Palaeoceanographic controls on spatial redox distribution over the Yangtze Platform during the Ediacaran–Cambrian transition

LAWRENCE M. OCH*, LORENZO CREMONESE*, GRAHAM A. SHIELDZ-ZHOU*†, SIMON W. POULTON‡, ULRICH STRUCK§, HONGFEI LING¶, DA LI¶, XI CHEN¶, CHRISTINA MANNING**, MATTHEW THIRLWALL**, HARALD STRAUSS†† and MAOYAN ZHU†

*Department of Earth Sciences, University College London, Gower Street, London WC1E 6BT, UK (E-mail: lawrence.och@gmail.com)
†State Key Laboratory of Palaeobiology and Stratigraphy, Nanjing Institute of Geology and Palaeontology, Chinese Academy of Sciences, Nanjing 210008, China
‡School of Earth and Environment, University of Leeds, Leeds LS2 9JT, UK
§Leibniz-Institut für Evolutions- und Biodiversitätsforschung, Museum für Naturkunde, 10115 Berlin, Germany
¶State Key Laboratory for Mineral Deposits Research, Department of Earth Sciences, Nanjing University, Nanjing 210093, China
**Department of Earth Sciences, Royal Holloway, University of London, Egham, Surrey TW20 0EX, UK
††Institute for Geology and Paleontology, Westfälische Wilhelms-Universität Münster, 48149 Münster, Germany

ABSTRACT

The Ediacaran–Cambrian interval was an eventful transitional period, when dynamic interactions between the biosphere and its physical environment allowed the Earth System to cross into a new state, characterized by the presence of metazoans, more equable climates and more expansive oxygenation of the oceans. Due to the retreat of widespread sulphidic conditions, redox-sensitive trace-metals could accumulate to a greater extent in ‘black shales’ deposited in localized anoxic/euxinic environments, such as highly productive ocean margins. This study investigates the concentrations of the redox-sensitive trace-metals molybdenum and vanadium in organic-rich sedimentary rocks from seven sections of the Yangtze Platform, slope and basin. Iron speciation analyses were carried out in order to distinguish oxic, anoxic-ferruginous and anoxic-sulphidic settings, while sulphur and nitrogen isotope ratios were measured to gain insight into sulphate and nitrate availability, respectively, in the context of changing redox conditions. The data herein demonstrate an overall increase in redox-sensitive trace-metal contents in black shales across the Ediacaran–Cambrian transition, but with marked temporal and spatial variability. Euxinia is evident in South China before 551 Ma in the Ediacaran, and again in the early Cambrian. However, some time-equivalent sections are not enriched in redox-sensitive trace-metals, and also exhibit contrasting S-isotope and N-isotope systematics. A more complex configuration of the Yangtze Platform, for example with vast intrashelf basins, together with changing (generally rising) eustatic sea-level may account for this variability. In this regard, it is proposed that a mid-depth sulphidic wedge, caused by nutrient upwelling over the south-east platform...
of today emerged after experienced profound modifications, until a surface System evolved in several broad steps and expe-
zoic intermittent feature even across the Neoprotero-
& Canfield, 2011) and euxinia was probably an
Brocks et al. 2003; Arnold et al. 2005; Sarkar et al., 2010; Poulton & Canfield, 2011) and euxinia would, in turn, have resulted in titration of dissolved iron from the water column.

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hed oxygen levels, termed the Great Oxidation Event (GOE) by Holland (2002), changed the face of the Earth. This first Oxygenation Event (e.g. Cloud, 1972; Holland, 1984, 2006; Des Marais et al., 1992; Bekker et al., 2004; Canfield, 2005) was originally believed to have led to at least moderate ventilation of the deep ocean, ending large-scale precipitation of banded iron formations ca 1.8 Ga (Holland, 1984, 2002; Isley & Abbott, 1999). Alternatively, Canfield (1998) proposed that the deep ocean developed widespread euxinia (anoxic-sulphidic water column conditions) after the GOE, because rising oxygen levels would have enhanced sulphate delivery to the ocean which then fuelled bacterial sulphate reduction (Canfield & Raiswell, 1999; Habicht et al., 2002; Strauss, 2004). Widespread euxinia would, in turn, have resulted in titration of dissolved iron from the water column.

Such anoxic and at least intermittently euxinic ocean after the GOE has been substanti-
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magnitude during the Ediacaran–Cambrian transitional interval (Fig. 1; Scott et al., 2008; Sahoo et al., 2012). A similar pattern is observed for other redox-sensitive trace-metals that are influenced minimally by detrital sources, such as vanadium and uranium (Fig. 1; Tribovillard et al., 2006; Och & Shields-Zhou, 2012; Sahoo et al., 2012; Partin et al., 2013).

Due to predictable differences between the behaviour of different palaeoredox proxies
under particular redox conditions, the use of a multiproxy approach is crucial to obtain a more complete picture of the NOE. This is especially true considering that past ocean and atmospheric composition is difficult to quantify directly, while trace-metal budgets may have been significantly different from those of the modern ocean (Fig. 1; Scott et al., 2008). In the present study a multiproxy approach has been applied involving trace element (Mo and V) concentrations, Fe speciation and stable isotopes ($\delta^{14}$N and $\delta^{34}$S), to investigate redox variability, and controls on such variability, across the Yangtze Platform, slope and basin during the Ediacaran–Cambrian transition. The well-preserved organic-rich sedimentary successions on the Yangtze Platform offer an excellent opportunity to study changes in biogeochemical cycling during the Ediacaran–Cambrian transition. Some of the earliest and best preserved Ediacaran and Cambrian fossil assemblages are found on the Yangtze Platform, whereby phosphatized, possible animal embryos from the Ediacaran Doushantuo Formation (Xiao & Knoll, 2000), and the Early Cambrian Chengjiang Fauna, one of the earliest Burgess-shale type fossil occurrences (e.g. Babcock et al., 2001; Hagadorn, 2002), are among the most intensely studied.

Redox geochemistry during the putative NOE has mainly been inferred from single sedimentary successions deposited on the Yangtze Platform (Canfield et al., 2008; Scott et al., 2008; Sahoo et al., 2012), while a relatively simple palaeobathymetry has been assumed in multi-proxy palaeoredox studies (e.g. Li et al., 2010; see also Cui et al., 2015, for a critical re-evaluation). The present study aims to consolidate previously documented changes in redox geochemistry, with estimates of redox reservoir sizes based on S isotopes, and nutrient/redox dynamics inferred from N isotopes and Fe speciation. Furthermore, this multi-proxy approach is used to shed light on the complex inter-relationships between biogeochemical cycling and the broad morphological and bathymetric changes that occurred on the Yangtze Platform during the Ediacaran–Cambrian transition.

GEOCHEMICAL PROXIES

Redox-sensitive trace elements

Molybdenum is present as molybdate ($\text{MoO}_4^{2-}$; Broecker & Peng, 1982) in oxic sea water and is readily sequestered by Mn/Fe-oxyhydroxides and subsequently released to pore waters under reducing conditions (Crusius et al., 1996; Goldberg et al., 2012). In the presence of free $\text{H}_2\text{S}$ (beyond a threshold concentration of ca 11 $\mu$M), molybdate can be converted to particle-reactive thiomolybdate ions ($\text{MoO}_3\text{S}_4^{2-x}$; Helz et al., 1996; Erickson & Helz, 2000) and subsequently scavenged by metal-rich particles, sulphur-rich organic molecules (Helz et al., 1996, 2011; Tribovillard et al., 2004) and iron sulphide minerals (Vorlicek et al., 2004; Poulson Brucker et al., 2012). In the modern ocean, oxic–suboxic facies represent the most important sink for Mo but, despite only making up ca 0.3% of the sea floor, euxinic environments account for ca 30 to 50% of Mo removal from the water column (Bertine & Turekian, 1973; Morford & Emerson, 1999; Anbar, 2004; Algeo & Lyons, 2006), illustrating that Mo is mainly enriched in deposited sediments under sulphidic conditions (e.g. Piper, 1994; Tribovillard et al., 2006).

In oxic sea water, vanadium is present as vanadate oxyanions (H$\text{VO}_4^{2-}$ and $\text{H}_2\text{VO}_4^-$) and readily adsorbs onto both Mn-oxyhydroxides and Fe-oxyhydroxides (Calvert & Piper, 1984; Wehrli & Stumm, 1989) and possibly kaolinite (Breit & Wanty, 1991). Mildly reducing conditions lead to the formation of vanadyl ions ($\text{VO}^{2+}$), related hydroxyl species ($\text{VO(OH)}_3^-$) and insoluble hydroxides $\text{VO(OH)}_2$. More strongly reducing conditions can lead V to be taken up by geoporphyrins or precipitated as solid oxide ($\text{V}_2\text{O}_3$) or hydroxide ($\text{V(OH)}_3$) (Breit & Wanty, 1991; Wanty & Goldhaber, 1992). In general, sedimentary V enrichment begins under denitrifying conditions (e.g. Piper, 1994; Tribovillard et al., 2006), making it more sensitive to less extreme redox conditions than Mo.

Although Mo and V are not dominantly delivered to the sediment in association with organic matter, concentrations often track TOC in anoxic non-sulphidic environments. However, under euxinic conditions, concentrations tend to be decoupled from TOC, resulting in higher Mo/TOC and V/TOC ratios (Tribovillard et al., 2006, and references therein). Molybdenum and, in a less efficient way, V also represent key bio-nutrients which are mainly involved in nitrogenase, an enzyme used in nitrogen-fixing bacteria (Stiefel, 1997; Anbar & Knoll, 2002; Eady, 2003; Zerkle et al., 2006; Canfield et al., 2010b). It has been hypothesized that such trace-metals might have been the limiting nutrients in the Precambrian ocean, as
opposed to phosphorus during the Phanerozoic (Anbar & Knoll, 2002; Planavsky et al., 2010; Zhang et al., 2014; see also Godfrey et al., 2013). Today, however, the budgets of Mo and V are not controlled by marine plankton, i.e. by their roles as micronutrients (e.g. Ho et al., 2003; Tribovillard et al., 2006), and Mo does not exhibit a nutrient-like distribution (concentrations remain relatively constant through the water column), while V exhibits only minor depletion in the photic zone (Collier & Edmond, 1984; Collier, 1985; Piper, 1994).

A complication with the use of Mo-based redox proxies is that the importance of euxinia on a global scale is often assessed from the level of Mo enrichment (or Mo isotope value) in euxinic black shales on a local scale (e.g. Scott et al., 2008). Local Mo enrichment in black shales is generally taken to indicate a lack of Mo sequestration on a global scale, while low Mo concentrations are taken to indicate that the global ocean Mo reservoir has been reduced due to widespread Mo sequestration. However, the absolute level of Mo-enrichment may also reflect the degree of euxinia, or other factors such as basin restriction or sedimentation rate. Therefore, for Mo to be an effective predictor of global euxinia, local redox conditions must be established using independent means, such as Fe speciation, while the nature of the basin and sedimentation characteristics need to be considered.

Iron speciation

Iron speciation analyses of marine sediments are increasingly being used to infer the redox conditions that prevailed in the water column above a depositional environment; for example in modern (Canfield et al., 1996; Raiswell & Canfield, 1996, 1998), Phanerozoic (Raiswell et al., 2001; Poulton & Raiswell, 2002), Proterozoic (Shen et al., 2002, 2003; Poulton et al., 2004, 2010; Canfield et al., 2007, 2008; Sahoo et al., 2012) and Archean (Reinhard et al., 2009; Kendall et al., 2010) settings. The Fe speciation scheme identifies several operationally derived iron pools (Poulton & Canfield, 2005), whereby iron species are characterized as either being unreactive (mostly silicate Fe) or highly reactive towards dissolved sulphide (Fe$_{\text{eq}}$). Highly reactive Fe includes four pools, namely pyrite Fe (Fe$_{\text{py}}$), magnetite Fe (Fe$_{\text{mag}}$), reducible-oxide Fe (Fe$_{\text{ox}}$) and carbonate-associated Fe (Fe$_{\text{carb}}$). The Fe$_{\text{py}}$/Fe$_{\text{HR}}$ ratio must be applied in combination with the Fe$_{\text{HR}}$/Fe$_{\text{T}}$ ratio in order to determine whether the deposition of a given marine sediment occurred under oxic, anoxic-ferruginous or anoxic-sulphidic conditions. In modern marine sediments deposited under an oxic water column, the Fe$_{\text{HR}}$/Fe$_{\text{T}}$ ratio does not exceed 0.38 (Raiswell & Canfield, 1998; Canfield et al., 2008; Poulton & Canfield, 2011). A higher proportion of highly reactive iron would indicate an anoxic water column and, for these samples, the Fe$_{\text{HR}}$/Fe$_{\text{HR}}$ ratio separates ferruginous (<0.8) from sulphidic (>0.8) bottom waters. Ratios between 0.7 and 0.8, however, are equivocal and do not necessarily exclude sulphidic conditions (März et al., 2008; Poulton & Canfield, 2011).

Stable isotopes

The removal of sulphur from the ocean is controlled by pyrite formation (which is often accompanied by a large isotopic fractionation), and the precipitation of evaporites (accompanied by negligible fractionations (e.g. Canfield, 2001). Today, the most important catalyst for sulphur isotope fractionation is the sulphur metabolism of microbes, especially during, but not restricted to, the process of sulphate reduction (Jones & Starkey, 1957; Harrison & Thode, 1958; Kaplan & Rittenberg, 1964) which has been active since the Archean (Shen et al., 2001; Shen & Buick, 2004; Archer & Vance, 2006). The following major controls on isotope fractionation during sulphate reduction can be formulated (Canfield, 2001): (i) when organic electron donors are used, lower specific rates of sulphate reduction lead to higher fractionations; (ii) lower fractionations (3 to 16‰; Kaplan & Rittenberg, 1964; Kemp & Thode, 1968) are achieved when H$_2$ is used as electron donor, particularly at low specific rates of sulphate reduction; (iii) small fractionations (<4‰) occur under sulphate-limiting conditions (ca 200 µM, e.g. Habicht et al., 2002); and (iv) high fractions reaching up to 70‰ occur when sulphate is more abundant (>1 mM; Canfield & Teske, 1996; Canfield et al., 2010a). Controls (iii) and (iv) are commonly used to estimate marine sulphate concentrations.

The stable isotope signature of nitrogen, δ$^{15}$N, can potentially give additional insights into the prevailing redox conditions in the water column, as well as the biochemical reactions involving nitrogen (Quan & Falkowski, 2009; Sigman et al., 2009; Boyle et al., 2013; Quan...
et al., 2013; Ader et al., 2014). Among all water column nitrogen reactions, incomplete denitrification gives the highest positive isotope fractionation, leaving the residual nitrate pool enriched in $^{15}$N, which is subsequently utilized by autotrophs to produce $^{15}$N-enriched organic matter (Sigman et al., 2009). Denitrification occurs at the suboxic/oxic boundary in the water column, and its intensity depends on the size of the nitrate pool and oxidative conditions. On the other hand, N$_2$ fixation by Mo-nitrogenase is carried out by diazotrophic cyanobacteria transferring atmospheric nitrogen into the oceanic pool, whereby the $\delta^{15}$N of newly fixed N is ca $-1_{\infty}$% (e.g. Zhang et al., 2014). Furthermore, as shown in a recent study by Zhang et al. (2014), the $\delta^{15}$N value resulting from nitrogen fixation by alternative (i.e. the V-only and Fe-only) nitrogenases can lead to significantly lower $\delta^{15}$N values (below $-2_{\infty}$%). Generally speaking, vigorous denitrification will increase the nitrogen pool isotopic signature, while intense nitrogen fixation (for example, in the case of sea water nitrogen depletion) will stabilize values at ca $0_{\infty}$%. For these reasons, under anoxic conditions nitrification (and thus denitrification) is inhibited, while under eutinic conditions, including in the photic zone, N$_2$ fixation becomes more highly favoured. In the latter scenario, nitrogen isotopic values can be negative (Higgins et al., 2012; Johnston et al., 2009; Meyers et al., 2012; Struck, 2012; Godfrey et al., 2013). In addition, higher $\delta^{15}$N values in well-preserved sediments can indicate the transition between anoxic and oxic conditions (in both directions), while lower values suggest either an anoxic or anoxic environment (Quan et al., 2008). Because denitrification during early diagenesis does not lead to significant isotopic fractionation in comparison to water column denitrification (Brandes & Devol, 2002), the $\delta^{15}$N signature can be a reliable redox proxy when used in combination with independent measures, such as the redox-sensitive trace-metals Mo and V described above. The present study discusses published $\delta^{15}$N profiles within this wider redox context (Cremonese et al., 2013, 2014). In addition, it has been hypothesized that eutinic conditions, evidenced by high Mo/TOC and high Fe$_{Py}$/Fe$_{HR}$ ratios, will affect the nitrogen isotope composition of organic matter in a systematic way, lowering $\delta^{15}$N to atmospheric values (Boyle et al., 2013). However, this idea can only be tested properly in integrated geochemical studies (e.g. Godfrey et al., 2013).

**GEOLOGICAL SETTING**

**The broader tectonic context**

The Neoproterozoic sedimentary successions of the Yangtze Platform were greatly influenced by the tectonic history of the South China craton, one of three major tectonic cratons in China (Fig. 2). The South China craton consists of the Yangtze and Cathaysia blocks, which amalgamated during the Sibao–Jinning orogeny at ca 900 Ma (Li et al., 1995, 2002, 2003b, 2005). During the break-up of Rodinia, whereby a plume-centre was possibly located under South China, inducing widespread granite intrusions (such as the 819 $\pm$ 7 Ma Huangling Granite in the Three Gorges Area) around the Yangtze block (Li et al., 1999, 2003a), major rift basins formed along the south-eastern and western margins of the South China craton (Li et al., 2003b; Wang & Li, 2003). The subsequent thermal subsidence created the necessary accommodation space for the Cryogenian–Silurian sediments that unconformably overlie Mesoproterozoic metamorphic rocks or early Neoproterozoic rift-related bimodal magmatic rocks, reflecting the different rifting phases (Wang & Li, 2003). The Yangtze Platform can be described schematically as comprising a large platform area, transitioning into a slope and basin (Fig. 2). However, more precise palaeobathymetric reconstructions have shown that a generally shallow-water platform was characterized by numerous intra-shelf basins and carbonate barriers on its margins (Zhu et al., 2003, 2007, 2013; Vernhet, 2007; Vernhet & Reijmer, 2010; Jiang et al., 2011).

The Neoproterozoic sedimentary successions of the Yangtze Platform, which despite the complex tectonic history of China remained relatively undeformed, can be subdivided into three main intervals: pre-glacial predominantly volcano-siliciclastic rocks (for example, the ca 750 Ma Liantuo Formation on the platform and the approximately contemporaneous more fine-grained Banxi Group in the basin), two Cryogenian glacial diamicite intervals (the Guicheng/Tiesiao/Chang’an formations and Nantuo Formation, respectively) separated by an interglacial unit (the Datangpo/Xiangmeng formations) and post-glacial Ediacaran marine carbonates and shales (the Doushantuo Formation and the Dengying/Liuchapo formations). The Early Cambrian successions are found over both the platform and basin areas, character-
ized by black shales with abundant carbonate concretions overlying phosphatic strata of the lowermost Cambrian; they can be very condensed on the platform margin and very thick on the platform (Och et al., 2013), in particular in the south-west (Yunnan & Guizhou provinces).

The Doushantuo Formation

The Doushantuo Formation is among the most extensively studied Neoproterozoic formations worldwide, notably because it accommodates the richest fossil record of this crucial time period, including acritarchs, multicellular algae and controversial fossil embryos (Xiao et al., 1998; Zhang et al., 1998; Xiao & Knoll, 2000; Condon et al., 2005; Jiang et al., 2006; Ling et al., 2007; Yin et al., 2007; McFadden, 2008; McFadden et al., 2008; Ohno et al., 2008; Lu et al., 2013; Zhu et al., 2013). Overlying the glacial diamicrites of the Nantuo Formation, the basal Ediacaran cap carbonate of the Doushantuo Formation represents the aftermath of the end-Cryogenian ‘Snowball Earth’ glaciation (Hoffman et al., 1998). Based on U–Pb age constraints (Condon et al., 2005) and an extremely negative δ13C excursion of arguable duration between ca 565 Ma and 550 Ma, the upper Doushantuo Formation has been correlated with the Johnnie Formation of Death Valley (USA), the Wonoka Formation of the Adelaide rift complex (Australia), the Shuram Formation in Oman, the post-Marinoan Windermere Supergroup (Canada), the Nama and Tsumeb groups of Namibia and south-east Siberia (Le Guerroué et al., 2006; Zhou & Xiao, 2007; Lu et al., 2013).

Although the Doushantuo Formation was deposited over the time span from 635 to 551 Ma (i.e. encompassing ca 90% of the Ediacaran Period; Condon et al., 2005), it nowhere exceeds a thickness of ca 320 m. Whether this reflects a condensed succession or major undetected breaks in sedimentation is presently unclear, although the former possibility is favoured (Zhou et al., 2007). Vernhet (2007) identified three different depositional environments in a study of several sections in Hunan, Guizhou and Hubei provinces, spanning the shallow, subtidal shelf environment, through intertidal or shoal settings, to deep-water basinal environments, illustrating the wide bathymetric range under which sedimentation of the Doushantuo Formation took place on the Yangtze Platform. Two and a half stratigraphic sequences (i.e. transgressive-regressive cycles), have been identified (Zhu et al., 2007, 2013; McFadden et al., 2008); they are, however, clearest in the shallower water facies and not easily discernible in the Yangtze Gorges Area (Zhu et al., 2013).

In the Yangtze Gorges Area, the type locality of the Ediacaran System (known as the Sinian in China), the Doushantuo Formation can be subdivided into four lithological members (Wang et al., 1998). Member I consists of a succession of cap carbonates (dolostone) which extend throughout the central and southern Yangtze Platform; they are characterized by tepee-like structures, sheet cracks, macropeloids, barite crystal fans and negative δ13C values (Jiang et al., 2003a, 2006; Zhou et al., 2004). A precise U–Pb age of 635.2 ± 0.6 Ma has been determined from an ash layer within the cap carbonate at one locality (Condon et al., 2005). In the Three Gorges Area, the cap carbonates have a thickness of ca 5 m and are thus relatively thin compared to some basal Ediacaran cap carbonates around the world (Hoffman et al., 2007). The overlying second member is between 80 m and 140 m thick and is composed of interbedded black shale, organic-rich calcareous mudstone and thinly bedded dolomicrite. An ash layer dating from 632.5 ± 0.5 Ma is situated a few metres above the cap carbonate (Condon et al., 2005). Abundant centimetre-sized chert nodules, decreasing up-section, occur in the lower-middle part of Member II and contain numerous acanthomorphic acritarchs and multicellular algae (Zhang et al., 1998; Yin et al.,

Fig. 2. Simplified geological map of South China with the locations of the studied sections in both, the Yangtze Platform and the basin (modified after Steiner et al., 2001; Ling et al., 2007) with a simplified stratigraphic overview of the Yangtze Platform (modified after Jiang et al., 2007). The smaller scale map shows the geological map of the Three Gorges Area (modified after McFadden et al., 2008) with a more detailed stratigraphy of the Precambrian-Cambrian transition. The pink shaded sections indicate occurrences of the main black shale successions. The yellow dots on both maps indicate the sections analysed for the present study whereas the Baigouyuan section (white dot) has been considered from earlier analysis (Wallis, 2006). Localities investigated during recent studies by Li et al. (2010) and Sahoo et al. (2012) have also been included for comparison. Shp = Shipai Formation; Cr = Cryogenian Period.

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Sections:

This study:

1. Maoshi
2. Huanglian
3. Longbizui
4. Xiaotan
5. Jiulongwan
6. Jijiapo
7. Jiuqunao

Sahoo et al., 2012:

8. Taoying
9. Wuhe
10. Yuanjia

Li et al., 2010:

11. Zhongling

Ediacaran

Huangling Granite

Palaeoproterozoic

Fault

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The sparse sedimentary structures include parallel laminations, crinkle laminations and rare intraformational breccias, indicating that wave and current activity were mostly absent during sedimentation of member II (Zhou et al., 2007). This is likely to have resulted from the Yangtze Gorges Area being: (i) a protected basin; and/or (ii) overlain by a water depth exceeding ca 100 m (i.e. below ‘storm-wave base’; Zhu et al., 2013).

Member III is between 30 m and 60 m thick and comprises medium to thick-beded dolostone with thin chert horizons and irregular chert nodules that grade up section into thin-beded limestone and dolomitic interbeds (i.e. ribbonites). Although most chert layers are diagenetic in origin, some contain well-preserved microfossils (Zhang et al., 1998; Xiao, 2004; Yin et al., 2007). Sedimentary structures include scour marks, crinkle laminations and low angle cross-bedding, while sandy layers capping limestone and dolomite bedding surfaces are common, indicating that deposition occurred at shallower water depths than the underlying Member II, possibly in shallow subtidal environments (Zhou et al., 2007). Member IV, commonly referred to as the Miaoho Member, comprises a succession of black shales and organic-rich mudstones. Sedimentary structures are absent apart from the fine lamination and abundant, sometimes huge (diameter >1 m), carbonate concretions of diagenetic origin (e.g. Lu et al., 2013; Zhu et al., 2013). Pyrite and barite are also common features of this member. An ash bed at the top of the Miaoho Member yields a U–Pb age of 551.1 ± 0.7 Ma, and a 3 to 4 cm thick bentonite bed that in some sections separates the Doushantuo Formation from the Dengying Formation has yielded an age of 550.4 ± 0.8 Ma (Condon et al., 2005).

Studied sections
Black shale successions of the Doushantuo Formation were studied, predominantly from Members II and IV, at several locations on the Yangtze Platform: Jiulongwan (Jiulongwan and Xiangdangping) and Jiuquanao (Jijiawan) in the Yangtze Gorges Area, Hubei Province, and at Maoshi (north-west of Zunyi, Guizhou Province). At the Jiulongwan section, a substantial part of Member II was sampled from directly above the cap carbonate throughout a tectonically disrupted succession of massive grey dolomite beds with interbedded black shales. Further outcrops sampled at Jiulongwan comprise the >10 m thick Miaoho Member, consisting of laminated black shales, barite and abundant huge carbonate nodules which lie between Member III and the wavy, shaly horizon with a pyrite-rich layer that constitutes the contact to the overlying Dengying Formation. Below the Miaoho Member, the top of Member III consists of dark grey dolomite which transitions downwards into paler carbonate ribbonites, followed by dolomite beds with some intercalated chert beds. The boundary between the Doushantuo Members II and III is obscured by poor outcrops and dense vegetation in this region.

At the Jiuquanao section, the lower black shales of the Doushantuo Member II and the black shales of the Miaoho (Doushantuo Member IV) have been investigated. Parts of Member III, particularly towards the top, consist of massive dolostone, which is considered to represent allochthonous blocks, due to either slumping and/or faulting (Lu et al., 2013).

The upper Doushantuo Formation, including the transition to the overlying Dengying Formation was also sampled at the Maoshi section, Guizhou Province, where it was presumably deposited in a lagoonal setting close to the margin of the Yangtze Platform (e.g. Jiang et al., 2011). Here, the upper Doushantuo Formation is characterized by carbonate-rich black shales that can tentatively be correlated with the Miaoho Member (Doushantuo Member IV). The black shale unit at the top of the Doushantuo Formation occurs in almost all palaeoenvironments of the Yangtze Platform and its sharp onset represents a flooding surface (Jiang et al., 2007) rather than a major sequence boundary (Zhu et al., 2007, 2013). Similar to the cap carbonates, it can be used as a regional stratigraphic marker. However, due to lithostratigraphic variability within the Doushantuo Formation throughout South China, it is unclear to what extent the subdivision into four members can be applied away from the southern limb of the Huangling granite in the Yangtze Gorges Area (Zhu et al., 2007, 2013).

The Dengying/Liuchapo formations
In contrast to the underlying Doushantuo Formation, a relatively short time interval, from ca 551 to 541 Ma, is represented by the Dengying Formation (Zhao et al., 1988). In places, this formation is much thicker (240 to 850 m), and can be subdivided into three members. In the Three Gorges Area these are: (i) the Hamajing Member...
at the base (20 to 190 m thick), consisting of light-grey, medium to thick-bedded dolomite with intercalated thin chert layers; (ii) the overlying Shibanantan Member (100 to 160 m), characterized by dark grey, thin-bedded limestone; and (iii) the Baimatuo Member (60 to 570 m), which is composed of light-grey, thickly bedded/massive dolomite. The tripartite subdivision of the Dengying Formation can be recognized in many places in South China, although a different terminology is sometimes used (Ding et al., 1992; Zhu et al., 2003).

A few samples near the top of the Dengying Formation, just below the Early Cambrian successions, notably the Jijiapo (Hubei Province) section, were collected and analysed but, due to their trace-metal content close to detection limit, they will not be discussed in the present study (see Och, 2011, for details). At the Maoshi section, the lower part of the Dengying Formation comprises a succession of alternating black carbonates, which get paler up-section, and sandy carbonates. At the Jijiapo section, Three Gorges Area, the base of the Hamajing Member consists of wavy beds of massive dolomite. The base of the Shibanantan Member is characterized by thinly bedded limestone followed by a succession of dark, macrocrystalline limestone beds, a few intercalated chert layers and chert nodules. The top of the Shibanantan Member is rich in chert nodules and possibly grades into the Baimatuo Member with low amplitude wavy bedding of the carbonates.

The depositional environment of the Dengying Formation is interpreted in terms of a prograding platform. Oolitic textures and oncolites in the dolomitic Hamajing Member are characteristic of a shallow, high-energy environment following black shale deposition at the top of the Doushantuo Formation, indicating a sea-level drop from the Doushantuo to the Dengying Formation (Zhu et al., 2007, 2013). Towards the south-east, the carbonate successions become gradually thinner, ultimately changing into the slope and basinal facies of the corresponding Liuchapo Formation, which is mainly composed of black silicified shales that in places reach into the early Cambrian (e.g. Wang et al., 1998; Chang et al., 2009).

Studied sections
At the Huanglian section, Guizhou Province, stratigraphic control is rather poor, but an equivalent level there to the Liuchapo Formation was sampled, consisting of black silicified shale below a black shale succession. The nearby Longbizui section, Hunan Province, on the other hand, offers an excellent transitional succession from cherty shales and bedded cherts of the predominantly Late Ediacaran Liuchapo Formation, to the black shales of the Early Cambrian Niutitang Formation.

Early Cambrian formations
The global standard stratotype-section and point (GSSP) for the base of the Cambrian System is the first appearance datum (FAD) of the trace fossil Trichophycus (Treptichnus) pedum (Brazier et al., 1994; Landing, 1994; Babcock & Peng, 2007). However, due to a lack of convincing evidence for the occurrence of this trace fossil in Cambrian sediments in South China and absolute age constraints (Compston et al., 1992, 2008; Yang et al., 1996; Jenkins et al., 2002; Jiang et al., 2009; Och et al., 2013), correlation of the Precambrian–Cambrian boundary focuses instead on the biostratigraphy of small shelly fossils (Steiner et al., 2007), as well as C-isotope stratigraphy (for example, at the Meishucun section sampled for this study; Cowie, 1985; Luo et al., 1992; Shields et al., 1999; Zhu et al., 2003). Because of the strongly varying abundance of small shelly fossils on the Yangtze Platform, phosphorite horizons occurring in the lowermost Cambrian have also been taken into account in the present study, where the Precambrian–Cambrian boundary was not obvious stratigraphically.

The Niutitang Formation
The Niutitang Formation, which is locally strongly condensed, comprises mainly black shales, siltstones, intercalated chert, organic-rich carbonates and phosphate nodules. It predominantly crops out within the transitional belt at the platform margin and, often uncomfortably, overlies the Dengying/Liuchapo formations. Phosphorite beds at the base of the Niutitang Formation and organic-rich black shales above can be used as marker horizons across the slope and basin of the Yangtze Platform; these have predominantly been studied because of an extreme enrichment of Ni and other metal sulphides, in particular Mo with concentrations up to several per cent (e.g. Och et al., 2013, and references therein).
Studied sections
The base of the Niutitang Formation has been sampled at the basinal Huanglian and Longbizui sections, and mainly consists of an uninterrupted black shale succession. In the more extensively sampled Longbizui section, two distinct layers of pyrite and carbonate concretions have been found above the Liuchapo–Niutitang boundary.

The Yanjiahe and Shuijingtuo formations
The Yanjiahe Formation, which covers the Precambrian–Cambrian transition south of the Huangling anticline in the Three Gorges Area, overlies pale, massive dolomites of the Dengying Formation (Baimatuo Member) above a sharp, wavy contact. The Yanjiahe Formation is ca 35 m thick and consists of dolomitic muddy limestone, calcareous black shale and some sandstone and chert (Ishikawa et al., 2008). The overlying Shuijingtuo Formation is ca 100 m thick and mainly consists of black shale with prominent carbonate concretions (Ishikawa et al., 2008). Mainly based on carbon isotope stratigraphy, it is likely that the upper part of the Yanjiahe Formation is equivalent to the Dahai Member of the Early Cambrian Zhujiaqing Formation (Li et al., 2013; Cremonese et al., 2014), while the presence of sponge spicules and trilobite fragments suggests that the overlying Shuijingtuo Formation could be considerably younger, possibly even Cambrian Stage 3.

Studied sections
The sampling of the Yanjiahe and Shuijingtuo formations was conducted at the Jijiapo and Jiuqunao section where the base of the Yanjiahe Formation is characterized by grey carbonate beds with thin chert intercalations. Abundant phosphatic cherts were recognized in the middle part of the formation, whereas massive dolomite beds, followed by organic-rich black dolomite interbedded with siltstone, including regularly shaped silicified carbonate nodules, occur in the upper part below the boundary to the Shuijingtuo Formation. The boundary zone is characterized by a thin (ca 10 cm) phosphorite bed, followed by a conglomeratic layer with rip-up clasts and framoidal pyrite. The basal Shuijingtuo Formation consists of black shales with abundant, metre-sized dolomite concretions and some intercalated grey massive dolomite beds.

MATERIALS AND METHODS
One hundred forty-three well-preserved predominantly black shale samples and some more or less organic-rich carbonate samples were collected at seven localities during two field seasons (2008 and 2009) on the Yangtze Platform, from the Doushantuo Formation Member II to the early Cambrian Shuijingtuo Formation. Sampling steps vary considerably between highly resolved segments of decimetre-scale to broad steps of over 10 m. All of the present samples were converted into powders using a tungsten carbide mill.

Geochemical analyses included X-ray fluorescence carried out on a Philips PW2400 spectrometer (PANalytical, Almelo, The Netherlands) for major and trace elements on 54 samples collected during the first field season (Maoshi, Huanglian and Longbizui). The 89 samples collected during the second field season in the Three Gorges Area (Jiulongwan, Jiuqunao, Yangjiahe and Jijiao) were dissolved afterashing at 550°C before analysis for bulk elemental composition using ICP-MS and Fe speciation. In detail, 5 ml of nitric acid was added to the ashed powder, before adding 2 ml of hydrofluoric acid and a few drops of HClO4. The sample was then dissolved on a hot plate. After evaporation, 2.5 ml of boric acid (50 g l−1) acid was added and evaporated to dryness on a hot plate to dissolve any aluminium hexafluorides. The samples were then taken up in 5 ml of hot 50% HCl. Iron speciation analysis was carried out on all samples according to the method outlined by Poulton & Canfield (2005). The present authors measured the concentrations of four different highly reactive iron phases (FeT(q)); pyrite iron (FePy), carbonate-associated iron (FeCarb), ferric oxides (FeOx) and magnetite iron (FeMag), and total iron content (FeT); FePy was determined via the chromous chloride technique of Canfield et al. (1986), while AVS (FeS) was only detected in trace amounts. The sequential extraction procedure of Poulton & Canfield (2005) was used to determine the other highly reactive iron phases, specifically FeCarb, FeOx and FeMag. To determine FeT, ca 100 mg sample powder was dissolved using the above-mentioned method for total dissolution and then diluted 50 times. All solutions were analysed for their respective iron contents using an atomic absorption spectrometer (AAS), with a relative standard deviation (RSD) of <5% for all stages. The same sample solutions were then evaporated to dryness and redissolved in
nitric acid in order to measure the trace-metal contents. The final solution, containing 0.15 ml HNO₃, 0.1 ml 500 ppb Re for instrument calibration and 3.75 ml H₂O, was analysed at the State Key Laboratory for Mineral Deposits Research, Nanjing University, using a Finnigan Element II inductively coupled plasma mass spectrometer (ICP-MS; Thermo Fisher Scientific Inc, Waltham, MA, USA). The precisions were generally better than 5% for the analysed elements based on long-term uncertainty of the laboratory measurement on a standard carbonate sample.

Samples which yielded enough Ag₂S residue (>0.03 g) after the sulphide extraction procedure were analysed for sulphide isotopes using a Finnigan MAT DeltaPlus plumbed to a Carlo Erba elemental analyser through a Conflo II interface (CE Elantech Inc, Lakewood, NJ, USA). All analytical work for sulphur isotopes was carried out at the Institute for Geology and

Fig. 3. Stratigraphy and geochemical profiles across the Doushantuo Member II at the Jiulongwan section, Hubei Province. Note that, due to extensive vegetation, considerable gaps along the stratigraphy had to be taken into account. The analysed samples are indicated on the left. The yellow shaded sections show deposition under possibly euxinic conditions indicated by Fe₉⁸/Fe₉₁ ratios above 0.7 (red dashed line). Anoxic conditions are indicated by Fe₉₁/Fe₉₅ ratios above 0.38 (black dashed line): TOC, total organic carbon; TS, total sulphur; Fe₉₅, total iron; Fe₉₁, highly reactive iron; Fe₉₈, pyrite iron.
Paleontology, Westfälische Wilhelms-Universität Münster, Germany. Nitrogen isotopic analyses have previously been published in Cremonese et al. (2014) and the methods are described therein.

The evaluation of TOC, total carbon (TC) and total sulphur (TS) was carried out using a Leco C/S analyser (Leco Corporation, St Joseph, MI, USA) at the Wolfson Laboratory, University College London. The TC and TS were measured directly after weighing between 100 mg and 300 mg of sample powder. The TOC content was determined after dissolving each sample with 10% HCl at room temperature until no effervescence was observed, followed by filtering and rinsing with H₂O prior to introducing the dried remains into the C/S analyser.

Fig. 4. The upper part of member III and the complete member IV (Miaohe) of the Doushantuo Formation at Jiu-longwan. See Fig. 3 for abbreviations and formatting. DNG = Dengying.
RESULTS

The Doushantuo Formation

The extensive Jiulongwan sections (JLW) in the Three Gorges Area, Hubei Province, span from Doushantuo Member II (DST II) to Doushantuo Member IV (DST IV). For these sections, Mo and V concentrations mostly remain well below average shale values, until the onset of DST IV (Figs 3 and 4). In this late Ediacaran black shale succession, Mo concentrations reach ca 370 ppm, with concomitant V concentrations of over 2400 ppm found just at the onset of black shale deposition in DST IV. Molybdenum correlates only moderately with V, with a more significant correlation within DST IV ($R^2 = 0.53$) than DST II ($R^2 = 0.25$). Total organic carbon shows no significant correlation with Mo, but is weakly correlated with V ($R^2 = 0.34$) within the lower DST II, where TOC contents are generally below 1% but with a few peaks of up to 2-4%. Within DST IV, no significant correlation is found between trace-metals and TOC contents, where TOC rises smoothly after the DST III/DST IV boundary, stabilizes between 4 wt% and 6 wt %, and reaches a maximum concentration of 14-9% towards the top. Total sulphur and pyrite sulphur correlate well throughout the section, but in particular within DST II, where less non-pyrite sulphur is found within DST IV. The Mo/TOC ratios within DST II are highly variable, ranging up to $4 \times 10^{-4}$. The Mo/TOC ratio reaches a maximum of $180 \times 10^{-4}$ at the bottom of DST IV and rapidly declines to oscillate between 0 and slightly above $40 \times 10^{-4}$. Iron speciation analysis suggests deposition under anoxic conditions, with DST II sediments indicating a ferruginous water column ($Fe_{HR}/Fe_T > 0.38$; $Fe_{py}/Fe_{HR} < 0.7$ to 0.8) and the overlying

Fig. 5. Stratigraphy and geochemical profiles for the black shales of the Doushantuo members II and IV at the Jiuqnao section. The distance between the members DST II and DST IV is unknown. Note that $Fe_{py}/Fe_{HR}$ ratios reach high values within DST IV but are not high enough to be identified as a euxinic depositional environment. See Fig. 3 for abbreviations and formatting. DNG = Dengying.
DST IV indicating at least intermittently euxinic conditions $\text{Fe}_{\text{HR}}/\text{Fe}_T > 0.38$; $\text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}} > 0.7$ to 0.8).

The pyrite $\delta^{34}$S profile is very variable but initially positive with values between 0‰ and 23‰ within DST II (Fig. 3). Further up in the Doushantuo Formation (Fig. 4), a decline occurs from ca $-7\%_\text{o}$ within the carbonates of DST III, to $-19\%_\text{o}$ within the upper part of the Miaohé black shale (DST IV), which is followed by a steep increase back to $-7\%_\text{o}$ below the Doushantuo/Dengying boundary. Within DST II, $\delta^{15}$N values are also positive and vary between 1.9% and 6.7‰ without any particular trend (Fig. 3). In the upper part of DST III the $\delta^{15}$N profile exhibits an increase from 1.3 to 4.3‰ towards the Miaohé Member, at which value they remain approximately except for a single, minor excursion down to 3.2‰ at the top (Fig. 4).

Throughout the Doushantuo Formation at the Jiuzi section (JJ), Hubei Province, very low redox-sensitive trace-metal concentrations close to average shale were measured (Fig. 5). Anoxic conditions can, however, be inferred from iron speciation analysis throughout the section but, although relatively high $\text{Fe}_{\text{py}}/\text{Fe}_{\text{HR}}$ ratios of between 0.6 and 0.7 are common within the black shale of DST IV, there is no convincing evidence for euxinic water column conditions. The $\delta^{34}$S values for pyrite show a sharp increase from $-10$ to $+10\%_\text{o}$ within DST II and remain at this approximate level. In the Miaohé Member, $\delta^{34}$S values are ca $+15\%_\text{o}$ and exhibit a decrease down to $+10\%_\text{o}$ 1 m below the Doushantuo/Dengying boundary. The $\delta^{15}$N profile basically shows a decrease from values between 5‰ and 6‰ within DST II, to values that decrease from 2.5 to 0.6‰ in the Miaohé Member.

The Maoshi section (MS), in Guizhou Province spans the boundary between the late Ediacaran Doushantuo and Dengying formations, and exhibits average shale Mo and V concentrations of ca 3 ppm and 100 ppm, respectively (McLennan, 2001; Fig. 6). The TOC contents are high in the upper Doushantuo Formation (1 to 3.4‰) and fall below 1‰ in the overlying Dengying Formation, with only a weak correspondence between peaks in TOC and TS. The correlation between the trace-metals and TOC and TS is weak but the highest Mo and V con-

Fig. 6. The stratigraphy and geochemical profiles at the Maoshi section, Guizhou Province. See Fig. 3 for abbreviations and formatting.
Concentrations are mostly concomitant with the highest TOC and TS values. There is a very strong correlation between TS and pyrite S, but with a significant excess of non-pyrite S. The Mo/TOC ratios show a slightly increasing trend towards the top of the Doushantuo Formation but remain very low, with a maximum of $3.4 \times 10^{-4}$. Iron speciation indicates anoxic-ferruginous conditions with possibly intermittent euxinia in the upper Doushantuo Formation, suggested by $\frac{Fe_{Py}}{Fe_{HR}}$ ratios of up to 0.77. The $\delta^{34}S$ values for pyrite recovered from the black shale of the upper Doushantuo Formation exhibit a decreasing trend from 18.5 to $-1.5\%_o$, while the $\delta^{15}N$ profile for the same samples is quite constant with high values of ca. $+8\%_o$.

The Late Ediacaran (Dengying/Liuchapo formations) and Early Cambrian formations

The Huanglian section, Guizhou Province, which covers the Precambrian–Cambrian boundary from the Liuchapo (equivalent) Formation to the Niutitang Formation, exhibits significant variability in trace-metal concentrations (Fig. 7). Maxima of 96 ppm for Mo and 9860 ppm for V at the base of the Niutitang Formation coincide with very high TOC contents of up to 9.6%, leading to a moderate correlation between Mo, V and TOC. Sulphur contents are relatively low and a moderate correlation between TS and pyrite S prevails, with substantial non-pyrite S at the base of the Niutitang Formation. Nevertheless, Mo concentrations correlate relatively well with TS ($R^2 = 0.67$), indicating that Mo is efficiently scavenged into sulphidic sediment even when pyrite formation is minor. A relatively high Mo/TOC ratio of $19 \times 10^{-4}$ is found within sediments of the upper Doushantuo Formation, while low values prevail within the Liuchapo Formation, increasing again at the base of the Niutitang Formation, where a maximum Mo/TOC ratio of $10 \times 10^{-4}$ is found. An anoxic-ferruginous depositional environment is clearly indicated by Fe speciation analysis, with $\frac{Fe_{Py}}{Fe_{HR}}$ ratios that vary considerably but remain low at between 0.01 and 0.56. Due to a lack of pyrite, only two samples could be analysed for $\delta^{34}S$, yielding $-4.3\%_o$ and $-6.3\%_o$. The $\delta^{15}N$ values are low, but positive, with values no higher than $2.4\%_o$ in the Liuchap Formation, decreasing to ca. $0\%_o$ in the Lower Cambrian Niutitang Formation.

Within the predominantly cherty shales and chert beds of the Liuchapo Formation at the Longbizui section (LBZ), a thin black shale interval is found where TOC and TS concentrations both reach over $3.1\%$ within a thin horizon, while Mo and U remain depleted and V is only slightly enriched, with concentrations of up to 340 ppm (Fig. 8). The $\frac{Fe_{Py}}{Fe_{HR}}$ ratios exceed 0.7 within an interval of 15 cm just below and within this organic-rich horizon, below which $\delta^{34}S_{pyrite}$ values range widely.
between +66‰ and a minimum of −0.38‰. The overlying Liuchapo Formation is characterized by TOC and TS contents below 1% and relatively low trace-metal concentrations, with V up to 1937 ppm and Mo up to 107 ppm, broadly tracking TOC and TS contents. The TS contents correlate well with pyrite S, although variable amounts of non-pyrite S (averaging 50% of TS) are observed. The Mo/TOC ratios are low prior to the Niutitang Formation, where they reach values of between 7.0 × 10⁻⁴ and 15.8 × 10⁻⁴. On the other hand, V/TOC ratios show a maximum of 553 × 10⁻⁴ just below the Niutitang black shales, and decrease to low values of between 26 × 10⁻⁴ and 190 × 10⁻⁴ further up. The Fe₉₉/Fe₉₅ ratios indicate anoxic conditions throughout the section, and Fe₉₅/Fe₉₉ ratios between 0.7 and 0.8 suggest possibly sulphidic conditions during extended intervals within the Niutitang black shale. Sulphide isotope values remain positive throughout the remaining section, δ³⁴S being enriched by between 9.3‰ and 29.6‰. The δ¹⁵N profile exhibits higher values of between 1‰ and 3.7‰ within the Liuchapo Formation, which decrease sharply to negative values of ca −2‰ at the Liuchapo/Niutitang Formation boundary.

The Jijiapo section (YS), Hubei Province, includes samples from the Lower Cambrian formations which are strongly depleted in Mo within the organic-poor Yanjiahe samples, but show enrichment within the overlying Shuijingtuo Formation, where Mo reaches concentrations of up to 37 ppm and Mo/TOC ratios up to 32 × 10⁻⁴ (Fig. 9). Good correlation is observed between TS and Mo contents (R² = 0.92),
whereas V concentrations are significantly better correlated with TOC ($R^2 = 0.61$). While TS correlates with pyrite sulphur, a great excess of non-pyrite sulphur was observed. Iron speciation analysis supports anoxic-ferruginous conditions without evidence for sustained euxinia. The δ$^{34}$S values within samples from the organic-poor Yangjiahe Formation are all positive, with an increase from values of ca $5\%$o to over $23\%$o beneath the overlying Shuijingtuo Formation, followed by a subsequent decrease down to $-5\%$o. The δ$^{15}$N profile shows constant values of slightly below $1\%$o within the Yanjiahe Formation except at the top, where values of between $-2.3\%$o and $+5.6\%$o occur within the lowermost Shuijingtuo Formation.

At the Jiuqnao section (JJL), redox-sensitive trace-metal concentrations are significantly enriched in the lower Cambrian black shales of the Shuijingtuo and Yanjiahe formations (Fig. 10). At the onset of the black shale succession of the Shuijingtuo Formation, Mo peaks at 213 ppm with a concomitant peak in V concentration of 910 ppm, before concentrations decrease to relatively constant values of ca 40 ppm and 150 ppm, respectively, with an overall good correlation between Mo and V ($R^2 = 0.95$). The Mo/TOC ratios track the Mo concentration profile, attaining a maximum of $36 \times 10^{-4}$. Total organic carbon, TS and pyrite sulphur also correlate well within the Early Cambrian formations, where a significant amount of non-pyrite sulphur is present. In the Shuijingtuo Formation, frequent euxinic conditions are plausible with $Fe_{Py}/Fe_{HR}$ ratios between 0.7 and 0.8. Interestingly, the maximum peaks in Mo and V

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contents occur at the stratigraphic level where FePy/FeHR ratios are particularly low. In the Shuijingtuo Formation, the one carbonate sample (JJL1) analysed for its δ34S signature within the lower Shuijingtuo Formation has a value of 18.6‰. Similarly, only a few δ15N values were measured, but they show a clear decrease from ca 0–8‰ within the carbonate samples at the bottom, to ca –0.5‰ in the TOC-rich black shale further up, before a slight increase (but remaining negative) for the rest of the section.

DISCUSSION

Figures 11 and 12 summarize the most important geochemical indicators considered in the present study. Figure 11 focuses on the Ediacaran and Cambrian strata found in the Three Gorges Area and also includes Mo/TOC and V/TOC ratios of the Baiguoyuan section north of the Huangliang granite (reported by Wallis, 2006). Figure 12 shows the data for the Precambrian/Cambrian transition at Huanglian and Longbizui which were deposited in a basin setting. The Xiaotan section, at the boundary between the Yunnan and Sichuan provinces (Fig. 2), contains an unusually expanded Early Cambrian succession to the north-west of the Yangtze Platform. The trace-metal concentrations and iron speciation analyses have previously been reported by Och et al. (2013) and the nitrogen isotope signature by Cremonese et al. (2013). These results are summarized here and include a previously unpublished δ34S profile (Fig. 13). The most significant geochemical feature is the relatively low FeHR/FeT ratio within the Yuanshan Formation, down to 0.2 at the top, which indicates that the sediments were potentially deposited beneath an oxic water column (see Supporting Information; Poulton & Canfield, 2011).

In the last few years, research on biogeochemical cycling on the Yangtze Platform during the Late Ediacaran has intensified. Li et al. (2010), Sahoo et al. (2012) and, more recently, Kikumoto et al. (2014), have contributed greatly to the growing understanding of the processes that prevailed during the Precambrian–Cambrian transition and presented datasets that complement the findings of the present study (Fig. 14). The study by Li et al. (2010) introduced the concept of a sulphidic wedge during deposition of the Miaohe Member (DST IV) to explain Mo enrichment at Jiuqunao, while little Mo
A multiproxy approach to ancient ocean chemistry

The present study aims to shed light on a crucial time in Earth history, the Neoproterozoic Oxygenation Event (NOE), when changing redox conditions in the ocean were succeeded by an unprecedented rise in the diversity and architectural complexity of organisms. Within this context and using a multiproxy elemental and isotopic approach, the palaeobathymetric situation and the degree of restriction of the regional and local marine environments on the Yangtze Platform during this transition can better constrain the framework for the NOE.

The Nantuo Formation, deposited during the end-Cryogenian ‘Marinoan’ Glaciation, represents the last stage of the rifting history of the Yangtze Platform, which left a sea floor with abundant horst and graben structures, onto which sediments of the Doushantuo Formation have been draped (e.g. Vernhet, 2007; Zhu et al., 2013). In addition, the Huangling granite intrusion and its subsequent erosion during multiple phases of glaciations are likely to have had a major impact on the bathymetry of the Three Gorges Area, in particular. A mosaic of different depositional environments can therefore be imagined, which has been confirmed by studies demonstrating significant lateral facies varia-

sequestration occurred at Zhongling ca 100 km in the south. These authors used a simplified platform-basin model and concept that has recently been disputed by Cui et al. (2015) on the basis of a revised chemostratigraphic framework. On the other hand, Sahoo et al. (2012) presented a substantial Mo enrichment further south and earlier in time during the deposition of DST II. While this phenomenon persists until the Cambrian within the Jiuqunao and Maoshi sections, a massive trace-metal enrichment is observed in the Jiulongwan and Baiguoyuan sections within the Maoshi Member alongside negative $\delta^{34}S$ values at Jiulongwan. Cry = Cryogenian Period; Nto = Nantuo Formation; Shp = Shipai Formation.

Fig. 11. Schematic summary plot for all sections sampled in the Three Gorges Area including the Maoshi section, Guizhou Province, which is situated ca 530 km to the south-west (see Fig. 2). The reddish shaded intervals indicate anoxic-ferruginous conditions and the yellow shaded intervals suggest a euxinic depositional environment. Note the very low Mo and V enrichment relative to TOC and the almost exclusively positive $\delta^{34}S$ values within the DST II. While this phenomenon persists until the Cambrian within the Jiuqunao and Maoshi sections, a massive trace-metal enrichment is observed in the Jiulongwan and Baiguoyuan sections within the Maoshi Member alongside negative $\delta^{34}S$ values at Jiulongwan. Cry = Cryogenian Period; Nto = Nantuo Formation; Shp = Shipai Formation.
tions, notably around the Three Gorges Area (e.g. Vernhet & Reijmer, 2010; Jiang et al., 2011; Zhu et al., 2013). The stratigraphic and palaeoenvironmental ambiguity, however, has contributed to a wealth of different ideas on the depositional environment of the Doushantuo Formation, i.e. whether it was deposited in a non-marine basin (Bristow et al., 2009), an intra-shelf basin (Vernhet & Reijmer, 2010) or a shelf lagoon (Jiang et al., 2011). The sections below begin by evaluating environmental conditions during deposition of the Doushantuo Formation, and then expand this palaeoenvironmental analysis to track environmental conditions into the Early Cambrian.

**The Ediacaran**

**Doushantuo Member II**

The short interval sampled for Doushantuo Member II (DST II) at Jiuqunao is geochemically similar to the more expanded profile analysed at Jiulongwan (Figs 4 and 5), where no enrichment in redox-sensitive trace-metals, comparable TOC and TS contents, generally anoxic non-sulphidic conditions and mostly positive $\delta^{34}S_{\text{pyrite}}$ values (above 10‰) are found. Previous studies of the Jiulongwan section agree with the present results (Bristow et al., 2009; Li et al., 2010) and suggest that a lack of trace-metal enrichment was a characteristic of the Yangtze Platform during the deposition of DST II. However, significantly

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**Fig. 12.** Mo and V concentrations normalized to TOC, pyrite sulphur and nitrogen isotopes for the basinal sections at Longbizui and Huanglian. Note that the Mo/TOC and V/TOC increase at the base of the Cambrian and the concurrent decrease of $\delta^{34}S$ and $\delta^{15}N$ values.

**Fig. 13.** Summary of geochemical results from the Early Cambrian Xiaotan section (Cremonese et al., 2013; Och et al., 2013). ZJQ = Zhujiaqing Formation; CLP = Canglangpu Formation.
higher enrichments of Mo and V in DST II have been reported from slope and basin sections, where Mo concentrations can attain values from a few tens of ppm (Wallis, 2006; Guo et al., 2007) up to almost 200 ppm (Sahoo et al., 2012).

Although there are possible indications of intermittent euxinia in DST II at Jiulongwan, suggested by high Fe$_{Py}$/Fe$_{HR}$ ratios ca 0-7, no Mo enrichment is apparent. Because Mo/TOC and V/TOC ratios remain very low throughout the lower part of the Doushantuo Formation, it is argued here that limited availability of Mo and V in the water column prevailed, at least locally, during deposition of DST II at Jiulongwan.

Independently, palaeogeographic, sedimentological and carbon isotope studies support a rimmed platform margin, which developed soon after deposition of the underlying cap carbonate (Jiang et al., 2011), and the formation of variably restricted basins on the Yangtze Platform (Jiang et al., 2011).

Fig. 14. Other geochemical profiles from previous studies carried out on the Yangtze Platform near the Three Gorges Area (see Fig. 2 for localities). (A) Li et al. (2010) and Kikumoto et al. (2014) also investigated the Jiulongwan section across the whole Doushantuo Formation and that data confirms the findings of the present study. The Doushantuo Formation at the Zhongling section, ca 100 km south of Jiulongwan, exhibits much lower Mo/TOC ratios and a $\delta^{34}$S profile that mirrors the trend at Jiulongwan up to DST Member II before showing a similarly decreasing trend within DST Members III and IV. (B) Sahoo et al. (2012) found elevated Mo/TOC ratios in the DST Member II and generally very low $\delta^{34}$S values in basinal sections.
is not indicated at Jiuqunao (Figs 4 and 5), which sited under euxinic conditions, whereas euxinia the Miaohe Member at Jiulongwan were deposited in a restricted environment with high concentrations of redox-sensitive trace-metals (Fig. 14), this indicates enhanced availability of sulphate within the open ocean, resulting in an increased potential for rates of sulphide production to overwhelm the influx of highly reactive Fe, thus leading to euxinic conditions (Poulton & Canfield, 2011) and, hence, extensive trace-metal drawdown. This condition is indicated by low Mo/TOC ratios in DST II on the shelf margin (Li et al., 2010; this study) and greatly elevated Mo/TOC ratios in basinal successions (Sahoo et al., 2012). The nitrogen isotopic record of DST II, where most values lie between 3‰ and 6‰ in the Jiulongwan section (see also Kikumoto et al., 2014), and at 6‰ at Jiuqunao section, both representing inner shelf sedimentary successions, indicate that denitrification and nitrogen fixation were well-balanced in the water column, consistent with stable nitrate supply (see also Ader et al., 2014). Together with the above mentioned positive δ34S signature measured in the Jiulongwan and Jiuqunao sections, this suggests that these successions were deposited in a restricted environment with low sulphate availability but a complete nitrogen cycle within a redox stratified water column.

**Doushantuo Member IV (Miaohé)**

In the Three Gorges Area, the black shales from the Miaohé Member at Jiulongwan were deposited under euxinic conditions, whereas euxinia is not indicated at Jiuqunao (Figs 4 and 5), which lies ca 30 km north-west of the Jiulongwan Formation. Intermittently euxinic conditions are also indicated within the uppermost Doushantuo Formation at Maoshi, which lies ca 530 km south-west of the Three Gorges Area in Guizhou Province, suggesting sporadic widespread euxinic conditions on the platform margins during deposition of the upper Doushantuo black shales. While the highest recorded Precambrian Mo concentrations are found at Jiulongwan (>300 ppm; Mo/TOC >180), the Jiuqunao and Maoshi sections exhibit only average shale Mo and V concentrations. In addition, sulphide isotope values are exclusively and distinctively negative in the Miaohé black shale at Jiulongwan, averaging −11.2‰, while at Jiuqunao and Maoshi, mostly positive values were found averaging +18.4‰ and +9.9‰, respectively, with similar trends in both sections (Fig. 11).

Therefore, Doushantuo Member IV (DST IV) at Jiulongwan was deposited under euxinic conditions, with seemingly unrestricted availability of sulphate and redox-sensitive trace-metals. Nitrogen isotopic signatures support stable nitrate availability and suggest that ‘normal’ marine production helped to fuel euxinia in this case. Hence, because a steady source of nitrate is required, and the present authors can envisage water column stratification, with euxinic bottom waters beneath oxic/dysoxic surface waters. Presumably, oxidized S, N and Mo originated from surface currents that flowed over the lip of the rimmed basin from the open ocean. The Miaohé Member sediments at the Jiuqunao and Maoshi sections, from the Three Gorges Area and Guizhou Province, respectively, do not exhibit any similar enrichment of redox-sensitive trace-metals, while pyrite is enriched in δ34S (Fig. 11), which suggests that the depositional environment of Jiulongwan and Maoshi could represent more restricted intra-shelf basins that remained isolated from oxygenated surface waters within the inner platform. Because the Jiulongwan and Jiuqunao sections are located close to one another, this finding implies considerable spatial redox complexity within the shelf lagoon.

The nitrogen isotopic signatures of these sections are considerably different: while δ15N values at Jiulongwan are relatively constant at ca 4‰, contrasting end-member δ15N values are found of ca 0.5‰ at Jiuqunao and 6.5‰ at Maoshi. At Jiuqunao, (de)nitrification seems to have been inhibited, possibly due to insufficient availability of nitrate which would also prevent the development of sustained euxinia. In that
case, the low $\delta^{15}$N values indicate that atmospheric nitrogen fixation fuelled productivity after nitrogen loss through organic matter burial. On the other hand, higher values at Maoshi suggest a more balanced nitrogen cycle in a redox stratified water column and the ability to, at least locally, sustain euxinic conditions (Boyle et al., 2013; Cremonese et al., 2013; Ader et al., 2014). Therefore, a gradually increasing exchange with the open ocean can be envisaged, as shown by the transition of pyrite sulphur isotopes from 20‰ to $-4$‰, in contrast to an average of 18.5‰ at Jiuquiao.

In summary, a rising eustatic sea-level would successively have allowed basins that were restricted or semi-restricted during deposition of DST II to have access to the open ocean, thus increasing the availability of redox-sensitive trace-metals to be scavenged by anoxic and particularly sulphidic bottom waters (Fig. 15). Low sulphate concentrations in the water column during the deposition of DST II were likely to be sufficiently counteracted by overall low Fe(II) concentrations at Jiuquiao, to allow episodic euxinia to develop (Poulton & Canfield, 2011). The euxinic sediments of the Miaohe Member at the Jiuquiao section are accompanied by very high Mo/TOC ratios and negative sulphide isotope values, suggesting a ready supply of redox-sensitive trace-metals and sulphate from the shallow, oxic ocean, probably overlying a sulphidic wedge (Fig. 15), as proposed by Li et al. (2010). The apparently close relationship between euxinic conditions and nitrate availability suggests that exchange with the generally oxygenated open ocean was essential to fuel euxinia during times of high productivity, potentially driven by upwelling of P-rich waters. Lower sea-level and basin restriction in these cases may have self-limited the spread of euxinia by limiting the supply.

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of fixed nitrogen from sources other than microbial nitrogen fixation.

The Baiguoyuan section

Previous studies of the Baiguoyuan section have focussed mainly on the black shale hosted Ag–V ore deposit, generally thought to be of sedimentary-diagenetic origin (Qian et al., 1995; Zhuang et al., 1995; Zhuang et al., 2013), especially the prominent peak in these intervals is not straightforward, peaks in Mo and V enrichments appear to be contemporaneous (Och et al., 1999). While TOC contents are within the same range as at Jiulongwan, the Miaohe Member at Baiguoyuan is poor in total sulphur (TS; Wallis, 2006). Not surprisingly, although Mo is still slightly enriched at Baiguoyuan with respect to average shale values, Mo concentrations do not exceed a tenth of the maximum content reached at Jiulongwan. On the other hand, V attains concentrations of more than 1 wt% at Baiguoyuan, which is about five times more than the maximum at Jiulongwan (Fig. 4; Wallis, 2006). The depositional environments at Baiguoyuan and Jiulongwan were presumably similar to suboxic trace-metal availability, but the former was deposited under an anoxic water column where euxinia never developed. Although neither sulphur nor nitrogen isotopic signatures are available for the Baiguoyuan section, it is difficult to formulate a more detailed explanation for the high V concentrations and further research on the section at Baiguoyuan is required. However, at this point these findings merely further corroborate a highly diversified physico-chemical environment on the Yangtze Platform with strongly varying trace-metal availability and redox conditions.

The Early Cambrian

Although the stratigraphic correlation between the Early Cambrian Jiuqunao and Jiijaipo successions is not straightforward, peaks in Mo and V enrichment appear to be contemporaneous (Och et al., 2013), especially the prominent peak in Mo concentrations and Mo/TOC ratios observed at the base of the Shuijingtuo Formation (Fig. 11). Maximum Mo and V concentrations are between five and ten times higher within the base of the Shuijingtuo Formation at Jiuqunao compared with Jiijaipo, but while V contents return to values close to average shale (e.g. McLennan, 2001), Mo decreases to values still significantly higher than average shale. One explanation is that euxinia only marginally developed during the deposition of the Shuijingtuo black shale at Jiijaipo but was pronounced at Jiuqunao, which is supported by iron speciation data. However, the δ34S excursion around the Yangjiahe/Shuijingtuo boundary at Jiijaipo section, where a euxinic interval is observed, does not seem to have led to the scavenging of trace-metals. The Mo enrichment only occurs above this interval, along with a significant decrease in δ34S. This indicates that higher sulphate concentrations were necessary to fuel sufficient sulphide production to draw down Mo. Because euxinic conditions are not indicated, it is likely that Mo precipitated during very early diagenesis close to the sediment–water interface. In addition, elevated sulphate concentrations probably coincided with higher molybdate concentrations in the water column.

The Mo and V profiles from the Huanglian and Longbizui sections, deposited in the basin, exhibit a peak around the Precambrian–Cambrian boundary, comparable to that observed at Jiuqunao and Jiijaipo (Figs 11 and 12). Hence, the present authors suggest that both environments experienced similar ocean chemistry, possibly due to further eustatic sea-level rise, which allowed for a connection between the deeper basin and the previously intra-shelf basins (e.g. Haq & Schutter, 2008). The predominantly high sulphur isotope values from Longbizui, however, may suggest a certain degree of sulphate limitation in the water column. Minimum δ34S values correlate with euxinic sediments and indicate that increases in oceanic sulphate concentrations may have intensified the production of H2S. A prominent geochemical feature at the Xiaotan section, which was deposited further inside the Yangtze Platform, is the possibility that the water column might have experienced oxic conditions above the last analysed black shale layer during deposition of the Yuanshan Formation (Fig. 13; Och et al., 2013). Moreover, δ15N stabilizes at values ca 2.5‰, recovering from values between 0‰ and 2‰, found within the anoxic-ferruginous interval (Cremone et al., 2013).

Although merely based on scarce data, the nitrogen isotope signatures at Jiijaipo, Jiuqunao and Longbizui exhibit their lowest values within the black shales deposited under euxinic conditions, which is in accordance with the model of Boyle et al. (2013), who hypothesized that euxinia should correlate with nitrogen fixation and low, atmospheric nitrogen isotope values (at least in the Precambrian; cf. Godfrey et al., 2013). In the Ediacaran section discussed above (for example, the black shales of DST IV at Jiulongwan), this was not the case. It appears that the main difference between periods of euxinic
environments in the Ediacaran and the Cambrian on the Yangtze Platform is that euxinia was less widespread and pervasive during deposition of the Doushantuo Formation, when it was established in the form of a sulphidic wedge in upwelling areas, carrying over in places to distal shelf lagoon areas such as Jiulongwan (Li et al., 2010). During the Early Cambrian, on the other hand, eunoxic conditions have been reported from both basin and platform sections (Canfield et al., 2008; Och et al., 2013, and references therein). At Jiuqunao and Jijiapo (Figs 9 and 10), after the positive peak in Mo and V, nitrogen isotopic signatures decrease to values between 0‰ and −1‰, consistent with an increase in N₂-fixation and/or photic zone anoxia, possibly facilitated by high concentrations of nitrogenase co-factors (Mo and V) (e.g. Canfield et al., 2010b; Zhang et al., 2014); N₂-fixation could then have contributed significantly to the increase in primary productivity, sustaining anoxia and sulphidic conditions. These features can also be observed at Jiuqunao and Jijiapo, although to a different extent due to the different palaeogeographic settings. Nevertheless, Mo and V peaks seem to be widespread on the Yangtze Platform, and the increase in bio-available Mo and V, in a dominantly ferruginous ocean might have triggered unprecedented primary production and organic matter delivery to the ocean floor (Anbar & Knoll, 2002), which created conditions favourable for intensified sulphate reduction and the emergence of widespread and sustained euxinia in the Early Cambrian, prior to the Cambrian bioradiation.

The effects on ecosystems of changing spatial redox distribution

It has been established that many extant animals are facultative anaerobes and so are not wholly dependent on oxygen (Martin & Müller, 2007; Budd, 2008); some are even able to withstand sulphidic conditions throughout their life cycle (Danovaro et al., 2010). Nonetheless, motility and other muscular activities of animals are repressed at low oxygen levels. Therefore, the finding of widespread anoxia and spatially extensive euxinia across the Ediacaran–Cambrian boundary seems to be incompatible with the ongoing Cambrian explosion, which is commonly believed to have been triggered by oceanic, and possibly atmospheric, oxygenation. One consequence of such spatial variability in redox conditions would be to create dynamically changing benthic ecosystems. Although preliminary, the present results may help to shed light on the distribution of benthic and, to a lesser extent, pelagic ecosystems on the Yangtze Platform. The most famous Ediacaran fronds are indisputably benthic but have only been reported in China from the Shibantan Member of the Dengying Formation, which is constrained by Fe speciation and rare earth element (REE)
distributions to have been deposited under largely oxygenated conditions (Ling et al., 2013; Duda et al., 2014); this is supported by examples of both body and trace fossils (Chen et al., 2013). As shown above, more distal equivalents such as the Liuchapo Formation, which are largely devoid of any diagnostic fossils, were deposited under anoxic and in some cases even euxinic conditions, thus restricting benthic ecosystems at this time to the generally shallower, inner Yangtze Platform.

The Miaohe biota (Steiner, 1994; Ding et al., 1996) is widely known but generally limited to just a few sections from the inner shelf lagoons close to the Jiulongwan section (the Miaohe section is ca 1 km distant and lithologically very similar) and Maoshi section (Guizhou Province). No Miaohe fossil assemblage has been reported from the equivalent horizon at the Jiulongwan section, which according to the present data experienced persistent sulphidic conditions. One possible interpretation of this is that dissolved sulphide was toxic to the Miaohe biota.

CONCLUSIONS

The multi-proxy approach, including the redox-sensitive trace-metals Mo and V, iron speciation, sulphide and nitrogen isotopes suggest a gradually increasing number of intra-shelf basins on the Yangtze Platform gaining access to the greater availability of trace-metals and sulphate present in the open ocean (Fig. 16). This effect can best be illustrated at the sections around Jiulongwan, where sea-level rise submerged the previously restricted rimmed basin during deposition of the Miaohe Member of the Doushantu Formation. There, Mo/TOC ratios suggest good communication with the open ocean, while the combination of Fe speciation and S isotopes suggests that euxinic conditions did not limit sulphate (or nitrate) availability. Although it is likely that the trace-metal and sulphate inventory increased gradually during the Precambrian–Cambrian transition, the present authors suggest that changing sea-level played an important role in the expression of the Neoproterozoic Oxygenation Event (NOE) at continental margins. However, although the change in biogeochemical cycling across this large transitional time interval follows a clear trend, the details rely on the current stratigraphic framework that may change with additional age constraints, and the signature of changing ocean chemistry and bathymetry cannot clearly be separated at present.

The apparent switch in nitrogen cycling, whereby δ15N values remained heavy during sulphidic periods in the Ediacaran, but with light values during euxinia in the Early Cambrian, possibly arises from sulphidic conditions being confined to sulphidic wedges in the upwelling areas during the Ediacaran. In contrast, productivity and subsequent sulphate reduction was probably fuelled by the nitrate reservoir of the open ocean during the early Cambrian. In the Early Cambrian, more widespread euxinic conditions across flooded shelves precluded incomplete denitrification, leading to lower δ15N values. A highly dynamic environment, both bathymetrically and chemically, including a trend to higher oxygen concentrations in the ocean interrupted by regional euxinic events, might have spurred on the emergence and diversification of animals for which the fossil record of the Cambrian Explosion can only represent a glimpse.

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**Supporting Information**

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

- **Table S1.** Elemental and isotopic data.
- **Table S2.** Iron Speciation.
- **Table S3.** Xiaotan section, Yunnan Province (see Och et al., 2013 for more details).