1 Introduction

Extensive hydrothermal chert vein systems containing abundant organic carbon are a unique phenomenon of early Archaean successions worldwide (Lindsay et al., 2005; Van Kranendonk, 2006; Hofmann, 2011). A dense stockwork of several hundred kerogen-rich hydrothermal chert veins that penetrate footwall pillow komatiitic basalts of the ca. 3.5 Ga Dresser Formation (Pilbara Craton, Western Australia; Fig. 1) are up to 2 km deep by 25 m wide (Hickman, 1973, 1983; Nijman et al., 1999; Van Kranendonk and Pirajno, 2004; Lindsay et al., 2005; Van Kranendonk, 2006; Van Kranendonk et al., 2008) (Fig. S1 in the Supplement). Depleted stable carbon isotope signatures (δ13C) of bulk kerogens (−38.1 to −24.3 ‰) and of organic microstructures (−33.6 to −25.7 ‰) in these hydrothermal chert veins, as well as remnants of what appear to be microbial remains, are consistent with a biological origin of the organic matter (Ueno et al., 2001, 2004; Glikson et al., 2008; Pinti et al., 2009; Morag et al., 2016). Problematically, however, simi-
larly depleted δ13C values (partly down to ca. −36‰ relative to the initial substrate) can also be formed through abiotic processes, for instance via Fischer–Tropsch-type synthesis (McCollom et al., 1999; McCollom and Seewald, 2006), and putative microbial remains are not always reliable (Schopf, 1993; Brasier et al., 2002, 2005; Schopf et al., 2002; Bower et al., 2016).

Organic biomarkers can help to trace life and biological processes through deep time and add important information on the origin of the organic matter, even in very old sedimentary rocks (Brocks and Summons, 2003; Summons and Hallmann, 2014). In Archaean rocks, however, molecular fingerprints in the conventionally analysed bitumen (i.e. the extractable portion of organic matter) are commonly blurred by thermal maturation and/or ancient or modern contamination (Brocks et al., 2008; Gérard et al., 2009; Brocks, 2011; French et al., 2015). In contrast, the non-extractable portion of organic matter, known as kerogen, tends to be less affected by thermal maturation and contamination and is considered to be syngentic with the host rock (Love et al., 1995; Brooks et al., 2003b; Marshall et al., 2007; Lockhart et al., 2008).

Catalytic hydropyrolysis (HyPy) is a powerful tool for sensitively releasing kerogen-bound hydrocarbon moieties with little structural alteration (Love et al., 1995). HyPy has been successfully applied to Archaean kerogens in rocks from the Pilbara Craton, liberating syngenetic organic compounds consistent with a biogenic origin (Brocks et al., 2003b; Marshall et al., 2007). However, this technique has not yet been used on kerogens contained in hydrothermal chert veins of this age.

Here, we present the results of analyses of kerogen embedded in a freshly exposed hydrothermal chert vein of the ca. 3.5 Ga Dresser Formation. Our analyses include field and petrographic observations, Raman spectroscopy and organic geochemistry (δ13C: kerogen-bound molecules via HyPy followed by gas chromatography – mass spectrometry (GC-MS) and gas chromatography – combustion – isotope ratio mass spectrometry (GC-C-IRMS)). To further constrain potential sources of the Dresser kerogen, we additionally applied HyPy on (i) excessively pre-extracted cyanobacterial biomass and (ii) produced abiogenic organic matter via Fischer–Tropsch-type synthesis using a hydrothermal reactor. Results of these investigations suggest that the Dresser kerogen has a primarily microbial origin. We hypothesize that biomass-derived organic compounds accumulated in anoxic aquatic environments and were then redistributed and sequestered by subsurface hydrothermal fluids (“hydrothermal pump hypothesis”).

2 Material and methods

2.1 Sample preparation

A fresh decimetre-sized sample of a Dresser chert vein was obtained from a recent cut wall of the abandoned Dresser Mine in the Pilbara Craton, Western Australia (GPS: 21°09′04.13″; S; 119°26′15.21″ E; Figs. 1, S1d; for geological maps, see Hickman, 1983; Van Kranendonk, 1999; Hickman and Van Kranendonk, 2012). The external surfaces (ca. 1–2 cm) of the sample block were removed using an acetone-cleaned rock saw and then used for the preparation of thin sections. The surfaces of the resulting inner block were extensively rinsed with acetone and then removed (ca. 1–2 cm) with an acetone-cleaned high precision saw (Buehler; Isomet 1000, Germany). The surfaces of the resulting sample were again rinsed with acetone and then crushed and powdered using a carefully acetone-cleaned pebble mill (Retsch MM 301, Germany).

2.2 Petrography and Raman spectroscopy

Petrographic analysis was conducted using a Zeiss SteREO Discovery.V8 stereomicroscope (transmitted and reflected light) linked to an AxioCam MRc5 5-megapixel camera.

Raman spectra were recorded using a Horiba Jobin Yvon LabRam-HR 800 UV spectrometer (focal length of 800 mm) attached to an Olympus BX41 microscope. For excitation an Argon ion laser (Melles Griot IMA 106020B0S) with a laser strength of 20 mW was used. The laser beam was focused onto the sample using an Olympus MPlane 100× objective with a numerical aperture of 0.9 and dispersed by a 600 L mm−1 grating on a charge-couple device (CCD) detector with 1024 × 256 pixels. This yielded a spectral resolution of < 2 cm−1 per pixel. Data were acquired over 10 to 30 s for a spectral range of 100–4000 cm−1. The spectrometer was calibrated by using a silicon standard with a major peak at 520.4 cm−1. All spectra were recorded and processed using the LabSpec™ database (version 5.19.17; Jobin Yvon, Villeneuve d’Ascq, France).

2.3 Raman-derived H / C data

H/C values were calculated based on integrated peak intensities of Raman spectra using the formula \[ \text{H/C} = 0.871 \cdot \frac{I_{DS}}{I_{G+D2}} - 0.0508 \] (Ferralis et al., 2016). The peaks were fitted in the LabSpec™ software (see Sect. 2.2) using the Gauss/Lorentz function.

2.4 Molecular analysis of the Dresser kerogen

All materials used for preparation were heated to 500°C for 3 h and/or extensively rinsed with acetone prior to sample contact. A laboratory blank was prepared and analysed in parallel to monitor laboratory contaminations.
We applied catalytic hydropyrolysis (HyPy) to release molecules from the Dresser kerogen and pre-extracted biomass of the heterocystous cyanobacterium *Anabaena cylindrica* SAG 1403-2 following previously published protocols (Snape et al., 1989; Love et al., 1995, 2005). HyPy allows the breaking of covalent bonds by progressive heating under high hydrogen pressure (150 bar). The released products are immediately removed from the hot zone by a constant hydrogen flow and trapped downstream on clean, combusted silica powder cooled with dry ice (Meredith et al., 2004). All hydropyrolyses were eluted from the silica trap with high-purity dichloromethane (DCM), desulfurized overnight using activated copper and subjected to gas chromatography–mass spectrometry (GC-MS). Before all experiments, the empty HyPy system was heated (ambient temperature to 520 °C, held for 30 min) to remove any residual molecules.

Isolation of the Dresser kerogen followed standard procedures (cf. Brocks et al., 2003b). 57 g of powdered sample was first demineralized with hydrochloric acid (24 h) and then hydrofluoric acid (11 days). The purified organic matter was then exhaustively extracted using three excess volumes of DCM and n-hexane, respectively (×3), ultrasonic swelling in pyridine (2 × 20 min at 80 °C), and ultrasonic extraction with methanol, DCM (×3) and DCM / methanol (1/1; v/v). The kerogen was subsequently extracted with n-hexane until no more compounds were detected via GC-MS. The pure kerogen was used for HyPy following an experimental protocol optimized for the analysis of Archaean kerogens (Brocks et al., 2003b; Marshall et al., 2007).

In order to monitor potential contamination, we applied HyPy to blanks before and after the kerogen run. The Dresser kerogen (131.06 mg) and blanks were sequentially heated in the presence of a sulfided molybdenum catalyst (10 wt %) and under a constant hydrogen flow (5 dm³ min⁻¹) using a two-step approach (cf. Brocks et al., 2003b; Marshall et al., 2007; Fig. S2). The low-temperature step included heating from ambient temperature to 250°C (300°C min⁻¹) and then to 330°C (8°C min⁻¹) to release any molecules that were strongly adsorbed to the kerogen and not accessible to solvent extraction. The silica powder was subsequently recovered from the product trap, and the trap was refilled with clean, combusted silica powder for the following high-temperature step. This step included heating from ambient temperature to 520°C (8°C min⁻¹) to release molecules covalently bound to the kerogen.

### 2.5 Molecular analysis of pre-extracted cyanobacterial biomass (*Anabaena cylindrica*)

We used cell material of the heterocystous cyanobacterium *Anabaena cylindrica* strain SAG 1403-2 as it fulfils all criteria for reference material (availability, well characterized, etc.). The material was obtained from the Culture Collection of Algae at the Georg-August-Universität Göttingen (Germany) and grown at the Christian-Albrechts-Universität Kiel (Germany). The axenic batch culture was maintained in 250 mL of BG11 medium free of combined nitrogen sources (Rippka et al., 1979) at 29 °C. A light : dark regime of 14 h : 10 h with a photon flux density of 135 µmol photons m⁻² s⁻¹ was provided by a white fluorescent light bulb. At the end of the logarithmic growth phase, cells were harvested by centrifugation, and they were lyophilized thereafter. Subsequently, an aliquot of the freeze-dried biomass was exhaustively extracted following the methodology described by Bauersachs et al. (2014).

HyPy was performed following an experimental protocol optimized for biomass applications (Love et al., 2005). Briefly, the material was heated in the presence of a sulfided molybdenum catalyst (1.5 times the weight of the bacterial extraction residue) and under a constant hydrogen flow (6 dm³ min⁻¹). The HyPy program included heating from ambient temperature to 260°C (300°C min⁻¹) and then to 500°C (8°C min⁻¹).

### 2.6 Fischer–Tropsch-type synthesis of organic matter under hydrothermal conditions

Fischer–Tropsch-type reactions were carried out based on McCollom et al. (1999). A mixture of 2.5 g oxalic acid (dihydrate, suprapur®), Merck KGaA), 1 g montmorillonite (K10, Sigma Aldrich; pre-extracted with DCM; ×4) and ca. 11 mL ultrapure water was heated to 175°C for 66 to 74 h in a sealed Morey-type stainless steel autoclave. The autoclave...
was rapidly (≤ 15 min) cooled to room temperature with compressed air.

Fluid and solid phases were collected and extracted with DCM (×4). The montmorillonite was removed by centrifugation and filtration with silica powder and sea sand (which were combusted prior to use). The obtained extracts were then concentrated by rotary evaporation (40 °C, 670 mbar) and subjected to GC-MS analysis. Analytical blank experiments were carried out with all reactants to keep track of contamination.

2.7 Gas chromatography – mass spectrometry

GC-MS analysis of HyPy products was carried out with a Thermo Scientific Trace 1310 GC coupled to a Thermo Scientific Quantum XLS Ultra MS. The GC instrument was equipped with a capillary column (Phenomenex Zebon ZB-5, 30 m, 0.25 µm film thickness, 0.25 mm inner diameter). Fractions were injected into a splitless injector and transferred to the GC column at 270 °C. He was used as carrier gas with a constant flow rate of 1.5 mL min⁻¹. The GC oven temperature was held isothermal at 80 °C for 1 min and then ramped to 310 °C at 5 °C min⁻¹, at which it was kept for 20 min. Electron ionization mass spectra were recorded in full-scan mode at an electron energy of 70 eV with a mass range of m/z 50–600 and scan time of 0.42 s.

2.8 Polyaromatic hydrocarbon ratios

Polyaromatic hydrocarbons (PAHs) are organic compounds consisting of multiple fused benzene rings. Methylation and isomerization characteristics of various GC-amenable PAHs are altered during maturation and thus provide a measure for assessing the thermal maturity of organic matter (Killops and Peters et al., 2005). The following PAH maturity parameters were used in this study:

1. methylphenanthrene ratio (MNR) = 2-MN / 1-MN (Radke et al., 1984),
2. methylphenanthrene index (MPI-I) = 1.5 · (2-MP + 3-MP) / (P + 1-MP + 9-MP) (Radke and Welte, 1983), and
3. computed vitrinite reflectance [Rv (MPI-I)] = 0.7 · MPI-I + 0.22 (according to P / MP > 1; Boreham et al., 1988).

MN stands for methylphenanthrene, P for phenanthrene and MP for methylphenanthrene.

2.9 Total organic carbon (TOC) and δ¹³C analyses (TOC and compound specific)

TOC, δ¹³C_TOC and compound-specific δ¹³C analyses were conducted at the Centre for Stable Isotope Research and Analysis (KOSI) at the Georg-August-Universität Göttingen, Germany. Stable carbon isotope data are expressed as delta values relative to the Vienna Pee Dee Belemnite (VPDB) reference standard.

The TOC content and δ¹³C_TOC values were determined in duplicate using an elemental analyser (NA-2500 CE-Instruments) coupled to an isotope ratio mass spectrometer (Finnigan MAT Delta plus). Ca. 100 mg of powdered and homogenized whole rock material were analysed in each run. For internal calibration an acetanilide standard was used (δ¹³C = −29.6 ‰; SD = 0.1 ‰). TOC content measurements showed a mean deviation of 0.1 wt %. The average δ¹³C_TOC value had a standard deviation of 0.3 ‰.

Compound-specific δ¹³C analyses were conducted with a Trace GC coupled to a Delta Plus isotope ratio mass spectrometer (IRMS) via a combustion interface (all Thermo Scientific). The combustion reactor contained CuO, Ni and Pt and was operated at 940 °C. The GC was equipped with two serially coupled capillary columns (Agilent DB-5 and DB-1; each 30 m, 0.25 µm film thickness, 0.25 mm inner diameter). Fractions were injected into a splitless injector and transferred to the GC column at 290 °C. The carrier gas was He at a flow rate of 1.2 mL min⁻¹. The GC oven temperature program was identical to the one used for GC-MS analysis (see above). CO₂ with a known δ¹³C value was used for internal calibration. Instrument precision was checked using laboratory standards. Standard deviations of duplicate measurements were better than 1.7 ‰.

3 Results

The studied hydrothermal chert vein is hosted in komatiitic pillow basalt that has undergone severe hydrothermal acid–sulfate alteration, producing a kaolinite–illite–quartz mineral assemblage (Van Kranendonk and Pirajno, 2004; Van Kranendonk, 2006) (Fig. S1). The sampled vein crops out in a recent cut wall of the abandoned Dresser Mine (Fig. S1) and consists of a dense chert (microquartz) matrix of deep black colour that contains kerogen and local concentrations of fresh (unweathered) pyrite crystals (Fig. 2a, b). There is no field and/or petrographic evidence for fluid-flow events that postdate the initial vein formation (brecciation textures, etc.).

Petrographic analysis and Raman spectroscopy revealed that kerogen (D bands at 1353 cm⁻¹, G bands at 1602 cm⁻¹) is embedded in the chert matrix (SiO₂ band at 464 cm⁻¹) (see Fig. 2c, for a representative Raman spectrum). The organic matter occurs as small clots (< 50 µm) of variable shape. Raman spectra of the organic matter exhibit relatively wide D and G bands (82 and 55 cm⁻¹ width, respectively) (Fig. 2c). The total organic carbon (TOC) content is 0.2 wt %, and the δ¹³C_TOC value is −32.8 ± 0.3 ‰ (Table 1). Raman-derived H / C ratios range between 0.03 and 0.14.

HyPy was applied to the isolated Dresser kerogen, as well as to preceding and subsequent analytical blanks (combusted sea sand). Hydropyrolysates of the preceding blank con-


Table 1. Stable carbon isotopic composition (δ\(^{13}\)C) of the total organic carbon (TOC) and n-alkanes released from high-temperature HyPy of the Dresser kerogen.

<table>
<thead>
<tr>
<th>n-C(_{12})</th>
<th>n-C(_{13})</th>
<th>n-C(_{14})</th>
<th>n-C(_{15})</th>
<th>n-C(_{16})</th>
<th>n-C(_{17})</th>
<th>n-C(_{18})</th>
<th>n-C(_{12-18}) (mean)</th>
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<td>-32.7</td>
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<td>-29.4</td>
<td>-31.2</td>
<td>-31.7</td>
<td>-31.4</td>
</tr>
<tr>
<td>SD</td>
<td>0.5</td>
<td>1.4</td>
<td>0.2</td>
<td>1.7</td>
<td>0.3</td>
<td>0.8</td>
<td>0.1</td>
<td>1.2</td>
</tr>
</tbody>
</table>

Table 2. Maturity indices (based on aromatic hydrocarbons) of the Dresser kerogen. MP/P: methylpheneanthrene/phenanthrene ratio; MPI-I: methylphenanthrene index (1.5·(2-MP + 3-MP)/(P + 1-MP + 9-MP); Radke and Welte, 1983); P/MP: phenanthrene/methylphenanthrene ratio; \( R_c \) (MPI-I): computed vitrinite reflectance (0.7·MPI-I + 0.22, according to P/MP > 1; Boreham et al., 1988); MNR: methylnaphthalene ratio (2-MN/1-MN; Radke et al., 1984).

<table>
<thead>
<tr>
<th>MP/P</th>
<th>MPI-I</th>
<th>P/MP</th>
<th>( R_c ) (MPI-I)</th>
<th>MNR</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.33</td>
<td>0.23</td>
<td>3.01</td>
<td>2.87</td>
<td>2.52</td>
</tr>
</tbody>
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4 Discussion

4.1 Maturity of the Dresser kerogen

The organic record of Archaean rocks is commonly affected by thermal maturation (Brocks et al., 2008; Brocks, 2011; French et al., 2015). The kerogen within the analysed Dresser sample is thermally mature and structurally disordered, as evidenced by relatively wide D and G bands in the Raman spectra (see Fig. 2c for a representative Raman spectrum). This is also supported by the absence of S1 bands at 2450 cm\(^{-1}\), high R1 ratios (0.98–1.05) and low FWHM-D1 values (68.64–76.12), consistent with prehnite–pumpellylite to lower greenschist metamorphism at a temperature of ca. 300 °C (cf. Yui et al., 1996; Delarue et al., 2016). All of these observations are well in line with published Raman spectra of Archaean organic matter from the same region (Ueno et al., 2001; Marshall et al., 2007; Delarue et al., 2016) and the general thermal history of the host rock (regional prehnite–pumpellylite to lower greenschist metamorphism; Hickman, 1975, 1983, 2012; Terabayashi et al., 2003).

Fitting of D5 peaks can be difficult for Raman spectra of highly mature organic matter (Ferralis et al., 2016). Therefore, the H/C ratios calculated herein (0.03–0.14) should be treated with caution. However, the Raman-based H/C ratios and the methylphenanthrene/phenanthrene value (MP/P = 0.33; Table 2) of the Dresser kerogen are in good accordance with data reported from more mature kerogens from the same region (ca. 3.4 Ga Strelley Pool Formation: 0.08–0.14 and 0.24–0.37, respectively; Marshall et al., 2007). The low methylphenanthrene index (MPI-I = 0.23) and high phenanthrene/methylphenanthrene index (P/MP = 3.01) result in a computed vitrinite reflectance (\( R_c \) (MPI-I)) of 2.87 (Table 2), indicating a thermal maturity far beyond the oil-generative stage (Radke and Welte, 1983; Boreham et al., 1988). However, it has to be considered that the MPI-I is potentially affected by (de-)methylation reac-
Figure 2. Petrographic observations on the hydrothermal chert vein. (a, b) Thin section photographs (a transmitted light; b reflected light) showing kerogen (brownish colours) and pyrite (arrows, black colours in a, bright colours in b) embedded within a fine-grained chert matrix. Note that the pyrite crystals (arrows) are excellently preserved and show no evidence of oxidation. (c) Representative Raman spectrum of kerogen (D band at 1353 cm$^{-1}$ and G band at 1602 cm$^{-1}$) present in the hydrothermal chert vein. Note the wide D and G bands (82 and 55 cm$^{-1}$ width, respectively), pointing to thermally mature and structurally disordered kerogen, and the absence of an S1 band (at 2450 cm$^{-1}$), consistent with prehnite–pumpellyite to lower greenschist metamorphism and a temperature of ca. 300°C (cf. Yui et al., 1996; Delarue et al., 2016).

The distribution of monomethylalkanes ≤ n-C$_{18}$ (Figs. S3, S4) released from the Dresser kerogen resembles high-temperature HyPy products from the Strelley Pool kerogen (Marshall et al., 2007; their Fig. 15). Such isomeric mixtures are typically formed during thermal cracking of alkyl moieties (Kissin, 1987) and are in good agreement with the estimated maturity range. Methylated aromatics such as methyl-naphthalenes and methylphenanthrenes have also been observed in other hydropyrolysates from Archaean kerogens that experienced low-grade metamorphism (Brocks et al., 2003b; Marshall et al., 2007; French et al., 2015). In all of these cases, the degree of alkylation varied with the exact thermal alteration of the respective kerogens (Marshall et al., 2007; French et al., 2015).

4.2 Syngeneity of the Dresser kerogen-derived compounds

The kerogen of the Dresser Formation exclusively occurs in the form of fluffy aggregates and clots embedded within a very dense chert matrix that is, once solidified, highly impermeable to fluids. The depositional age of the formation is constrained to 3481 ± 3.6 Ma (Van Kranendonk et al., 2008), and the investigated chert vein shows no evidence for disruption by post-depositional hydrothermal fluids. This has also been described for other hydrothermal chert veins of the Dresser Formation, where the kerogen has been interpreted as being syngenetic (i.e. formed prior to host rock lithification; Ueno et al., 2001, 2004; Morag et al., 2016). Furthermore, the maturity of the embedded kerogen is in good accordance with the thermal history of the host rock. An introduction of solid macromolecular organic matter from stratigraphically younger units in this region during later fluid-flow phases, as proposed for the younger Apex chert (Olcott-Marshall et al., 2014), can therefore be excluded.

As the bitumen fractions of Precambrian rocks are easily biased by the incorporation of contaminants during later
Figure 3. Total ion currents (a–c; on the same scale) and ion chromatograms selective for alkanes (d–e; m/z 85). (a) High-temperature HyPy (330–520°C) product of analytical blank (combusted sea sand) obtained prior to HyPy of the Dresser kerogen. n-Alkanes in the range of n-C_{14} to n-C_{24} (maxima at n-C_{15}) represent HyPy background contamination. (b) High-temperature HyPy (330–520°C) product of the Dresser kerogen. Note the sharp decrease in abundance of n-alkanes beyond n-C_{18} (see arrows). δ^{13}C values of n-C_{12} to n-C_{28} show a high similarity to the δ^{13}C_{TOC} value (−32.8 ± 0.3 ‰), further confirming syngeneity. (c) Blank subtraction (b minus a) showing that contaminants have no major impact on the n-alkane pattern yielded during the high-temperature HyPy step of the Dresser kerogen (b). (d) HyPy products of cell material of the heterocystous cyanobacterium Anabaena cylindrica. Note the sharp decrease in abundance of n-alkanes beyond n-C_{18} (see arrows), similar to the n-alkane distribution of the Dresser kerogen (b). (e) Products of experimental Fischer–Tropsch-type synthesis under hydrothermal conditions; abiogenic n-alkanes show a unimodal distribution that is distinctly different from the Dresser kerogen. Black dots: n-alkanes (numbers refer to carbon chain lengths); N: naphthalene; MN: methylnaphthalenes; BiPh: 1,1′-biphenyl; DMN: dimethylnaphthalenes; DBF: dibenzofuran; MAN: methylacenaphthene; P: phenanthrene; crosses: siloxanes (GC column or septum bleeding).
stages of rock history (Brocks et al., 2008; Brocks, 2011; French et al., 2015), studies have increasingly focussed on kerogen-bound compounds that are more likely to be syngenetic to the host rock (Love et al., 1995; Brocks et al., 2003b; Marshall et al., 2007; French et al., 2015). Potential volatile organic contaminants adhering to the kerogen are removed through excessive extraction and a thermal desorption step (\(\sim 350 \, ^\circ\text{C}\)) prior to high-temperature HyPy (550 \(^\circ\text{C}\); cf. Brocks et al., 2003b; Marshall et al., 2007). The recurrence of few \(n\)-alkanes in the range of \(n\)-C14 to \(n\)-C24 (maximum at \(n\)-C15) in high-temperature HyPy blank runs obtained immediately before and after the actual sample run (Figs. 3a, S2, S3) indicates minor background contamination during pyrolysis, with a source most likely within the HyPy system. However, these contaminants do not significantly affect the \(n\)-alkane pattern yielded by high-temperature HyPy of the Dresser kerogen, as evidenced by blank subtraction (Fig. 3c).

Contamination of the sample can be further deciphered by the presence of polar additives or hydrocarbons that are not consistent with the thermal history of the host rock. Plastic-derived branched alkanes with quaternary carbon centres (BAQCs), a common contaminant in Precambrian rock samples (Brocks et al., 2008), have not been detected (Fig. S7). Traces of functionalized plasticizers (phenols and phthalic acid; Figs. S2, S6, S8) are unlikely to survive (or result from) HyPy treatment. These compounds are therefore considered as background contamination introduced during sample preparation and analysis after HyPy. The observed siloxanes (Fig. 3, S2) probably originate from the GC column or septum bleeding and are unlikely to be contained in the sample. All of these compounds occur only in low or trace abundances and can be clearly distinguished from the ancient aliphatic and aromatic hydrocarbons contained in the Dresser kerogen.

Contamination by (sub-)recent endoliths can be excluded as sample surfaces have been carefully removed and there is no petrographic indication for borings or fissures containing recent organic material (Fig. 2). HyPy of untreated or extracted biomass would yield a variety of acyclic and cyclic biomarkers (cf. Love et al., 2005). However, high-temperature HyPy of the Dresser kerogen almost exclusively yielded \(n\)-alkanes, minor amounts of monomethylalkanes and various aromatic hydrocarbons (Figs. 3b, c, S2–S6), while hopanoids or steroids were absent (Figs. S8, S9). This is in good agreement with the maturity of the Dresser kerogen and previous HyPy studies of Archaean rocks (Brocks et al., 2003b; Marshall et al., 2007; French et al., 2015).

Accidental contamination of the kerogen by mono-, di- and triglycerides (e.g. dust, skin surface lipids) can also be ruled out as HyPy treatment of these compounds typically results in \(n\)-alkane distributions with a distinct predominance of \(n\)-C16 and \(n\)-C18 homologues, corresponding to the \(n\)-C16 and \(n\)-C18 fatty acid precursors (Craig et al., 2004; Love et al., 2005; unpublished data from our own observations). At the same time, the observed distribution of kerogen-derived \(n\)-alkanes, with a distinct decrease beyond \(n\)-C18 (Fig. 3b, c), is notably similar to high-temperature HyPy products of kerogens isolated from the ca. 3.4 Ga Strelley Pool Formation of the Pilbara Craton (Marshall et al., 2007; their Fig. 14). Marshall and co-workers considered these \(n\)-alkanes unlikely to be contaminants as they were released only in the high-temperature HyPy step. Furthermore, the stable carbon isotopic composition of \(n\)-alkanes in the Dresser high-temperature pyrolysate (\(-29.4 \pm -33.3 \, ^\circ\text{C}\); mean \(-31.4 \pm 1.2 \, ^\circ\text{C}\)) is very similar to the \(\delta^{13}\text{C}_{\text{TOC}}\) signature of the sample (\(-32.8 \pm 0.3 \, ^\circ\text{C}\); Figs. 3, S10; Table 1), indicating that these compounds were generated from the kerogen. Hence, we consider the compounds released from the Dresser kerogen during high-temperature HyPy (Fig. 3b, c) as syngenetic.

High-temperature HyPy of the Dresser kerogen yielded a variety of aromatic hydrocarbons, which are orders of magnitudes lower or absent in all other pyrolysates (Figs. 3, S2, S4–S6). It also produced significantly higher amounts of \(n\)-alkanes than the low-temperature step, and these further showed a distinct distribution pattern (i.e. a step \(\leq n\)-C18; Figs. 3, S2, S3). The lack of even low quantities of \(n\)-alkanes that typically accompany bond cleavage in kerogen pyrolysis was also observed by Marshall et al. (2007) in their HyPy analysis of cherts from the Strelley Pool Formation. As a possible explanation for the lack of \(n\)-alkenes, these authors suggested that the \(n\)-alkanes were not cracked from the kerogen but were rather trapped in closed micropores until the organic host matrix was pyrolytically disrupted. Alternatively, however, unsaturated cleavage products may have been immediately reduced during HyPy by the steadily available hydrogen and catalysts. Indeed, it has been demonstrated that double bonds in linear alkyl chains are efficiently hydrogenated during HyPy, even in the case of pre-extracted microbial biomass (e.g. Love et al., 2005; our Fig. 3d). We consider both as plausible scenarios to explain the absence of \(n\)-alkenes in the Dresser chert hydropyrolysate.

4.3 Origin of the Dresser kerogen: hydrothermal vs. biological origin

The \(\delta^{13}\text{C}_{\text{TOC}}\) value of \(-32.8 \pm 0.3 \, ^\circ\text{C}\) is consistent with carbon fixation by photo- or chemoautotrophs (cf. Schidlowski, 2001). However, organic compounds exhibiting similar \(^{13}\text{C}\) depletions (partly down to ca. \(-36 \, ^\circ\text{C}\) relative to the initial substrate) could also be formed abiotically during the serpenization of ultramafic rocks (Fischer–Tropsch-type synthesis; McCollom et al., 1999; McCollom and Seewald, 2006; Proskurowski et al., 2008). A further constraint on the origin of the Dresser kerogen is provided by the distinct decrease in \(n\)-alkane abundance beyond \(n\)-C18 observed in the high-temperature HyPy pyrolysate (Fig. 3b, c). This distribution resembles HyPy products of pre-extracted recent cyanobac-
terial biomass, which also shows a very pronounced restriction in carbon number to homologues ≤ n-C_{18} (Fig. 3d).

The pre-extracted cyanobacterial cell material and the abiotically produced n-alkanes studied as reference samples experienced no thermal maturation. It can nevertheless be expected that burial, and thus heating, of *Anabaena* biomass (Fig. 3d) would initially liberate lower n-alkane homologues from their predominant n-alkyl moieties while, up to a certain point, retaining the distinct step at n-C_{18}. Experimental maturation of an immature kerogen revealed the preservation of distinct alkyl-chain length preferences even after 100 days at 300°C (Mißbach et al., 2016; e.g. a step beyond n-C_{31} in their Fig. 2). Further maturation would ultimately lead to a unimodal distribution of short-chain n-alkanes and erase the biologically inherited pattern (cf. Mißbach et al., 2016). In contrast, abiotically synthesized extractable organic compounds show a unimodal homologue distribution from the beginning (Fig. 3e) and will retain it, while thermal maturation would gradually shift the n-alkane pattern towards shorter homologues. It can therefore be expected that organic compounds cleaved from an abiotic “Fischer–Tropsch kerogen” – whose existence has not been proven yet – would also exhibit a unimodal distribution. Consequently, the distinctive distribution of n-alkanes released from the Dresser kerogen (i.e. the step ≤ n-C_{18}; Figs. 3b, c) can be regarded as a molecular fingerprint relating to a biosynthetic origin of the organic matter.

Highly 13C-depleted methane in primary fluid inclusions in hydrothermal chert veins of the Dresser Formation (δ^{13}C ≤ 56 ‰) was taken as evidence for biological methanogenesis and thus the presence of Archaea (Ueno et al., 2006). Our δ^{13}C_{TOC} value (−32.8 ± 0.3 ‰; Table 1; Fig. S10) would generally be consistent with both bacterial and archaeal sources (cf. Schidlowski, 2001). Archaea only synthesize isoprene-based compounds (Koga and Morii, 2007; Matsumi et al., 2011). Straight-chain (acytyl-based) hydrocarbon moieties such as fatty acids – the potential precursors of the kerogen-derived n-alkanes – are to current knowledge being formed only by Bacteria and Eukarya, where they typically function as constituents of membranes or storage lipids (cf. Erwin, 1973; Kaneda, 1991). The formation of these lipids is tightly controlled by different biosynthetic pathways resulting in characteristic chain-length distributions. In bacterial lipids, carbon chain lengths typically do not extend above n-C_{18} (cf. Kaneda, 1991). Consequently, given the lack of convincing evidence for the presence of Eukarya as early as 3.5 Ga (cf. Parfrey et al., 2011; Knoll, 2014; French et al., 2015), the most plausible source of kerogen-occluded n-alkanes in the Dresser hydrothermal chert is Bacteria. Strongly reducing conditions during the deposition of the Dresser Formation are indicated by widespread pyrite, Fe-rich carbonates and trace element signatures (Van Kranendonk et al., 2003, 2008). Potential microbial sources for the Dresser kerogen therefore may have included anoxygenic photoautotrophic, chemooautotrophic and heterotrophic microorganisms.

### 4.4 The “hydrothermal pump hypothesis”

Our results strongly support a biological origin of the kerogen found in the early Archaean hydrothermal chert veins of the Dresser Formation. We explain this finding by the redistribution and sequestration of microbial organic matter that may have formed in a variety of Dresser environments through hydrothermal circulation (proposed herein as the “hydrothermal pump hypothesis”; Fig. 4). Higher geothermal gradients prior to the onset of modern-type plate tectonics at ca. 3.2–3.0 Ga (Smithies et al., 2005; Shirey and Richardson, 2011) were possibly important drivers of early Archaean hydrothermal systems. In fact, the Dresser Formation was formed in a volcanic caldera environment affected by strong hydrothermal circulation, where voluminous fluid circulation locally caused intense acid–sulfate alteration of basalts and the formation of a dense hydrothermal vein swarm (Nijman et al., 1999; Van Kranendonk and Pirajno, 2004; Van Kranendonk, 2006; Van Kranendonk et al., 2008; Harris et al., 2009). In addition to the high crustal heat flow, the absence of thick sedimentary cover may have facilitated the intrusion of seawater into the hydrothermal system. The associated large-scale assimilation of particulate and dissolved organic matter and its transport and alteration by hydrothermal fluids (Fig. 4) therefore appears to be a plausible mechanism that may, at least partly, explain the high amounts of kerogen in early Archaean hydrothermal veins.

The hydrothermal pump hypothesis requires a source of organic matter during the deposition of the Dresser Formation (Fig. 4). Whereas contributions from extraterrestrial sources, as well as from Fischer–Tropsch-type synthesis linked to the serpentization of ultramafic rocks, cannot be excluded, our results indicate a primarily biological origin for the kerogen contained in the chert veins (Fig. 4). The inferred biogenicity is also in line with the consistent δ^{13}C offsets between bulk kerogens (ca. −20 to −30 ‰) and carbonate (ca. ±2 ‰) in Archaean rocks (Hayes, 1983; Schidlowski, 2001). Prokaryotic primary producers and heterotrophs may have flourished in microbial mats (Dresser stromatolites; Walter et al., 1980; Van Kranendonk, 2006, 2011; Philippot et al., 2007; Van Kranendonk et al., 2008), the water column (planktic “marine snow”; Brasier et al., 2006; Blake et al., 2010) and even hot springs on land (Djokic et al., 2017). Another biological source for the ancient organic matter could have been chemooautotrophs and heterotrophs thriving in subsurface environments, such as basalts (Banerjee et al., 2007; Funes et al., 2008) and hydrothermal vent systems (Shen et al., 2001; Ueno et al., 2001, 2004, 2006; Pinti et al., 2009; Morag et al., 2016) (Fig. 4). All of these systems are not mutually exclusive and the largely anoxic conditions would have encouraged a high steady-state abundance of organic matter in the aquatic environment (Fig. 4).
Figure 4. The “hydrothermal pump hypothesis”. Organic matter was predominantly biologically produced and heterotrophically processed by Bacteria and, possibly, Archaea. Additionally, Fischer–Tropsch-type synthesis of organic matter linked to the serpentinization of ultramafic rocks (McCollom et al., 1999; McCollom and Seewald, 2006) may have occurred locally. Primary producers (chemoautotrophs, anoxygenic photoautotrophs) and heterotrophs may have flourished in surface waters (planktic “marine snow”), at the water–rock interface (microbial mats and/or biofilms) and in cryptic environments (e.g. within basalts and hydrothermal vent systems). After accumulating in different Dresser environments, the organic matter was redistributed and sequestered in veins by hydrothermal fluids.

Dissolved organic matter (DOM) in modern seawater may resist decomposition over millennial timescales (Druffel and Griffin, 2015). In recent hydrothermal fields, however, organic matter becomes thermally altered and redistributed (Simoneit, 1993; Delacour et al., 2008; Konn et al., 2009). Laboratory experiments using marine DOM indicate that thermal alteration already occurs at temperatures > 68–100 °C and efficient removal of organic molecules at 212–401 °C (Hawkes et al., 2015, 2016). It has been argued, however, that such DOM removal may also be due to transformation into immiscible material through, for example, condensation (Castello et al., 2014) and/or defunctionalization reactions (Hawkes et al., 2016). These processes, however, are as yet poorly understood. In the Dresser Formation, hydrothermal temperatures ranged from ca. 300 °C at depth to 120 °C near the palaeosurface, causing propylitic (ca. 250–350 °C) and argillic (including advanced argillic: ca. 100–200 °C) alteration of the host rocks (Van Kranendonk and Pirajno, 2004; Van Kranendonk et al., 2008; Harris et al., 2009). Given this variety of thermal regimes, and the generally anoxic nature of early Archaean seawater (e.g. Van Kranendonk et al., 2003, 2008; Li et al., 2013), it is likely that some of the organic substances underwent in situ alteration, but not complete oxidation, during hydrothermal circulation. The entrained organics would have been trapped in the chert that instantaneously precipitated from the ascending hydrothermal fluids due to subsurface cooling (cf. Van Kranendonk, 2006).

In summary, the hydrothermal pump hypothesis proposed here (Fig. 4) includes (i) a net build-up of organic matter in different Dresser environments under largely anoxic conditions, (ii) a large-scale assimilation of particulate and dissolved organic matter from various biological sources and its subsurface transport and alteration by hydrothermal fluids, and (iii) its sequestration within hydrothermal chert veins as kerogen. This model explains the presence of abundant organic carbon in early Archaean hydrothermal veins, as well as its morphological, structural and isotopic variability observed in the Dresser hydrothermal chert veins (Ueno et al., 2001, 2004; Pinti et al., 2009; Morag et al., 2016). It does not, however, help to pinpoint the formation pathways of distinct carbonaceous structures, as for instance those preserved in the Dresser Formation (Glikson et al., 2008) or the younger Apex chert (e.g. Schopf, 1993; Brasier et al., 2002, 2005; Schopf et al., 2002; Bower et al., 2016). Further work is necessary to test whether consistent molecular and compound-specific isotopic patterns can be generated from a larger set of Archaean kerogens.

5 Conclusions

Kerogen embedded in a hydrothermal chert vein from the ca. 3.5 Ga Dresser Formation (Pilbara Craton, Western Australia) is syngenetic. A biological origin is inferred from the presence of short-chain n-alkanes in high-temperature HyPy pyrolysates, showing a sharp decrease in homologue abundance beyond n-C18. HyPy products of pre-extracted recent bacterial biomass exhibited a similar restriction to carbon chain lengths ≤ n-C18, whereas abiotic compounds experimentally formed via Fischer–Tropsch-type synthesis exhibited a unimodal distribution. A biological interpretation for Dresser organics is further consistent with the δ13C_TOC value (−32.8 ± 0.3 ‰) and the stable carbon isotopic composition of n-alkanes in the Dresser high-temperature pyrolysate (−29.4 to −33.3 ‰; mean −31.4 ± 1.2 ‰). Based on these observations, we propose that the original organic matter was primarily biologically synthesized. We hypothesize that microbially derived organic matter accumulating in anoxic aquatic (surface and/or subsurface) environments was
assimilated, redistributed and sequestered by hydrothermal fluids (“hydrothermal pump hypothesis”).

Data availability. Data are available upon request.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/bg-15-1535-2018-supplement.

Author contributions. JPD, JR and MJVK conducted the field work and designed the study. JR conducted petrographic analyses. NS performed Raman spectroscopic analyses. MR and JPD conducted pyrolysis experiments. HM conducted Fischer–Tropsch-type synthesis. TB prepared cyanobacterial cell material. JPD, HM, MR and VT performed biomarker analyses. JPD wrote the manuscript. All authors discussed the results and provided input to the manuscript.

Competing interests. The authors declare that they have no conflict of interest.

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