic compositions is consistent with early diagenetic formation in sediments from (either abiotic or microbial) sulfate reduction. In the same sample, another group of 8 targets defines a linear array with a larger range of mass-dependent ($\delta^{34}$S from 4.0$_{\text{vo}}$ to 13.4$_{\text{vo}}$) and mass-independent ($\Delta^{34}$S from 4.8$_{\text{vo}}$ to 9.9$_{\text{vo}}$) fractionation (Fig. 9). This more isotopically heterogeneous group of pyrites generally occurs as isolated, individual grains within the clastic matrix, sometimes with rounded margins, suggestive of a detrital component in this sample. The large range in $\delta^{34}$S coupled with strongly positive S-MIF suggests metabolic recycling of sulfur (disproportionation followed by sulfate reduction) that left the atmosphere as $S^0$ (Pavlov & Kasting, 2002; Ono et al., 2003; Kaufman et al., 2007). However, the rounded grain morphologies (that may indicate a detrital component) suggest that this record of the sulfur cycle may predate the deposition of the host rock.

The roughly time-equivalent sample of the Jeerinah Fm from the RHDH2a core (285.3) shows a similar separation in sulfur isotopic compositions (Fig. 9B), with one small group of six pyrite targets having $\delta^{34}$S between $-0.8_{\text{vo}}$ and $1.8_{\text{vo}}$ and $\Delta^{34}$S between $0.8_{\text{vo}}$ and $4.0_{\text{vo}}$ and a larger group defining a linear array with $\delta^{34}$S from $2.2_{\text{vo}}$ to $15.3_{\text{vo}}$ and $\Delta^{34}$S from $6.9_{\text{vo}}$ to $14.4_{\text{vo}}$ (Fig. 9). The petrographic difference between these two groups of Jeerinah Fm pyrite in the RHDH2a sample is not as clear as they are for WRL-1. Pyrite targets in RHDH2a 285.3 are generally relatively large (10s to >100 μm), commonly subangular to partially rounded, and associated with pyrobitumen and Fe- and Mg-bearing chlorite.

Intriguingly, the sample of the Jeerinah Fm (RHDH2a 285.3) interpreted to have been deposited closer to the basin margin is dominated by pyrite with diverse sulfur iso-
topic compositions distributed along a linear array of mostly positive $\Delta^{33}S$ and $\delta^{34}S$, whereas the sample from the depocenter (WRL-1 679.9) is dominated by pyrite with $\Delta^{33}S$ and $\delta^{34}S$ closer to zero. It may be that this latter variety of pyrite (with near-zero $\Delta^{33}S$ and $\delta^{34}S$) represents relatively strong sulfate limitation in the depocenter, whereas the former variety of pyrite (with wide-ranging $\Delta^{33}S$ and $\delta^{34}S$) represents conditions of sulfur cycling restricted to the margin (e.g., metabolic recycling with less sulfate limitation). If so, then pyrite in WRL-1 679.9 with $\Delta^{33}S$ and $\delta^{34}S \geq 5$ may indeed be detrital, having washed into deeper water from the basin margin—consistent with our petrographic observations.

Previous $\delta^{34}S$ measurements of Jeerinah Fm pyrite using laser ablation showed 6–7 mm-scale heterogeneity, interpreted as strong evidence for microbial sulfate reduction (Kakegawa et al., 2000; Kakegawa & Nanri, 2006). The sample of the Jeerinah Fm analyzed in this study exhibits even larger, micrometer-scale heterogeneities in $\delta^{34}S$, $\Delta^{33}S$, and $\Delta^{36}S$ (e.g., RHDH2a 285.3#16 and #19; Fig. S19). Over a distance of ~20 μm, from the core of target RHDH2a 285.3#16 to the rim, $\delta^{34}S$ increases by 6.2‰ (from 3.5 to 9.7‰, 2SD) and $\Delta^{33}S$ increases by 3‰ (from 7.8 to 10.8‰, 2SD). Another target (#19) in the same sample (RHDH2a 285.3) shows differences of 12.7‰ (2.6–15.3‰, 2SD) in $\delta^{34}S$, 7.5‰ (6.9–14.4‰, 2SD) in $\Delta^{33}S$, and −5.5‰ (−5.8 to −11.3‰, 2SD) in $\Delta^{36}S$ over only a distance of only ~40 μm. Not only is this pyrite grain extremely heterogeneous, it exhibits the largest S-MIF yet reported from any rock (Philippot et al., 2012). Although much larger gradients in $\delta^{34}S$ (>30‰) have been observed at smaller (<4 μm) spatial scales using the extremely high spatial resolution SIMS techniques we reported in an earlier study of

Fig. 9 Sulfur three-isotope data from this study shown for individual samples in ABDP-9 (A), RHDH2a (B), WRL-1 (C), and SV-1 (D) drill cores. External precision is smaller than symbols.
Paleoproterozoic pyrite (Williford et al., 2011), to our knowledge the gradients in $\Delta^{33}S$ and $\Delta^{36}S$ we report here for the Jeerinah Fm are the largest so far observed.

Isotopic relations correlated with magnetic anomalies, but otherwise similar to those we report from the Jeerinah Fm, have recently been reported from Archean pyrite nodules in the Barberton Greenstone Belt using scanning high-resolution superconducting magnetic (SQUID) microscopy SQUID-SIMS, and these were interpreted to have resulted from dynamic mixing of locally heterogeneous and compositionally evolving sulfur sources, and metasomatic alteration (including pyrite to pyrrhotite transformations indicated by magnetic domains in pyrite) that postdated deposition by hundreds of Myr (Fischer et al., 2014). Although the resolution of the SQUID-SIMS technique (50-µm SQUID step size, ~25-µm SIMS analytical area) reported by Fischer et al. (2014) is insufficient to assess whether the isotopic heterogeneities reported here (e.g., over scales of ~20 µm in a ~50-µm grain) are associated with magnetic anomalies, we suggest that some of the variability we observe may be driven by similar mechanisms to those hypothesized in that study. Similarly, distinction of zonation of minor elements in pyrite (e.g., As, Ni, Co) by electron probe microanalysis (EPMA) can be achieved with a lateral resolution on the micrometer scale. This method, coupled to SIMS, has been used to distinguish multiple generations of pyrite with correlated elemental and isotopic heterogeneities in individual grains (Williford et al., 2011) and nodules (Marin-Carbonne et al., 2014).

Accordingly, we apply extraordinary care in target selection and analysis in order to understand the degree to which post-depositional processes have affected our samples. To assess the confidence with which we have identified our sulfur isotope analysis targets as pyrite, we analyzed pyrite and pyrrhotite standards by EDS. Although EDS does not provide the accuracy in atom% associated with wavelength-dispersive spectroscopy (e.g., with EPMA), pyrite and pyrrhotite are quickly and clearly distinguishable by their relative Fe and S peak heights using this technique. At 15 keV, the S/Fe peak height ratio is typically 4 for pyrite and 2 for pyrrhotite. We find no evidence of pyrite to pyrrhotite transformations in the analytical targets reported here.

Associations between pyrite, pyrobitumen, and chlorite in the Jeerinah Fm suggest that thermochemical sulfate reduction associated with metamorphism/metamoturbation may have played at least a secondary role in pyrite formation; that is, a ‘primary’ episode of early diagenetic, microbial sulfate reduction may have been followed by a later (perhaps by 100s of Myr), ‘secondary’ episode of abiogenic sulfate reduction accompanied by evolution of pyrobitumen from kerogen and metamorphic chlorite transformations. Indeed, anomalous sulfur isotope fractionation effects completely decoupled from atmospheric chemistry – such as those observed in laboratory experiments with thermochemical sulphate reduction (Lasaga et al., 2008; Watanabe et al., 2009) – may explain some of the intragrain heterogeneity reported here. Partial dissolution, remobilization, and local redeposition could also have contributed to these heterogeneities, although we find no evidence for introduction of secondary minor elements (e.g., Ni, Co, As) during late diagenesis as has been reported for other sedimentary pyrite with strong sulfur isotope zonation (Williford et al., 2011).
Atmospheric and metabolic effects on sulfur isotope systematics

Pyrite grains in the oldest samples analyzed in this study, from the ~2.7 Ga Tumbiana Fm, have universally negative $\delta^{34}S$, as low as $-9\%_\text{ref}$, but they show very little heterogeneity in $\Delta^{33}S$ (-0.3 to 0.9). This is consistent with previous bulk sulfur isotope analysis of pyrite in the Tumbiana Fm (Thomazo et al., 2009a) and the Manjeri and Cheshire Fms of Zimbabwe (Thomazo et al., 2013), and indicates that the dampened S-MIF characteristic of the Mesoarchean (Farquhar et al., 2007b) persisted into the early Neoarchean. Organic matter in samples of the Tumbiana Fm with $\delta^{13}C$ as low as $-52.4\%_\text{ref}$ supports the earlier contention that methanotrophy may have played a role in the relatively small range of S-MIF observed between ~3.2 and 2.7 Ga, and below we discuss similarly coupled carbon and sulfur isotope systematics evident in the younger Neoarchean record from Australia.

Slopes of linear $\Delta^{33}S/\delta^{34}S$ arrays for Jeerinah Fm pyrite targets in the WRL-1 (depocenter) and RHDH2a (margin) are indistinguishable (0.5; Fig. 9), but the $\Delta^{33}S$ intercept is higher (6.9) for RHDH2a pyrites compared to WRL-1 pyrite (3.1). We interpret the similar slope to indicate stability of atmospheric MIF-S production (on a time scale) during the deposition of the Jeerinah Fm. The mechanism behind the different intercepts is more difficult to understand in the depositional context. A higher $\Delta^{33}S$ intercept suggests an increased role of $S_0$ relative to $SO_4^{2-}$ in the formation of the RHDH2a Jeerinah pyrite compared to the WRL-1 pyrite. This conclusion is based upon a model of S-MIF production in which positive $\Delta^{33}S$ is carried from the atmosphere to the ocean by elemental sulfur, and negative $\Delta^{33}S$ by sulfate (Pavlov & Kasting, 2002; Ono et al., 2003).

Sulfur four-isotope compositions in the Carawine Dolomite (RHDH2a 252.2; Fig. 10B) depart from the $\Delta^{36}S/\Delta^{33}S$ array observed for other samples in this study (and previous studies; e.g., Farquhar et al., 2000; Ono et al., 2003). Intriguingly, organic matter in this sample of the Carawine Dolomite has lower typical $\delta^{13}C$ than any other sample in this study (avg. $-49.5\%_\text{ref}$), adding to the growing body of evidence for a connection between methane and S-MIF. Rotation of the $\Delta^{36}S/\Delta^{33}S$ array away from a typical slope of $-0.9$ toward a slope of $-1.5$ has been interpreted as evidence for changing atmospheric production of S-MIF, with an ‘organic haze’ as the hypothesized mechanism (Domagal-Goldman et al., 2008; Thomazo et al., 2009a; Zerkle et al., 2012; Farquhar et al., 2013). Zerkle et al. (2012) defined the $\Delta^{36}S/\Delta^{33}S_{\text{dev}}$ index to describe this behavior, and using bulk measurements, found that the greatest deviation ($\Delta^{36}S/\Delta^{33}S_{\text{dev}} \geq 2$) occurred in samples with $\delta^{13}C_{\text{org}} < -40\%_\text{ref}$. A subset of the samples analyzed in bulk by Zerkle et al. (2012) were analyzed by Farquhar et al. (2013) for sulfur four isotopes using the same instrument (CAMECA IMS 1280) as we used in this study, but in a different laboratory with different analytical conditions (e.g., electron multiplier rather than Faraday cup detector for $^{36}S$). These $\textit{in situ}$ data were consistent with the findings of Zerkle et al. (2012), and the authors further argued for two distinct pathways of Archean pyrite formation (see below; Farquhar et al., 2013).

Samples analyzed in this study with the lowest $\delta^{13}C$ values of organic matter (and average $\delta^{13}C$ of organic matter $<40\%_\text{ref}$ Table 3), including the Carawine Dolomite, Jeerinah Fm, and the lowest sample of the Mount McRae Shale, also exhibit the highest values of $\Delta^{36}S/\Delta^{33}S_{\text{dev}}$ (as high as 2.2, 1.7, and 2.5, respectively; supplementary data). Sulfur isotope data from the Carawine Dolomite depart from the Archean array with (loose) linearity. Similar systematics have been observed in Phanerozoic fossils preserved by pyritization of organic matter (Ono et al., 2006). Accordingly, the departure from the Archean $\Delta^{36}S/\Delta^{33}S$ array observed in the Carawine Dolomite could result from mass-dependent effects expressed through reaction networks associated with microbial sulfur metabolism (Farquhar et al., 2003, 2007a; Johnston et al., 2007, 2008; Zerkle et al., 2010). Archaeal anaerobic methanotrophy (ANME) and bacterial sulfate reduction are coupled by consortia in modern ecosystems (Orphan et al., 2001a,b), and the data presented here may indicate a similar connection between these organisms in the Neoarchean as recently proposed by Thomazo et al. (2013).

Sulfur isotope systematics in the ~2.5 Ga Mount McRae Shale

Multiple lines of geochemical evidence from samples of the Mount McRae Shale recovered from the ABDP-9 core, including sulfur three-isotope measurements, were used to argue for oxygenation in the surface ocean (Anbar et al., 2007; Kaufman et al., 2007). In particular, a major transition from pyrite with uniformly positive $\delta^{34}S$ and $\Delta^{33}S$ below 153 m in the ABDP-9 core to pyrite with negative $\delta^{34}S$ and positive $\Delta^{33}S$ was interpreted to represent emergence of an oxidative sulfur cycle, and a shorter interval between 153 and 135 m with negative values of both $\delta^{34}S$ and $\Delta^{33}S$ was interpreted to represent increasing seawater sulfate concentrations due to oxidative weathering (Kaufman et al., 2007).

In multiple analyses of pyrite from only three samples, we observe the entire range of sulfur isotopic heterogeneity so far detected by bulk analysis in the ABDP-9 core (Figs. 9 and 10). Our data are broadly consistent with the earlier bulk measurements and interpretations of the sulfur
isotopic composition of pyrite in the Mount McRae Shale (Ono et al., 2003; Kaufman et al., 2007), but the high spatial resolution, *in situ* nature of our analysis, and the addition of $\Delta^{36}$S measurements offer powerful new windows into the biogeochemistry of this unit. In the oldest sample, taken 8 m below the ‘153 m transition’ (161.02 m), $\delta^{34}$S varies from −0.9 to 18.7‰, $\Delta^{32}$S from −1.6 to 10.9‰, and $\Delta^{26}$S from −7.3 to 2.6‰. The range in $\delta^{34}$S and $\Delta^{33}$S encompasses the range reported from bulk measurements below 153 m in the ABDP-9 core. Pyrite in this sample exhibits strong linearity in $\Delta^{33}$S/$\Delta^{36}$S ($r^2 = 0.99$), with a slope of −0.9, typical for the Archean ($\Delta^{26}$S = −0.88$\Delta^{34}$S + 1.39; Fig. 10A). A shift in the $\Delta^{36}$S intercept in the $\Delta^{36}$S/$\Delta^{33}$S array to 1.4‰ can be driven by changing fractionation during sulfate reduction (Ono et al., 2006), but increasing fractionation would tend to shift the intercept in the negative direction (Zerkle et al., 2012). Values of $\delta^{34}$S are high, and almost uniformly positive, implying low seawater sulfate concentrations and a greater relative contribution of elemental sulfur (by way of polysulfide intermediates; Farquhar et al., 2013) than sulfate to pyrite formation.

Of nine targets in ABDP-9 161.02 chosen for sulfur four-isotope analysis, eight have $\Delta^{34}$S/$\Delta^{33}$Sdev > 1.2, and one has $\Delta^{36}$S/$\Delta^{34}$Sdev = 2.5, comparable to the high values observed in the a Upper Nauga Fm, similar in age (~2.5 Ga) to the Mount McRae Shale (Zerkle et al., 2012). Values of $\Delta^{36}$S/$\Delta^{34}$Sdev vary from 0.9 to 2.5 (with an average of 1.5), and this sample also has the lowest $\delta^{13}$C (~−42‰) recorded in the ABDP-9 core. Critically, however, the slope of the $\Delta^{36}$S/$\Delta^{34}$S array in this sample is not rotated from typical Archean values, but ‘anomalous’ $\Delta^{36}$S/$\Delta^{33}$Sdev values are driven by a change in $\Delta^{36}$S intercept.

Target 23 in ABDP-9 161.02 exhibits a large gradient from the interior to the exterior of the nodule: the interior has $\delta^{34}$S = 0.5‰ and $\Delta^{33}$S = 1.2‰ whereas the exterior has $\delta^{34}$S = 9.6‰ and $\Delta^{33}$S = 7.3‰ (Fig. S19). Only the exterior of the nodule (exposed by repolishing the sample between sulfur three- and four-isotope analysis sessions) was analyzed for $\Delta^{36}$S, so we do not know whether or not a gradient also exists for $\Delta^{34}$S/$\Delta^{33}$S in this target. The large (9‰) gradient in $\delta^{34}$S, distributed over only a few micrometer, suggests Rayleigh fractionation during formation of the nodule.

A recent and detailed petrographic and geochemical analysis of nodular associations of organic matter, pyrite, and siderite in Mississippian shales from Wardie, Scotland (Bojanowski & Clarkson, 2012), reports textures similar to those we observe in ABDP-9 161.02 (see especially Fig. 4 in that publication). Based upon $\delta^{13}$C values in siderite >5‰, as well as organic matter associations, these authors suggest that the precipitation of Wardie siderite relied on alkalinity produced during methanogenesis, initiated in the sulfate reduction zone after rapid consumption of sulfate. We propose a similar mechanism: microbial sulfate reduction proceeded under low sulfate concentrations typical of Archean seawater until sulfate was exhausted, at which point methanogenesis and disimilatory iron reduction provided the conditions required for siderite formation. This is consistent with the gradient we observe in $\delta^{34}$S, but it does not immediately explain the gradient in $\Delta^{33}$S from the core to rim of a nodule, observed, for example, in ABDP-9 161.02 target 23 (Fig. S19). Such a gradient, similar to those observed in larger pyrite nodules of Archean age (Fischer et al., 2014; Marin-Carbonne et al., 2014), requires local heterogeneities in $\Delta^{33}$S of pore fluids, changes in S-MIF production on the time scale of nodule growth, or an as yet unconstrained biological effect.

The model proposed by Farquhar et al. (2013) suggests two distinct pathways for pyrite formation that depend upon sulfate availability, and that the relative contribution of these pathways determines the final $\Delta^{33}$S of pyrite: when the ‘direct’ reaction of relatively abundant and soluble sulfate (e.g., from an atmospheric exit channel as $\text{H}_2\text{SO}_4$ with
negative $\Delta^{32}S$) with sulfide dominates, pyrite with low $\Delta^{32}S$ is produced. When the ‘indirect’ reaction of insoluble elemental sulfur (e.g., from an atmospheric exit channel as $S_8$ with positive $\Delta^{32}S$) with sulfide dominates, soluble polysulfide intermediates with high $\Delta^{33}S$ are produced and impart this signature to later formed pyrite. We further suggest here the logical extension of this argument, that changing relative expression of these two pathways in closed system conditions could result in a positive gradient in $\Delta^{33}S$ within a growing grain or nodule as a local reservoir of low $\Delta^{33}S$ sulfate is depleted and the relative dominance of high $\Delta^{33}S$ polysulfide intermediates in pyrite formation increases.

In the ‘transition’ sample, taken from 148.37 m in the ABDP-9 core, $\delta^{34}S$ varies from −7.7 to 4.8‰, $\Delta^{33}S$ from −3.9 to 6.1‰, and $\Delta^{36}S$ from −3.1 to 4.5‰. In the youngest sample (111.22 m), $\delta^{34}S$ is universally negative, varying from −11.8 to −2.7‰, whereas $\Delta^{32}S$ is positive from 1.7 to 4.3‰, and $\Delta^{36}S$ varies from −3.5 to 0.6‰ (Table 4, Figs 9A, 10A). Decreasing values of $\delta^{34}S$ in these three samples of the ABDP-9 core, as well as decreasing trends in $\Delta^{32}S$, are consistent with a secular increase in seawater sulfate during deposition of the Mount McRae Shale, perhaps due to oxidative continental weathering as previously proposed by Kaufman et al. (2007). Pyrite grains from the two younger samples of the Mount McRae Shale, from 148.37 and 111.22 m in the ABDP-9 core, have $\Delta^{36}S/\Delta^{32}S_{Ago}$ values that are significantly lower than those in the oldest sample (avg. = 0.7 vs. 1.5; $P < 0.02$ in a two-tailed Student’s t-test), but the populations from the two younger samples are statistically indistinguishable from each other (supplementary data). This is consistent with a transition in the expression of $\Delta^{36}S/\Delta^{32}S$ at 153 m depth in the core (Kaufman et al., 2007). Indeed, $\Delta^{36}S/\Delta^{32}S$ relations for individual pyrite grains in all three samples of the Mount McRae Shale analyzed in this study are strongly linear ($\bar{r}^2$ from 0.89 to 0.99), and slopes decrease from −0.88 (161.02 m) to −1.07 (148.37 m) to −1.26 (111.22 m).

Finally, two distinct populations of pyrite with distinct sulfur isotope systematics were observed in ABDP-9 148.37 (Fig. S8). Some pyrite targets (e.g., ABDP-9 148.37a targets 11 and 14) are distinguished by their nodular occurrence in intimate association with a rim of chlorite and sometimes calcite. Other pyrite in the sample is not associated with chlorite, and commonly has rounded and/or broken margins, consistent with a detrital origin. Chlorite-associated pyrite has higher $\delta^{34}S$ and $\Delta^{32}S$ (>0‰) than non-chlorite-associated pyrite. The original sulfur source would have left the atmosphere as zero valent, elemental sulfur, requiring multiple metabolic pathways to pyrite formation including disproportionation and sulfate reduction.

CONCLUSIONS
Spatially resolved, in situ carbon and sulfur isotope analysis of organic matter, carbonate minerals, and pyrite, supported by light and electron microscopy as well as elemental mapping deepen our understanding of Neoarchean biogeochemistry. The approach described here offers an unprecedented combination of spatial resolution, precision and accuracy for carbon and sulfur isotope analysis of micrometer-scale domains of sedimentary organic matter, carbonate minerals, and pyrite, and maximizes potential information recovery in sample limited scenarios (such as scientific drilling or Mars sample return, e.g.). Although SIMS analysis yields extraordinary information density when done correctly, it is labor intensive relative to conventional, or ‘bulk’ analysis. For this reason, and also because of the inherent sampling biases that can be associated with resolution-limited spatially resolved techniques, a combination of in situ and bulk analysis (when sample sizes permit) is the best approach to maximize information recovery. As was the case for this study, bulk analysis is particularly useful to guide the selection of samples to analyze in greater detail. Key findings are as follows:

1 A new method for the simultaneous collection of $^{13}$CH/$^{12}$C data, along with a new set of organic standards that can be used to correct for the H/C effect is described. Based upon the data we report here and the earlier reports from another laboratory, we conclude that a failure to account for this effect can lead to inaccuracies of up to 4‰.

2 This new method for SIMS $\delta^{13}C$ analysis of organic matter affords reproducibilities of 0.3‰ routinely (2SD; comparable to bulk analysis by continuous flow techniques) with a 6-µm-diameter beam and 0.8‰ (2SD) with a 1- to 3-µm-diameter beam. The sample volume consumed by each SIMS $\delta^{13}C$ analysis with a 6-µm-diameter beam (~24 mm$^2$) is $10^{-8}$ that typically required for conventional analysis by continuous flow techniques.

3 We have demonstrated $\delta^{13}C$ measurements with sub-perm error at count rates as low as 30% of the bracketing standard (i.e., >50% carbonate) due in part to the low conversion of inorganic carbon to $C^-$ anions, relative to organic carbon. Conversely, SIMS $\delta^{13}C$ analyses of sedimentary carbonates are strongly affected by small organic inclusions although correlations between $^{12}$C count rate and $\delta^{13}C$ can be used to estimate $\delta^{13}C$ of carbonates in a given sample.

4 The remarkable consistency of averaged SIMS data and earlier bulk data from other laboratories bolsters earlier interpretations of biogeochemical changes observed in Neoarchean drill core material from Australia, including evidence for biotic and abiotic oxidative processes at least 100 Ma prior to the Great Oxidation Event. Fur-
thermore, this consistency lends confidence to the fidelity of the analytical techniques reported here.

5 In situ textural and organic carbon isotope evidence for multiple microbial metabolisms, including phototrophy and methanotrophy, is preserved at the millimeter scale in a single sample of the ~2.7 Ga Tumbiana Fm.

6 Pyrobitumen nodules in samples of the Jeerinah Fm and Mount McRae Shale are enriched in $^{13}C$ relative to co-occurring kerogen, and associations with uraniferous minerals suggest that this may be due to radiolytic alteration.

7 Coupled, in situ carbon and sulfur isotope analysis of organic matter and pyrite in the ~2.7 Ga Tumbiana Fm are consistent with earlier reports of correlations between low $\delta^{13}C$ organic matter and a dampened S-MIF signal, possibly driven by the temporary accumulation of an organic haze. In the ~2.6 Ga Jeerinah Fm and Carawine Dolomite, the range of S-MIF expanded considerably, but anomalous behavior of the minor sulfur isotopes ($^{33}S$ and $^{36}S$) at the scale of individual pyrite grains is associated with extremely low $\delta^{13}C$ of organic matter in intimate association with the pyrite. These strong ties between carbon and sulfur cycling demonstrated for the first time in situ, at the microbial scale, likely continued through the entire Neoarchean.

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REFERENCES


**SUPPORTING INFORMATION**

Additional Supporting Information may be found in the online version of this article:

Data S1. Supplementary materials.