The aftermath of the CPE and the Carnian–Norian transition in northwestern Sichuan Basin, South China

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Abstract: The northwestern Sichuan Basin (South China) was a portion of eastern Tethys where, during the Late Triassic, a sharp lithological transition from oolitic–bioclastic limestones and sponge reef mounds to dark grey terrigenous clays, siltstones and sandstones is visible in several localities. The timing and significance of this major facies transition is still unclear. Here we report new biostatigraphic and carbon stable isotope data from Hanwang and Jushui, northwestern Sichuan Basin. Sporomorphs, ammonoids, conodonts and halobiid bivalves show that the lithological change is Late Carnian to Early Norian in age. This amended age determination facilitates recalibration of the magnetostratigraphy in the area allowing correlation between the Late Triassic of the Sichuan Basin and the Astrochronostratigraphic Polarity Time Scale. A carbon stable isotopic perturbation across the Carnian–Norian Boundary (CNB) is missing in our sections, or it is concealed because of the mixed organic matter sources. Our findings pinpoint the position of the CNB to a short stratigraphic interval of c. 12 m thickness in the Sichuan Basin. The studied sections greatly extend the palaeogeographical documentation of the CNB and provide novel information on biostatigraphy and chemostatigraphy that should be considered in defining the best position of the Norian Global Stratotype Section and Point.

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At least two main episodes of biotic crisis and environmental change are documented during the Late Triassic, the Carnian Pluvial Episode (CPE) (Simms & Ruffell 1989, 2018) and the end-Triassic mass extinction, which is one of ‘big five’ mass extinctions in the Phanerozoic (Raup & Sepkoski 1982). Both are associated with major negative δ13C perturbations (carbon isotope excursions; CIEs) (Whiteside et al. 2010; Dal Corso et al. 2018, and references therein). The CIE at the Carnian Pluvial Episode has been so far documented in a few bulk organic carbon and δ–alkane records (Dal Corso et al. 2012, 2015, 2018; Mueller et al. 2015; Sun et al. 2016; Miller et al. 2017; Baranyi et al. 2018; Shi et al. 2018) and also reported in carbonate rocks (Sun et al. 2016, 2018), and the carbon isotope record of the end-Triassic mass extinction has been extensively studied (e.g. Hesselbo et al. 2002; Schoene et al. 2010; Whiteside et al. 2010). In both cases, the biotic and environmental changes were possibly related to emissions of CO2 as a consequence of the eruption of a large igneous province (Furin et al. 2006; Schoene et al. 2010; Whiteside et al. 2010; Dal Corso et al. 2012, 2015, 2018; Sun et al. 2016, 2018).

The stratigraphic interval after the CPE is relatively poorly studied, and the definition of the boundaries between the three Upper Triassic stages (Carnian, Norian and Rhaetian) is still a matter of debate (Lucas 2018). Two main Global Stratotype Section and Point (GSSP) candidate sections of the Carnian–Norian Boundary (CNB) have been studied and proposed for their biostatigraphy and carbon stable isotopic stratigraphy: Black Bear Ridge in British Columbia (e.g. Orchard et al. 2001; McRoberts 2007, 2011; Orchard 2007, 2014; Willford et al. 2007; Zonneveld et al. 2010; Onoue et al. 2016) and Pizzo Mondello in Sicily (e.g. Muttoni et al. 2001; Nicora et al. 2007; Mazza et al. 2010, 2012, 2018; Balini et al. 2012). Magnetostratigraphy is available from Pizzo Mondello (Muttoni et al. 2004), and it has been suggested that the Norian GSSP should be located at the first occurrence (FO) of the conodont Metapolygnathus parvus (Mazza et al. 2018; Rigo et al. 2018). Halobia faunas (bivalves) at Pizzo Mondello were studied and correlated to ammonoid biozones by Levera (2012), showing that Halobia australica is the best halobiid marker for defining the base of the Norian GSSP.

We here document stratigraphic sections that are younger than the Ma’antang section (Shi et al. 2018) including the aftermath of the CPE and CNB interval from the Longmen Shan area in the northwestern Sichuan Basin, a region far from the two proposed GSSP localities of Black Bear Ridge and Pizzo Mondello (BBR and PMS in Fig. 1a).

The Sichuan Basin hosts a thick succession of Upper Triassic rocks, from Carnian ramp carbonates to Carnian–Norian marginal marine and continental siliciclastic rocks (Li et al. 2003; Wu 2009). Their sequence stratigraphic correlation has been discussed by He et al. (2011) and Mei & Liu (2017), mainly combining outcrop with core data. Fragmentary biostratigraphic data include ammonoids and bivalves (Wu 1989; Wang 1992; Shi et al. 2017), foraminifera (He 1980), conodonts (Shi et al. 2017; Jin et al. 2018) and palynofloras (Li & Wang 2016; Li et al. 2016, 2017a). The CNB in the area has so far been positioned on the basis of these scarce biostratigraphic data and is still poorly constrained. Recently, Li et al. (2017b) investigated the magnetostratigraphy of an overlying siliciclastic succession and constrained its age from Norian to Rhaetian. Zhang et al. (2015) and Shi et al. (2017) provided magnetostratigraphic and biostratigraphic data from the Qingyan Gou section in the Sichuan Basin and considered the sharp
lithological transition from carbonates to shales as related to the CPE. However, Jin et al. (2018) reported a Tuvalian 3 (late Carnian) conodont association from the carbonate units of the Qingyan Gou section, which would exclude the presence of the CPE. In an effort to better constrain this stratigraphic interval, new ammonoids, bivalves, conodonts and palynomorphs were collected and determined from the sections considered in this study. Carbon isotopic composition of organic matter (OM) was analysed on bulk-rock and wood, to assess the stable isotope record in this stratigraphic interval in the Sichuan Basin and compare it with that in other regions. Although not coincident with major shifts in the isotope record or major extinction events, the CNB is placed in a time when important biotic changes were occurring; for instance, the major differentiation of conifers (Miller 1982; Renner 2009). Therefore, investigations aimed at better characterizing this interval are needed. Results presented in the present paper go in this direction and clarify the Upper Triassic stratigraphic framework of the northwestern Sichuan Basin, allowing better correlations with sections in the western Tethys. Our findings help shed light on a relatively neglected interval of geological time and may provide a contribution to the process of identifying a GSSP for the Norian.

**Geological setting**

The northwestern Sichuan Basin is a portion of the Longmen Shan foreland basin that now crops out as part of the Longmen Shan thrust belt, at the boundary between the western Sichuan Basin and Songpan–Ganzi fold belt (Fig. 1b). It was surrounded by emerged land during the Late Triassic and was connected to the Paleotethys through a large bay (Fig. 1c; Deng et al. 1982; Wu 1984). At the end of the Middle Triassic, collision between the North China and South China plates resulted in an extensive unconformity (Zhang et al.
1996; Li et al. 2003, 2011a) with the addition of NE-striking, NW-dipping extensional faults with strike-slip components (Li et al. 1995). This unconformity marks the transition from the Paleozoic–Middle Triassic passive continental margin to the Longmen Shan foreland basin (Li et al. 2003, 2011a). In the early Carnian, a transgression connected the coastline of the Ganzi open marine region toward the western Sichuan Basin as far as Chengdu (Deng et al. 1982; Wu 1989) (Fig. 1c), and a carbonate ramp, dipping west and surrounded by locally uplifted areas, formed in the middle to western Sichuan Basin (Wu 2009; Li et al. 2011b, 2014). The Triassic stratigraphy in this region includes thick marine carbonates (c. 350 m in thickness) from the Ladinian (Middle Triassic) to Carnian (Late Triassic), and Carnian to Norian siliciclastic rocks (Wu 1989; Li et al. 2003; Mei & Liu 2017; Shi et al. 2017; Jin et al. 2018).

Three sections have been studied in this area. The Qinyang Gou section (HWQ) (31°27′46.85″N, 104°09′35.40″E), which is 9 km, and the Guanyin Ya section (HWG) (31°28′18.4″N, 104°08′50.4″E), which is around 1.5 km, are situated in the Hanwang region. The third section is Jushui (JS) (31°30′44.6″N, 104°13′54.5″E) near the town of Jushui (Fig. 1c). All of these sections were previously studied for their lithostratigraphy and microfacies (Wu 2009; Jin et al. 2015, 2018; Shi et al. 2017), biostatigraphy (Wu 1989; Wang 1992; Shi et al. 2017; Jin et al. 2018) and palaeomagnetic stratigraphy (Zhang et al. 2015).

The Middle Triassic in the northwestern Sichuan Basin

The uppermost Middle Triassic stratigraphic unit in the NW of the Sichuan Basin is the Tianjingshan Formation (T₃ₕ), which is mainly composed of stromatolitic–bioclastic limestones or dolostones. The presence of limestone coexisting with dolostone is considered a lithological marker to identify T₃ₕ. Such a lithology is markedly different from the overlying Carnian rocks, which comprise oolitic–bioclastic limestones and overlying hexactinellid sponge-bearing limestones (Wu 1989; Wendt 2001). The depositional environment of T₃ₕ is interpreted as an evaporitic tidal flat (Wu 1989, Wu 1989) assigned T₃ₕ to the Ladinian (Middle Triassic) on the basis of bivalve and brachiopod faunas. However, He (1980) reported a Carnian foraminiferal association including Diplolitmina sp., Turritellita sp., Ophthalmidium exiguum, Eoguttulina sp., Palaeomiliolina tibetica, Involutina sp., P. tenus, Aulostorus sp., Gaudryina sp. and Nodosariae (Astacolus, Frondlicularia, Lenticulina, Nodosaria, etc.), in the upper portion of the unit. Furthermore, ammonoids (Thisbites sp.) and conodonts (Neogondolella polygnathiformis, N. navicula) from the study by Wang & Dai (1981) suggest an upper Carnian age, but those fossils were not illustrated and are more typical of the overlying Ma’antang Formation (T₃ₘ) (e.g. Wang 1992; Shi et al. 2017). The top of the Tianjingshan Formation is marked by an extensive, basin-scale unconformity surface within the stromatolitic limestone (Li et al. 2003, 2011a). This unconformity is clearly seen in the lowest part of the HWG section (Fig. 2).

The Upper Triassic (Carnian and Norian) in the northwestern Sichuan Basin

The Tianjingshan Formation is overlain by the Ma’antang Formation (T₃ₘ) of Carnian age (Wu 1989; Wang 1992). The Carnian succession consists of two parts (Li et al. 2003, 2014). In our studied sections, the lower part is c. 55 m thick, and it begins with oolitic–bioclastic limestones, followed by hexactinellid sponge-bearing limestones and sponge mounds in an upward-deepening succession (Jin et al. 2018). Its sedimentary environment comprises a carbonate ramp (Wu 2009; Jin et al. 2018). The upper part is c. 65 m thick and is composed of dark grey shales, mudstones and siltstones (Li et al. 2003). Its sedimentary environment is controversial, having been interpreted as a gulf (Wu 1989), foredeep (Li et al. 2014) and lagoon (Shi et al. 2015). In this paper we use the facies model of Li et al. (2014) (Fig. 2). The abrupt lithological change from limestone to shale has been associated with the CPE (Shi et al. 2009, 2017; Wang et al. 2015). On the basis of conodont biostratigraphy, Jin et al. (2018) suggested instead that the whole T₃ₘ transition in the Hanwang area is Tuvallian (late Carnian) and the carbonate crisis in this region is thus unrelated to the CPE.

The T₃ₘ is subdivided into four units (Shi et al. 2017; Jin et al. 2018): Unit 1 comprises oolitic limestones; Unit 2 consists of bioclastic limestones; Unit 3 is mainly composed of microbial sponge mounds; Unit 4 consists of siltstones and shales (Fig. 2). The oolitic–bioclastic units (Unit 1 and Unit 2 in Fig. 2) in the Hanwang area have been assigned to the lower Carnian (Julian), based on ammonoids Protrachyceras cf. victoria, Chionites sp., Sandlingites sp. and Californites? sp. (Wu 1989) and conodont Paragondolella (= Quadratella) polygnathiformis. Recently, Jin et al. (2018) suggested that the conodont associations from the HWQ section consisted of Carnepigondolella angulata, Hayasihella tuvaluica, Paragondolella noah, Carnepigondolella orchardi, Epigondolella miettoi and Carnepigondolella cf. orchardi. These are typical late Tuvallian (late Carnian) conodonts (Maenza et al. 2012; Rigo et al. 2012, 2018). Wang (1992) reported ammonoids above the sponge mounds (lower part of Unit 4) in the Hanwang area that were determined as Thisbites borelli, T. haushoferi, T. cf. borni, T. agricolus, Anatomites herici, Discotropites plinii, Timmottropites dubvoasus, Anathibites sp., Thibites sp. and Javavites sp. In particular, the ammonoid Discotropites plinii defines the uppermost late Carnian (Tuvallian 3) biozone (Krystyn 1982).

The overlying thick succession (c. 500 m thick in the studied area) of coarser terrigenous rocks is the Xiaotangzi Formation (Fig. 2). Li et al. (2003) attributed this stratigraphic interval to the Xujiahe Formation (T₃ₜ). In the present study, we use the term ‘Xujiahe Formation’ only for the upper, coarse-grained siliciclastic unit within the Late Triassic of the Sichuan Basin.

The Xiaotangzi Formation (T₃ₜ) is dated as Norian based on ammonoids (Wu 1989) and bivalves (Gou 1998). The uppermost T₃ₜ is characterized by quartz-lithic arenites with calcite cement. Its sedimentary environment is thought to vary from shelf to littoral (Mei & Liu 2017). To date, the biostratigraphy has been poorly determined across the CNB interval. Some researchers have proposed the occurrence of coarse quartzose sandstone as a marker of the CNB (Deng et al. 1982; Li et al. 2003; Shi et al. 2015). Zhang et al. (2013) suggested that a local unconformity exists between the T₃ₘ and T₃ₜ, and this unconformity could correspond to the CNB (Fig. 2). In some localities, such as the Emei area (Fig. 1c), the upper part of T₃ₜ consists of shales and siltstones, and this unit is locally named the Kuahongdong Formation (Wu 1989). Li & Wang (2016) investigated the palynological assemblages in the western Sichuan Basin, including the area of this study. Those researchers recognized three assemblages: a Dicotyphellitidæ–Corollisporitidæ–Michrystràdium (DCM) assemblage mainly occurring in the Ma’antang Formation, a Dicotyphellitidæ–Kertosinopporidæ–Canalizonopporidæ (DKC) assemblage mainly occurring in the Ma’antang Formation and a Dicotyphellitidæ–Lunzisporitidæ–Chasmatoportidæ (LDC) assemblage that is the main association in the Xiaotangzi and Xujiahe formations.

In summary, only sparse biostratigraphic data from the Ma’antang, Xiaotangzi and Xujiahe formations are available, which suggests that the Carnian and Norian are both present in the studied area.

Magnetostratigraphy in the HWQ section, Hanwang area

The HWQ section has been also investigated for its magnetic polarity pattern (Zhang et al. 2015) and tentative correlations with
the Triassic Astrochronostratigraphic Polarity Time Scale (APTS) have been proposed (Zhang et al. 2015; Shi et al. 2017). The latest proposed correlation implies that the HWQ section spans the lower to upper Carnian with a short interruption owing to a hiatus, which was ascribed to a karstic surface, in the middle of our Unit 3 (corresponding to the middle of Unit 2 in the HWQ section of Shi et al. 2017). The above-cited researchers supposed that the late Julian is missing, on the basis of their magnetostratigraphy. However, no petrographic and field evidence of karst was found by Jin et al. (2018).

Methods

Ammonoids, bivalves and conodonts

Ammonoids were collected from the HWQ and JS sections (A–H horizons in Fig. 3). Tens of bivalves were collected from the HWQ and JS sections in 2012 (Ba and Bb horizons in Fig. 3), but only a few of them were illustrated by Shi et al. (2017). Collected specimens were cleaned and photographed at the Department of Geosciences, Padova University. Two rock samples, 3–5 kg each, were also collected from the lowest part of the JS section for conodont biostratigraphy (positions a and b in Fig. 3). The conodont samples were etched with formic acid, and the residues were dried at the Institute of Sedimentary Geology, Chengdu University of Technology. Conodonts were concentrated by heavy liquid treatment, separated manually by binocular microscope and photographed with a CamScan MX 2500 SEM in the Department of Geosciences, Padova University.

Palynology

Three palynological samples (PA–PC) were collected in the HWG and one (PD) in the HWQ section (Fig. 3). The samples were ground and c. 10 g of powder was treated with 10% HCl to remove the sulphates and carbonates, followed by 47% HF at 55°C, to remove silicate minerals. Residues were washed with deionized water until neutrality was reached, and were then sieved at 15 µm. The residue was stored in glycerin jelly. All the slides were studied and photographed in the Geoscience Department of Padova University.

Organic carbon isotopes

Seventy-six bulk-rock samples and 30 wood fragments from the HWQ section, 51 bulk-rock samples from the HWG section, and 40 bulk-rock samples and five wood fragments from the JS section were collected to be analysed for their δ13C. (From now on, the notation δ13Corg will be used for the stable isotopic composition of organic carbon.) Bulk-rock samples were first cleaned with deionized water, then oven-dried at 50°C for one night and pulverized using an agate mortar. Subsequently, the carbonates
were removed by putting c. 2 g of powder into a polypropylene Falcon tube with 50 ml of 10% HCl. Desulphurization was carried out for sulphurous samples with 37% HCl poured into the same tubes at 55°C for 4 h. The process was repeated until the solution lost its yellow colour. The residuals were rinsed with deionized water and centrifuged until neutrality was reached, then oven-dried again. Approximately 1–5 mg of sample were weighed into tin capsules. Wood samples were separated from bulk-rock, etched with 10% HCl and oven-dried. Approximately 0.04 mg of treated wood was weighted into tin capsules. All samples were run in duplicate. The δ¹³Corg analyses were performed in the Department of Geosciences, University of Padova, with a Thermo Flash 2000 Elemental Analyzer linked to a Thermo-Delta VAdvantage isotopic ratio mass spectrometer. Results were calibrated to the international VPDB scale, by analysing contemporaneously two international standards CH-7 (−32.151‰) and CH-6 (−10.449‰). A quality control standard was also run along with the samples (ZER sucrose from a C3 plant). The accuracy was c. ±0.15‰ (1σ) during the period of the analyses.

Results

Biostratigraphy

Conodonts

Three conodonts were found in two samples from the JS section (Fig. 3). One conodont species, Paragondolella polygnathiformis (Fig. 4A), occurs in a sample within the top of stromatolite bed. In sample b (Fig. 3), above the stromatolite, the conodont species Paragondolella praelindae (Fig. 4B) was found. All conodonts show a Colour Alteration Index (CAI) of 1.

Ammonoids

Ammonoids were collected from five horizons in units 3 and 4 of the HWQ section (A–E in Fig. 3) and three horizons (F–H) of the JS section (Fig. 3). Ammonoids occur along with a diverse fauna that includes brachiopods and molluscs, including thin-shelled halobiid bivalves. Ammonoids are sometimes preserved with their original aragonitic shell, sometimes shells are replaced by calcite and suture lines are seldom visible. The distribution of all ammonoids from the HWQ and JS sections is displayed in Table 1. Most specimens belong to a small (<2 cm) taxon of ammonoids with a keeled rounded venter and ventrolateral shoulders, with a moderately open umbilicus of <30% of the shell diameter. These ammonoids are densely ribbed (Fig. 5B1, B2, G1 and H), showing 10–12 ribs per quadrant on the ventrolateral margin that terminate before the keel on faint bullae or swellings. This taxon corresponds to ‘Discotropitid (n. gen. n. sp. 1)’ of Shi et al. (2017) and is here identified as ‘Tropitid n. gen. n. sp.’ Some specimens expose the suture line (Fig. 5G1), which is ammonitic, confirming the attribution of this taxon to the tropitid family. Tropitid n. gen. n. sp. is by far the most abundant ammonoid in horizons A, B, C and D of the HWQ section, and F, G and H of the JS section.
Involuted, smooth ammonoids were also collected that are attributed to the family Juvavitidae. A specimen from horizon A at HWQ (Fig. 5A1) is c. 3 cm in diameter and is preserved as a compressed replaced shell. The venter and suture lines are not visible; however, weak ribs are visible in the outer part of the flank that disappear in the umbilical part. This specimen was determined as *Gonionotites* sp. Several specimens were collected of an ammonoid taxon with smooth, involute shell and an elliptical whorl section that we tentatively attribute to the genus *Griesbachites* (e.g. Fig. 5A2, F1 and F2). These have a rounded venter without ornamentation, and flanks with few coarse and simple ribs or plicae that near the venter are adorally projected and terminate without bullae or knots. One specimen from the JS section exhibits the suture line partially (Fig. 5F1) at a diameter of c. 2 cm. This suture line has at least three lobes and three saddles, all deeply denticulated and squared, the first lobe being deeper than all others. This suture line and the weak ornamentation compare well with that of *Griesbachites laevis* illustrated by Tozer (1994), in which the ventrolateral bullae that are typical in large specimens of *Griesbachites* are instead missing or minimally developed. The ‘Juvavitid, n. gen.’ of Shi et al. (2017) most probably belongs to this taxon.

A fragment of a strongly ribbed, involuted ammonoid, preserved as an external mould, was collected from horizon E of the HWQ section (Fig. 5E1). This specimen compares well with the inner whorls of *Guembelites philostrati* illustrated by Krystyn (1982, plate 13, fig. 6) from the Jomsom section in the Nepalese Himalaya. From the same horizon, we also collected a fragmented specimen of *Thisbites* (Fig. 5E2). This ammonoid shows a dense ribbing, with 17 ribs being counted per quadrant at a diameter of c. 1 cm. Ribs are falcoid, mostly simple, a few are bifurcated close to the umbilical margin, and the ribs terminate on strong tubercles alongside a keeled venter. This specimen can be clearly distinguished from *Discotropitid n. gen. n. sp.* 1 because of the denser ribs, which terminate on strong tubercles instead of the weak bullate swellings of *Tropitid n. gen. n. sp.* Because of the dense ribbing and strong tubercles, this *Thisbites* has been determined as *Thisbites cf. petralis* (see Tozer 1994, plate 108, figs 3 and 4).

Finally, a nearly complete specimen of an ammonoid with distinctive ornamentation comes from horizon G of the JS section (Fig. 5G2). This specimen, c. 1.5 cm in diameter, has a narrow, keeled venter bordered by bullated tubercles on the ventrolateral shoulders. The umbilicus is well open and exposes ribbed inner whors. The last whorl, probably a habitation chamber, is not ribbed, and shows only very weak tubercles elongated radially that mimic faint ribs. Of the suture line, only part of an entire saddle is visible. We attribute this ammonoid to a species of *Hadrothisbites* different from *H. taylori* of Tozer (1994).

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Table 1. Late Triassic ammonoids in northwestern Sichuan Basin, China

<table>
<thead>
<tr>
<th>Ammonoid species</th>
<th>Horizon</th>
<th>Occurrence</th>
<th>Section</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tropitid n. gen. n. sp.</td>
<td>A</td>
<td>XX</td>
<td>HWQ</td>
</tr>
<tr>
<td>Tropitid n. gen. n. sp.</td>
<td>B</td>
<td>XX</td>
<td>HWQ</td>
</tr>
<tr>
<td>Tropitid n. gen. n. sp.</td>
<td>C</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Tropitid n. gen. n. sp.</td>
<td>D</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Tropitid n. gen. n. sp.</td>
<td>F</td>
<td>X</td>
<td>JS</td>
</tr>
<tr>
<td>Tropitid n. gen. n. sp.</td>
<td>G</td>
<td>XX</td>
<td>JS</td>
</tr>
<tr>
<td>Tropitid n. gen. n. sp.</td>
<td>H</td>
<td>XX</td>
<td>JS</td>
</tr>
<tr>
<td>Gonionotites sp.</td>
<td>A</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Griesbachites sp.</td>
<td>A</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Griesbachites cf. laevis</td>
<td>F</td>
<td>X</td>
<td>JS</td>
</tr>
<tr>
<td>Guembelites philostrati</td>
<td>E</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Thisbites cf. petralis</td>
<td>E</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Hadrothisbites n. sp.</td>
<td>G</td>
<td>X</td>
<td>JS</td>
</tr>
</tbody>
</table>

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Fig. 4. Scanning electron microscope photomicrographs of conodonts from the Jushui section. (a) *Paragondolella polygnathiformis*; (b) *Paragondolella praetinaidae*. For each specimen: (1) upper view, (2) lower view, (3) lateral view. ‘a’ and ‘b’ refer to the sampling horizons (see Fig. 3). CAI=1 All scale bars represent 200 μm.

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Fig. 5. Examples of ammonoids from the Jushui and Jomsom sections. (A) *Tropitid n. gen. n. sp.* 1 from horizon A of the HWQ section. (B) *Tropitid n. gen. n. sp.* 2 from horizon A of the JS section. (C) *Tropitid n. gen. n. sp.* 3 from horizon A of the JS section. (D) *Tropitid n. gen. n. sp.* 4 from horizon A of the JS section. (E) *Thisbites cf. petralis* from horizon E of the HWQ section. (F) *Thisbites cf. petralis* from horizon E of the HWQ section. (G) *Hadrothisbites n. sp.* from horizon G of the JS section.
The association of tropitids, *Hadrothisbites* n. sp., *Gonionotites* sp. and *Griesbachites* sp. at HWQ and JS indicates a late Tuvalian (late Carnian) age. In particular, tropitids are known so far only from the upper Carnian. It should be noted, however, that the tropitids illustrated in the present paper are a new taxon, whose distribution is thus unknown. The ammonoids from horizon E of the HWQ section suggest instead an early Norian age: *Guembelites philostrati* is so far known only from the Norian, whereas the species of genus *Thisbites* occur in a short stratigraphic interval across the Carnian–Norian boundary; *T. petralis*, however, is known only from the lower Norian (Tozer 1994).

**Bivalves**

Halobiid bivalves occur at several levels (Ba and Bb) within both the HWQ and JS sections (Fig. 3). Most of the specimens are preserved as partial skeletons of recrystallized calcite, which expose the outer surface of a middle skeletal layer. Although this surface
(between the shell layers) does preserve some of the original ornament of the individuals (e.g. commarginal growth banding and at least the general course and direction of the radial ornament), details of the external expression of radial ribs from the outer skeletal layer are obscured, precluding definitive species-level identification.

Based on shell outlines, commarginal growth banding, the angular breadth of the anterior tube and generalized course of the ribbing of the specimens, it appears highly probable that they belong to a single species, here referred to as Halobia cf. septentrionalis Smith (Fig. 6), and may therefore may serve as a robust correlation datum between the two sections. The HWQ and JS specimens are similar to those that have been attributed to both Halobia qaboensis Zhang and Halobia septentrionalis Smith. There is a strong possibility of these two taxa being synonymous. Chen & Ba (1986, plate 2, figs 6, 8, 11 and 14) illustrated Halobia septentrionalis Smith 1927 from the middle to late Carnian of Zedang (south Xizang). This species is known from several localities throughout North America, and is perhaps best represented in southeastern Alaska and western Canada in the upper Carnian (K. macrolobatus ammonoid zone) as discussed by McRoberts (2010, 2011). This morphotype is also similar to Halobia qaboensis Zhang (of Zhang et al. 1985, plate 33, figs 1–9) known from the Bolila Formation, near Kangba and Chengdu. However, the preservation of Zhang’s illustrated specimens is rather poor, making direct comparisons with H. septentrionalis not possible. Additionally, the HWQ and JS halobias may be compared with those Ma et al. 1976 (plate 33, fig. 6) referred to as Halobia rugosa Gümbel from the Yunnanyi Formation, Xiangyun County, Yunnan. More recent evaluations of Halobia rugosa (see McRoberts 2011; Levera 2012) suggest that ‘H. rugosa’ from the late Carnian are distinctively different from true Halobia rugosa, which is earliest Carnian (=Trachyceras ammonoid zone) from the western Tethys. Therefore, it is possible that the Yunnanyi ‘H. rugosa’ are also conspecific with those described as Halobia cf. septentrionalis in this paper (Fig. 6).

The morphological affinities of the HWQ and JS Halobia cf. septentrionalis suggest a Tuvalian age and are most probably a mid- to upper Tuvalian equivalent to the Anatropites (= K. macrolobatus) or slightly older T. subulatus (= T. welleri) ammonoid zones (sensu Silberling & Tozer 1968; Kozur 2003; McRoberts 2010).

Spores and pollen

The sporomorphs were collected from PA–PC and PD positions from the HWG and HWQ sections (Fig. 3), respectively. The preservation of the palynological material is unfortunately not good, because of degradation on the exine. Representative sporomorphs are shown in Figure 7, and the distribution of all sporomorphs is shown in Table 2.

The spores from the HWQ section are composed of levigated and ornamented Azonotriletes Deitiodospora sp., Concavisporites spp., Dicryophyllidites sp., Dicyophyllidites harrisi, Todisporites sp., Punctatisporites spp., Converrucosporites spp., Furtucosiropites sp., Pustulatisporites sp. and Leiotriletes sp.; cingulati spores Annulispora sp., Stereisporites sp. and Assevertospora sp.; zonate spore Kraeusselisporites sp.; cavatomoletes Aratrisporites spp.; and a cingulicavate form Denosporites sp. Pollen grains are rare, including alete forms Psophosphaero sp. and Araucariae sp.; and monocolpate forms Riccispores sp., Cyadopites sp. and Chasmatisporites spp. Bisaccates Piceites sp. and Pinuspollenites enodatus, and green algae spores Tasmanites sp. are present.

The spores from the HWQ section consist of Azonotriletes Concavisporites sp., Converrucosporites sp. and Anapiculatisporites sp.; zonate spore Kraeusselisporites sp.; and cavatomoletes Aratrisporites sp. The pollen grains contain the monocolpate

Fig. 6. Photographs of Halobia cf. septentrionalis Smith from the HWQ (Ba) and JS (Bb1–3) sections. Ba, articulated valve pair; Bb1, left valve; Bb2, right valve; Bb3, left valve. All scale bars represent 1 cm.
Cydopites sp. and Chasmatosporites spp., bisaccates Pinuspollenites sp. and Alisporites sp.; and algae spores Botryococcus sp. and Tasmanites sp.

**δ¹³C of bulk organic matter from woods and bulk-rock**

The δ¹³Corg values at the HWG section range from −29.12 to −24.39‰, those from the HWQ section range between −27.47 and −24.29‰, and those from the JS section vary from −27.47 to −23.90‰. In the HWQ section, 30 wood fragments were collected, and their values range from −25.35 to −22.37‰. The values of five wood samples from the JS section range between −24.29 and −22.07‰. The δ¹³C data from limestone, terrigenous rocks and wood are distinct (Fig. 8).

The mean value of δ¹³Corg in limestones from the three sections is −26.6‰, whereas terrigenous rocks of the Ma’antang Formation...
have an average δ¹³Corg value of −24.8‰. The wood fragments of the HWQ and JS sections are −24.0‰ on average: more positive than but largely overlapping with the δ¹³Corg in terrigenous rocks (Fig. 8). All sections exhibit a weak positive trend in limestone units 1–3 (Fig. 2), and very negative values of about −29‰ occur at the stromatolite in the basal part of the HWQ section above the unconformity surface (dashed circle in Fig. 2).

In the HWQ section, two positive isotopic excursions (PIE1 and PIE2) are observed. PIE1 is within the limestone unit, whereas PIE2 coincides with the lithological change from limestone to shale and siltstone. PIE2 can be observed in a comparable position also at the stromatolite in the basal part of the HWG section above the unconformity surface (Fig. 2).

Table 2. Late Triassic palynomorphs in the Hanwang area, northwestern Sichuan Basin

<table>
<thead>
<tr>
<th>Palynomorph</th>
<th>Samples</th>
<th>Occurrence</th>
<th>Section</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alisporites sp.</td>
<td>PD</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Anapiculatisporites sp.</td>
<td>PD</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Annulispora sp.</td>
<td>PA</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Aratrupisporites spp.</td>
<td>PA, PB, PD</td>
<td>XX</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Aracruariacites sp.</td>
<td>PA, PB, PC</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Asseretoispora sp.</td>
<td>PB</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Bisaccates</td>
<td>PA, PB</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Botyrococcus sp.</td>
<td>PA</td>
<td>X</td>
<td>HWQ</td>
</tr>
<tr>
<td>Chaumatopisporites sp.</td>
<td>PB, PC, PD</td>
<td>XX</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Concavisporites sp.</td>
<td>PA, PB, PC, PD</td>
<td>X</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Converrucosisporites sp.</td>
<td>PA, PD</td>
<td>XX</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Cyadogites sp.</td>
<td>PA, PB, PD</td>
<td>XX</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Detitoidospore sp.</td>
<td>PC</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Densoisporites sp.</td>
<td>PA</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Dictyophyllidites harrissii</td>
<td>PB</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Dictyophyllidites sp.</td>
<td>PC</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Kraeuselisporites sp.</td>
<td>PA, PB, PD</td>
<td>X</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Leioiricites sp.</td>
<td>PA</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Pecetea sp.</td>
<td>PA</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Pinuspollinites enodatus</td>
<td>PB, PD</td>
<td>X</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Psophoaphaera spp.</td>
<td>PA, PB</td>
<td>XX</td>
<td>HWG</td>
</tr>
<tr>
<td>Punctatisporites spp.</td>
<td>PA, PB, PC</td>
<td>XX</td>
<td>HWG</td>
</tr>
<tr>
<td>Pustulatisporites sp.</td>
<td>PC</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Riccisporites sp.</td>
<td>PC</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Stereisporites sp.</td>
<td>PB</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Tasmanites sp.</td>
<td>PA, PD</td>
<td>X</td>
<td>HWG, HWQ</td>
</tr>
<tr>
<td>Todisporites sp.</td>
<td>PB</td>
<td>X</td>
<td>HWG</td>
</tr>
<tr>
<td>Verrucosisporites sp.</td>
<td>PC</td>
<td>X</td>
<td>HWG</td>
</tr>
</tbody>
</table>

Shi et al. (2017) illustrated an ammonoid association from the JS section that is similar to the one illustrated in the present paper (Fig. 3). The ammonoids illustrated by Shi et al. (2017) were collected on a large but isolated outcrop on the eastern valley flank. However, their stratigraphic position could be determined with good approximation by tracing beds in the field (Fig. 2). These ammonoids include both smooth-shelled juvavids and abundant tropitids; Shi et al. (2017) interpreted this ammonoid association as representing the lower Tuvalian. Because of a richer ammonoid association presented herein, and an improved determination of smooth-shelled taxa, we are now confident that this ammonoid association instead indicates a late Tuvalian (late Carnian) to early Norian age. This ammonoid biochronology is consistent with our age determination of halobid bivalves and conodonts.

The palynological associations also suggest a late Carnian–Norian age. Pollen and spores from the PA horizon in the HWQ section and the PD horizon are similar to the DKC assemblage of Li & Wang (2016) from the Kuahongdong Formation dated to late Carnian–early Norian. Pollen and spores from the PB horizon show the appearance of some important taxa such as Psophoaphaera and Riccisporites, which are similar to the DLC assemblage of Li & Wang (2016) from the Xiaotangzi and Xujiahe formations, which those researchers assumed to be Norian–Rhaetian. Moreover, the distinctive palynological associations described in the western Tethys during the Carnian Pluvial Episode (e.g. Roghi 2004; Roghi et al. 2010; Mueller et al. 2016) seem to be absent, further confirming that the stratigraphic succession in Hanwang is younger than the CPE.

**The amended age of stromatolitic beds of the uppermost Tianjingshan Formation**

The conodonts Paragondolella polygonaformis (Fig. 4a) and Paragondolella praelindae (Fig. 4b) were found within or just above the stromatolitic beds of the JS section (Fig. 3). This interval
was attributed to the uppermost Tianjinshan Formation and was dated to the Middle Triassic (Wu 1989). However, *P. polygnathiformis* and *P. praelindae* are exclusive to the Julian (lower Carnian) to lower Tuvalian (upper Carnian; e.g. Mazza et al. 2012). *P. polygnathiformis* first appears in the Carnian GSSP section of Stuores Wiesen, very close to the Ladinian–Carnian boundary (Mietto et al. 2012), and is routinely used as a datum to define the Carnian. Therefore, the uppermost Tianjinshan Formation at the JS section should be assigned a Carnian age. This age determination would agree with those of, for example, He (1980) and Wang & Dai (1981).

A revised magnetostratigraphic correlation of the HWQ section

The most recent magnetic polarity timescale for the late Triassic (Kent et al. 2017) is mostly based on data from the Newark Basin, correlated with the biostratigraphy of western Tethys. For the Tuvalian to Ladinian (i.e. late Carnian to early Norian) interval a biostatigraphically calibrated record of magnetic polarity reversals has been investigated in the Pizzo Mondello section (Sicily; Mutti et al. 2001, 2004; Mazza et al. 2012, 2018).

The new findings of conodonts, ammonoids and halobiid bivalves at HWQ allow an amended magnetostratigraphic correlation with western Tethys and the Newark polarity timescale. However, biostatigraphic age determinations are not fully in agreement with each other at HWQ. This led us to provide two correlation proposals, each having its advantages and drawbacks.

At Pizzo Mondello, among many other taxa, the platform conodont elements *Hayashiella tuvalica, Carnepigondolella zoae, C. angulata* and *Paragondolella noah* disappear at the top of the PM3r magnetozone, whereas the co-occurrence of *Epigondolella miettoi* and *Carnepigondolella orchardi* has a narrow distribution in the magnetozone PM4n, immediately below the proposed CNB defined by the FO of *Metapolygnathus parvus* (Mazza et al. 2018; Rigo et al. 2018) (Fig. 9). The conodonts *Epigondolella miettoi* and *Carnepigondolella orchardi* were found by Jin et al. (2018) in the HWQ section within HQ2n, below horizons with the upper Tuvalian to lower Ladinian ammonoids. This implies that the whole magnetostratigraphic record at the HWQ section is younger than suggested by Zhang et al. (2015) and Shi et al. (2017). In the HWQ section, Shi et al. (2017) identified a karstic surface (i.e. hiatus) within HQ2r. It was suggested that the record of geomagnetic reversals at HWQ is thus discontinuous, and the late Julian is truncated (fig. 10 of Shi et al. 2017). However, Jin et al. (2018) did not find any detectable omission or erosive surface, which could correspond to a hiatus, in this interval. In light of the conodont biostratigraphy, and assuming near-constant sedimentation rates, the magnetozones in the HWQ section can be correlated to the magnetozones at Pizzo Mondello (Option A in Fig. 9). Assuming near-constant sedimentation rates means that the thickness ratios of magnetozones at HWQ are similar to those of the Newark magnetic polarity timescale.

The Option A correlation implies that the ammonoid associations of horizons A–D, which contain tropitids and *Halobia cf. septentrionalis*, are Norian. The CNB interval is to be placed between 6 and 9.5 m from the base of the HWQ section. However, this correlation implies low sedimentation rates through the section, generally <10 m Ma⁻¹, which seem to be unlikely in a foreland
basin system and more typical of starved open ocean environments below the carbonate compensation depth (e.g. Enos 1991).

An alternative correlation is proposed that accounts for increasing sedimentation rates at the HWQ section that could be expected in a foreland basin (Option B in Fig. 10). The Option B correlation maintains that ammonoid horizons with tropitids and Halobia cf. septentrionalis are within the Carnian, but requires that the thickness of magnetozones at HWQ is not proportional to their duration, and these thicknesses do not correlate to those of magnetozones at Pizzo Mondello. The Carnian Ma’antang Formation is the initial sedimentation stage of the Longmen Shan foreland basin (Li et al. 2003, 2014). The deep marine shales and mudstones of the upper Ma’antang Formation (Fig. 2) contain deep-water fossils, suggesting that the basin was underfilled at this stage (Li et al. 2003, 2014), but then, with the siltstones and sandstones of the Xiaotangzi Formation, sedimentation was keeping up with subsidence. Along with the development of the foreland basin, the sedimentation rates were increasing, and on the basis of the upper Triassic thickness of Li et al. (2003) and the geological timescale of Cohen et al. (2013), we calculated average net sedimentation rates in the studied area of c. 25 to c. 50 m Ma$^{-1}$ in the Carnian, c. 162 m Ma$^{-1}$ in the Norian and >139 m Ma$^{-1}$ in the Rhaetian. This suggests that sedimentation rates were increasing near the CNB in the Sichuan Basin. All calculations of net sedimentation rates do not consider compaction and denudation rates, hence the original accumulation rates may have been higher. In view of ammonoids, bivalves and sedimentation rate, the reverse polarity interval PM4r at Pizzo Mondello may correlate to the

Fig. 9. Correlation of the HWQ section with the Newark Astrochronostratigraphic Polarity Time Scale (APTS) (Kent et al. 2017) and Pizzo Mondello, Sicily (Muttoni et al. 2004), based on conodont biostratigraphy and magnetostratigraphy (Zhang et al. 2015; Jin et al. 2018). The Carnian–Norian boundary (CNB) interval (in grey) is placed between 6 and 9.5 m. According to this proposal, the CNB (in grey) coincides with the boundary interval of Mazza et al. (2012); that is, the interval between the FO of Metapolygnathus parvus (conodont) and the FO of Halobia austriaca (bivalve) at Pizzo Mondello. This boundary interval was then correlated assuming that HQ2r above the conodonts at the HWQ section coincides with PM4r at Pizzo Mondello. With this correlation, the bivalve Halobia cf. septentrionalis would occur in the Norian at HWQ.
interval between HQ3r and the top of HQ4r at the HWQ section. According to Option B, the CNB interval at the HWQ section could be positioned between 14.5 and 26.5 m. The drawback of this correlation (Option B) is that the thickness ratios of magnetozones are not preserved, and as a result the correlation is outside of the range of biostratigraphic constraints. Further biostratigraphic investigations are needed to finally assess which of the offered correlation options is better.

The sources of OM in the Sichuan Basin

The $\delta^{13}$C$_{org}$ in sedimentary rocks can result from the combination of OM from diverse sources, in particular marine or freshwater–terrestrial (Fogel & Cifuentes 1993). A possible variability of OM sources in the investigated sections is demonstrated in Figure 8, where the average carbon isotopic compositions of different components from the Sichuan Basin are compared with those of coeval localities worldwide, where the provenance of OM could be suggested on the basis of the sedimentary environments. The $\delta^{13}$C$_{org}$ of sandstone across the CNB interval is more positive than samples from coeval deep marine settings of Black Bear Ridge, but overlaps with that of wood samples from both the Sichuan Basin and the Southern Alps. In contrast, the average $\delta^{13}$C$_{org}$ of limestones in the Sichuan Basin, which bear little terrigenous material and no wood, is closer to the carbon isotopic composition of the open marine Black Bear Ridge section. These different $\delta^{13}$C$_{org}$ values could be easily explained if we assume that OM in limestones is mostly marine in origin, whereas the wood-bearing sandstones contain mostly terrestrial organic matter.

Recent marine OM, which is mainly derived from phytoplankton and benthic and macroscopic algae (Fry et al. 1977), has a $\delta^{13}$C$_{org}$ less negative than $-23.0\%$ (Fry et al. 1977; Anawar et al. 2010). However, the terrestrial OM is more $\delta^{13}$C-enriched than marine OM during the Triassic (Hayes et al. 1999; Nordt et al. 2016). OM remineralization, metabolic fractionation and diagenesis could alter the isotopic composition of OM (e.g. McArthur et al. 1992; Goñi et al. 1997; Anawar et al. 2010), but the impact of diagenetic factors should not be important in our case, because we consider only relative changes within bulk-rock and wood, which were affected by similar diagenetic influences in each study area.
Wood fragments from the HWQ and JS sections have more positive δ¹³Corg than marine OM (Fig. 8), within a wide range from −25.35 to −22.07‰. Although wood from the Dolomites has a slightly more positive δ¹³Corg (Dal Corso et al. 2011), this could be ascribed to an undetected difference in taxonomic composition. Triassic land plants were C3 plants, and today C3 plants have an average δ¹³Corg of c. −26‰. Even though the average δ¹³Corg in the Longmen Shan area is −24.0‰, slightly more positive than present values, it is still within the range of what would be expected for C3 plants (Gröcke 2002; Badeck et al. 2005, and references therein). The wide range of δ¹³Corg from wood fragments reflects a natural dispersion that is expected for plants (e.g. Arens et al. 2000; Dal Corso et al. 2011). Owing to its intrinsic variability, an erratic trend is observed in the δ¹³Corg of wood of the HWQ section (Fig. 2).

The mean value of δ¹³Corg in terrigenous rocks (Unit 4 in Fig. 2) is between those of wood and bulk OM of the limestone units (Fig. 8), albeit closer to the δ¹³Corg of wood. These foredeep-deposited terrigenous rocks have been suggested to be related to the uplift of Longmen Mountains and other surrounding palaeolands (Yang et al. 2008; Liu et al. 2009; Shi et al. 2015), thus the δ¹³Corg of bulk organic carbon in this unit may reflect the mixing of C3 plant material with marine OM. The different OM source with distinctive ranges of δ¹³Corg value from limestone and terrigenous units may explain PIE2 (Fig. 2).

Differences in organic carbon sources may also explain other minor features of the δ¹³Corg record that could not be correlated between the three study sections. In the HWQ section, the well-bedded sponge mound layers are inter-reef deposits that correspond to the mound-shaped sponge reefs in the JS and HWG sections (Wu 1989). A slight negative perturbation of c. 1.5‰ in magnitude (CIE1 in Fig. 2) is recorded within the inter-reef deposits, which is however, not reproduced in the other two sections of this study. We suggest that this local isotopic shift may be related to accumulation of marine OM in this inter-reef environment rather than to a regional or global modification of the organic carbon pools.

Is there a carbon isotopic oscillation at the CNB?

In our correlation Option A (Fig. 11a), a strong isotopic excursion of δ¹³Ccarb nearly coincides with the CNB. However, δ¹³Ccarb from carefully chosen brachiopod shells shows no detectable excursion. This has been explained as the result of early diagenesis of carbonate sediments rich in organic carbon during shallow burial (Jin et al. 2012).
The δ¹³C_carb record from brachiopods is considered to represent a more pristine record of seawater δ¹³C than bulk carbonate (Brand 1989; Brand et al. 2003; Korte et al. 2005). Although a 1‰ magnitude shift is still found in brachiopod data near the CNB (Fig. 11a), this may actually derive from the natural range of isotopic variability of brachiopod shells (Korte et al. 2017). A PIE in the δ¹³C_carb across the CNB interval has been reported in many locations (Black Bear Ridge, Onoue et al. 2016; Guri Zi, Muttoni et al. 2014; Pizzo Mondello, Muttoni et al. 2004; Mazza et al. 2010; Wadi Mayhah, Sun et al. 2018) (only the BBR and PMS sections are displayed in Fig. 11a). This PIE has different shapes and magnitudes and has been reported only from a deep-water setting (Fig. 11a).

According to correlation Option B (Fig. 11b), a minor negative shift (CIE2) of c. 1‰ is observed at 15 m, at the base of the CNB interval, which is difficult to correlate with Black Bear Ridge because biostratigraphic constraints in this part are poor. Moreover, this negative shift is found in an interval of facies transition characterized by the increase in terrigenous input and therefore from the CNB interval in South China. The only counterpart with detailed biostratigraphy is Black Bear Ridge, Canada (Williford et al. 2007). A small negative isotopic perturbation, c. 0.5‰ in magnitude, was found at the proposed position of the CNB at Black Bear Ridge and was discussed in relation to a fossil turnover observed in the section (Williford et al. 2007). In our correlation Option A (Fig. 11a), an opposite shift (PIE1) to more positive values c. 2‰ in magnitude occurs in the δ¹³C_org across the CNB, which is part of a general positive trend within the limestone unit, reproduced in all study sections (Fig. 2).
might be related to the mixing of OM from continental and marine sources. As a consequence, the general flat trend of $\delta^{13}$C$_{org}$ values within the CNB interval (Fig. 11b, grey band) could have recorded a mixed-carbon stable isotopic composition, and the real signal of $\delta^{13}$C$_{org}$ across the CNB, which could have been masked in our sections.

The end of the Carnian Pluvial Episode in the northwestern Sichuan Basin

The most negative values of $\delta^{13}$C$_{org}$ are recorded in the lowest part of the HWQ section, followed by a long-lasting positive trend (Fig. 2). As this trend is detected within the limestone units (Units 1–3), we assume that the OM source was not changing too much in this interval. The $\delta^{13}$C$_{org}$ values of limestones in the Sichuan Basin overlap with those during the fading phase of the CPE from the adjacent Nanpanjiang Basin (Fig. 8). We tentatively correlate the carbon isotope records between these two areas (Fig. 12).

The biostratigraphic of the HWQ section is still incomplete. The lower Ma’antang Formation yielding most negative $\delta^{13}$C$_{org}$ values might correspond to approximately the 34 m level in the Long Chang section of Sun et al. (2016) and thus to the fading phase of the carbon isotopic excursion (CIE) of the Carnian Pluvial Episode. This trend was recognized clearly in the Ma’antang section, northwestern Sichuan Basin (Shi et al. 2018), and it is identifiable in the basal part of our three sections (Fig. 2). The distinctive CIE was also reported in other parts of Tethys (Dal Corso et al. 2012, 2015, 2018; Mueller et al. 2015, 2016; Miller et al. 2017; Baranyi et al. 2018; Sun et al. 2018) and it is partly missing at the HWQ section because it corresponds to a hiatus marked by the basal karstic surface (Figs 2 and 12), which has been interpreted as the forebulge unconformity of the Longmen Shan foreland basin (Li et al. 2003, 2011a). In the eastern Sichuan Basin, the whole Tianjinshan Formation and the overlying Ma’antang Formation were eroded or were never deposited (Li et al. 2003; Mei & Liu 2017).

Conclusions

New biostratigraphic data from the Carnian and Norian of the northwestern Sichuan Basin are provided in this paper. On the basis of an existing magnetostatigraphy and new biostratigraphic age determinations, two possible correlations between the Astrochronostratigraphic Polarity Time Scale of the Triassic and the magnetostatigraphy of the HWQ section are proposed, but further biostratigraphic investigations are needed to identify a preferred correlation. In Option A correlation, the CNB is located between 6 and 9.5 m. In Option B correlation, the CNB could be positioned between 14.5 and 26.5 m of the HWQ section. Furthermore, the occurrence of Carnian conodonts at the base of the JS section implies that some stromatolitic dolostones–limestones in the Sichuan Basin are Carnian rather than Ladinian.

Carbon stable isotopes from wood and bulk organic matter are here measured for the first time in eastern Tethys near the CNB. The recorded variability of $\delta^{13}$C$_{org}$ between different lithologies is explained by varying organic matter sources. A long-lasting positive trend has been recognized within a uniformly calcareous interval, which can be attributed to the slow recovery from the isotopic perturbations of the CPE and can be correlated with the isotopic record of the Nanpanjiang Basin of South China. Instead, the isotopic record of the CPE in the western Tethys and continental Europe is more complex, featuring repeated negative isotopic excursions, and its correlation with the Sichuan Basin is still uncertain.

Overall, our findings identify the northwestern Sichuan Basin as a relevant area for the study of the CNB. In particular, Carnian–Norian ammonoids, conodonts and bivalves from the Sichuan Basin are from a much more easterly position than other successions so far considered for the definition of the CNB. These new biostratigraphic data may thus prove useful for a better-informed placement of the Norian GSSP.

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