Diversified calcimicrobes in dendrolites of the Zhangxia Formation, Miaolingian Series (Middle Cambrian) of the North China craton

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Abstract

As a type of non-laminated microbial carbonates, dendrolites are dominated by isolated dendritic clusters of calcimicrobes and are distinct from stromatolites and thrombolites. The dendrolites in the upper part of the Miaolingian Zhangxia Formation at Anjiazhuang section in Feicheng city of Shandong Province, China, provide an excellent example for further understanding of both growth pattern and forming mechanism of dendrolites. These dendrolites are featured by sedimentary fabrics and composition of calcified microbes as follows. (1) The strata of massive limestones, composed of dendrolites with thickness of more than one hundred meters, intergrade with thick-bedded to massive leiolites, forming the upper part of a third-order depositional sequence that constitutes a forced regressive systems tract. (2) A centimeter-sized bush-like fabric (shrub) typically produced by calcified microbes is similar to the mesoclot in thrombolites but distinctive from clotted fabrics of thrombolites. This bush-like fabric is actually constituted by diversified calcified microbes like the modern shrub as a result of gliding mobility of filamentous cyanobacteria. Such forms traditionally include: the Epiphyton group (which actually has uncertain biological affinity), the Hedstroemia group which closely resembles modern rivulariacean cyanobacteria, and the possible calcified cyanobacteria of the Lithocodium–Bacinella group. (3) Significantly, dense micrite of leiolite is associated with sponge fossils and burrows, and is covered by microstromatolite. The Lithocodium–Bacinella group is a controversial group of interpreted calcified cyanobacteria in the Cambrian that has also been widely observed and described in the Mesozoic. Therefore, dendrolites with symbiosis of leiolites in the studied section provide an extraordinary example for further understanding of growing style of bush-like fabrics (shrubs) of the dendrolites dominated by cyanobacterial mats. Furthermore, the present research provides some useful thinking approaches for better understanding of the history of the Early Paleozoic skeletal reefs and the microbe–metazoan transitions of the Cambrian.

Keywords: Calcified microbe, Dendrolite, Leiolite, Epiphyton, Lithocodium–Bacinella, Cambrian, Miaolingian, Zhangxia Formation

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1 Introduction
Although microbialites (Burne and Moore 1987) are formed by the action or influence of microbes, they do not always preserve direct, or diagenetically robust, evidence for their mode of formation, thus, they are believed to be among the most intractable sedimentary rocks to study (Bosence et al. 2015). Microbial carbonates (calcified microbialites) are the spectacular examples of “microbially-induced sedimentary structures” (MISSs; Noffke and Awramik 2013). Microbial carbonates have been classified into four types on the basis of microstructure, i.e. stromatolites marked by laminated structures, thrombolites associated with clotted fabrics, leiolites commonly expressed as structureless, and dendrolites characterized by dendritic fabrics (Burne and Moore 1987; Riding 1991a, 2000, 2011a).

Among a variety of microbial sedimentary structures in carbonates, dendrolite is characterized by the following particularities and complexities (Riding 1991a). Firstly, the dendrolites have a non-laminated microstructure with relatively large (cm scale) clots dominated by isolated dendritic clusters of calcimicrobes (mainly grouped into Epiphyton; Riding 1991a, 2000; Howell et al. 2011). Dendrolites are commonly confused with thrombolites due to similar clotted fabrics (Chen et al. 2014a, 2014b; Gong 2016; Yan et al. 2017), and defined as dendritic thrombolite (Qi et al. 2014). Epiphyton framstone (Han et al. 2009), or Epiphyton buildup (Woo et al. 2008; Woo and Chough 2010). Secondly, similar to the modern dendrolitic cone (Bradley et al. 2017) or dendrolitic microbial mat (Suosaari et al. 2018), the dendrolites of the Cambrian are dominated by diversified filamentous microbes. Thirdly, it seems that dendrolites have a distinctive distribution during the Cambrian Miaolingian (Riding 2000; Rowland and Shapiro 2002). The reason for the special temporal distribution is not still thoroughly understood. Fourthly, the biological affinity of the main calcimicrobes belonging to the Epiphyton group in the Cambrian dendrolites remains uncertain (Riding 1991b, 1991c, 2011b; Luchinina and Terleev 2008, 2014; Luchinina 2009).

The common calcimicrobes belong to the Epiphyton group, the Hedstroemia group, and the Lithocodium–Bacinella group. Although results of several researchers demonstrated that biological affinity of fossils of the Epiphyton group belongs to the cyanobacteria (e.g. Zhang et al. 1985; Jiang and Sha 1996; Laval et al. 2000; Woo et al. 2008; Han et al. 2009; Woo and Chough 2010; Chen et al. 2014a, 2014b; Säsänen et al. 2014; Gong 2016; Ezaki et al. 2017), some researchers did not include it in cyanobacteria (Luchinina and Terleev 2008, 2014; Luchinina 2009; Adachi et al. 2014, 2015; Liu et al. 2016). There are numerous arguments about the affiliation of calcified cyanobacteria with the Lithocodium–Bacinella group (Cherchi and Schroeder 2006; Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013). The sedimentary fabrics of the studied Cambrian microbial reefs show similar characteristics to the Lithocodium that either formed through decomposition of sponges (Kwon et al. 2012; Chen et al. 2014a, 2014b; Lee et al. 2014a, 2014b, 2015; Li et al. 2015; Park et al. 2015; Coulson and Brand 2016; Shen and Neuweiler 2018; Cordie et al. 2019) or from sponge fibers of keratose demosponges (Luo and Reitner 2014, 2016; Coulson and Brand 2016; Lee and Riding 2018; Shen and Neuweiler 2018; Lee et al. 2019). These arguments enhance the sophistication of the studied Middle Cambrian dendrolites at the Anjiazhuang section that provides an example for further understanding of the growth pattern and forming mechanism of calcified microbes. Furthermore, the establishment of the Lithocodium or Lithocodium-like fabric in the Cambrian dendrolites on the North China craton may provide some useful thinking approaches and research clues for further research of the early history of the Paleozoic skeletal reefs (Rowland and Shapiro 2002; Kiessling 2009; Lee et al. 2015, 2019; Ezaki et al. 2017; Lee and Riding 2018; Cordie et al. 2019) and the microbe–metazoan transitions (Chen et al. 2019).

2 Geological setting
Phanerozoic sedimentation in the North China craton began in Cambrian Epoch 2 times. The Cambrian sediments unconformably overlie the Precambrian sediments in a similar style as described in North America (Peters and Gaines 2012). Consequently, approximately 700 m thick Cambrian strata were deposited on the North China Platform, consisting of a mixed succession of red beds and carbonate rocks from late part of Epoch 2 to early Miaolingian, a carbonate succession dominated by oolitic grainstones during the main period of the Miaolingian, and a succession of carbonate rocks predominated by carbonate mud during the Furongian (Wang et al. 1989; Meng et al. 1997; Shi et al. 1997; Feng et al. 2004; Mei 2011a; Ma et al. 2017; Latif et al. 2018; Riaz et al. 2019a, 2019b).

The Miaolingian Series in the North China craton comprises Maozhuang, Xuzhuang, Zhangxia and Gushan formations (e.g. Peng and Zhao 2018), and can be subdivided into four third-order depositional sequences (Ma et al. 2017; Latif et al. 2018; Riaz et al. 2019a, 2019b). The oolitic-grain bank that developed during the forced regression (Fig. 1) laid the foundation of ramp-type carbonate platform (see Pratt et al. 2012). It is characterized by shallow shelf ooidal shoals, flanked seawards by a ramp and landwards by tidal flats (Fig. 2a) (e.g. Feng et al. 1990, 2004; Meng et al. 1997). Moreover, abundant microbial reefs (Riding 2002a; Rowland and Shapiro 2002; Kiessling 2009; Chen et al. 2014a, 2014b; Lee et al.
dominated by microbial carbonates were developed in the shallow shelf covered by ooidal shoals (Fig. 2a), which are similar to "algal flats" (Jiang and Sha 1996; Sha and Jiang 1998).

Similar to other regions of the North China craton, the Zhangxia Formation at the Anjiazhuang section in the western part of Shandong Province consists of a third-order sequence (DS3: Fig. 1) that is further subdivided into three subsequences (I to III in Fig. 3). The top and bottom of each subsequence is marked by drowning unconformities (e.g. Schlager 1989, 1999; Gómez and Fernández-López 1994; Mei 1996, 2010). These boundaries depict rapid transgressions by directly overlying

Fig. 1 Diagram depicts the sedimentary succession of the Cambrian in North China craton. Biostratigraphic succession of trilobites is adapted from Peng (2009). The scheme-1 for the lithostratigraphic division is adapted from Lu et al. (1994), Meng et al. (1997) and Feng et al. (1990, 2004). Scheme-2 is from Xiang et al. (1999). Scheme-3 is used in present paper, and is modified from scheme-1 on the basis of sequence-stratigraphic division according to succession of sedimentary facies. Nine third-order sequences (DS1 to DS9) are discerned in the Cambrian System in the North China craton, which are divided by two types of sequence boundaries, i.e. the exposure punctuated surface (SB) and the drowning unconformity surface (arrows). The gray rectangle marks the target strata in present study.
shelf mudstones on high-energy oolitic grainstones with the intergrowth of massive bioherms comprising microbial carbonates. The strata in the upper part of subsequence III of the Zhangxia Formation at the studied section, recording the relative sea-level fall, can be distinguished into three parts (b, c, and e in Fig. 3a): (1) the lower part is marked by a set of oncolitic- and oolitic-grainstones with thickness of about 15 m (Fig. 3b); (2) the middle part is constituted by a set of massive oolitic grainstones with approximately thickness is 30 m (Fig. 3c); (3) the upper part is about 125 m thick and comprises of massive biothermal limestones dominated by dendrolites with the intergrowth of leiolites (Fig. 3e), which is the key target of this study. These are remarkable deposits especially for the diversified cyanobacteria composition and the heterogeneous sedimentary fabrics in dendrolites with the intergrowth of leiolites, which further fit the concept of microbial reefs (Rowland and Shapiro 2002; Kiessling 2009; Lee et al. 2014a, 2014b, 2015, 2019; Lee and Riding 2018; Cordie et al. 2019).

Fig. 2 a Sedimentary pattern for the late depositional period of the Cambrian Miaolingian Zhangxia Formation in the North China craton (adapted from Feng et al. 1990, 2004); b Geographical outline of Shandong Province with the section location; c Regionally geological feature in the southern part of Feicheng city with the section location
3 Materials and methods

Dendrolites with the intergrowth of leiolites in the massive limestones are most abundant in the upper part of subsequence III of the Miaolingian Zhangxia Formation at the Anjiazhuang section (Figs. 2, 3). The macro- and mesostructures of these massive limestones were observed on pale weathered surfaces (Fig. 3e). Detailed observations were conducted on the well-exposed outcrops, where 118 samples were collected from massive limestones, oolitic grainstone and oncolitic grainstone. Microfabrics were observed in thin sections with a Zeiss Axio Scope A1 microscope. High-resolution images were made for dendrolites and leiolites under the microscope.

4 Results

4.1 Fundamental features of the dendrolite

4.1.1 Macroscopic feature

The term dendrolite was firstly introduced by Riding (1991a). It has a distinctive form different from the laminated stromatolite and clotted thrombolite (Riding 1991a, 2000). Typical samples of dendrolites were collected from the Miaolingian Zhangxia Formation at the Anjiazhuang section in the western part of Shandong Province (Fig. 2b).

Macroscopically, the dendrolites of the Zhangxia Formation at the Anjiazhuang section are composed of large-scale (cm-sized) and small-scale (mm-sized) dark...
clots of dense micrites (Fig. 4). These clots verify the initial definition of thrombolite (Aitken 1967) that led to an important distinction between thrombolites and stromatolites (Kennard and James 1986). These special clots are bush-like (shrubs), and are made up of isolated dendritic clusters of calcimicrobes (including *Epiphtont*), which occur in the following forms: (1) irregular bodies of semi-connected or isolated shape (Fig. 4a), (2) relatively regular bodies of large-scale fan-like or chamber-like shape (cm-sized; Fig. 4b), and (3) small-scale (mm-sized) regular or irregular isolated bodies of chamber-like shape (Fig. 4c).

### 4.1.2 Microscopic feature

Diversified calcimicrobe composition and heterogeneous sedimentary fabrics in dendrolites of the Miaolingian Zhangxia Formation are shown in Figs. 5, 6, 7, 8, 9, 10. The bush-like fabrics (or shrubs) of the dendrolites in mm- to cm-level size are made up of isolated dendritic clusters of calcimicrobes (Figs. 5, 6, 7, 8, 9, 10). Microscopically, these calcimicrobes belong to fossils of the *Epiphyton* group (Riding 1991b, 1991c, 2000; Luchinina and Terleev 2008, 2014; Woo et al. 2008; Han et al. 2009; Luchinina 2009; Woo and Chough 2010; Yan et al. 2017) with lengths up to 500 μm and diameters from 50 μm to 70 μm, and depict the structure of bifurcating branches of dense micritic rods. These typical fossils of the *Epiphyton* commonly form cm-size or even larger shrub thalli (Fig. 5a), fan-like thalli (Fig. 5b), and isolated chamber-like thalli (Fig. 5c). The *Epiphyton* along with *Korilophyton* ("K" in Fig. 5c, d; solid micrite, but with short branches; Riding 1991b, 1991c) can form isolated chamber-like thalli. These thalli of calcimicrobes seem to grow in a matrix of dense micrites as well as a particular matrix of microspars that are similar to the “Bacinnella-like” fabric (Schlagintweit and Bover-Arnal 2013) or the *Bacinnella* (Cherchi and Schroeder 2006; Rameil et al. 2010).

More importantly, four special microfabrics are observed within the dendrolites (Fig. 5). Firstly, it is the possible *Lithocodium* (Cherchi and Schroeder 2006; Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013) which shows the characteristic of microtubular fabric in dark dense micrites. The dark micritic aggregations are...
possibly filamentous networks that are filled by microspars. These aphanitic microtubules have diameters of 10–20 μm (“L” in Fig. 5c, d), and form an irregular-shaped mm-leveled tubular structure with relatively clear boundaries. For microproblematica that may be calcified cyanobacteria, there is continuing discussion regarding the biological affinity of *Lithocodium* than that of *Epiphyton*. One approach is to group these forms into a special sedimentary fabric that is marked by encrusting masses of irregular open micritic networks, e.g. the *Lithocodium*-like fabric rather than a possible microbial fossil (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013). Secondly, the “Bacinella-like” fabric as shown in Fig. 5c and d is microspar separated by thin films (~1 μm thick) of dark micrite which is relatively enriched in organic substances. These typical fabrics have been elucidated as microproblematica of the *Lithocodium–Bacinella* group (Cherchi and Schroeder 2006; Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013), rather than a type of fossil calcimicrobe. Thirdly, special crystals of dolomites have the size of 10–20 μm with blurred core and clear rim (Fig. 5c). Although these crystals can be grouped into dolomitic cements (Choquette and Hiatt 2008) and/or dolomitic precipitates (Roberts et al. 2013), their genetic relationship with the dendrolites needs to be further investigated. Fourthly, the irregular-shaped mm-leveled fenestral cavities with clear boundaries (“F” in Fig. 5c, d) are observed. These fenestral cavities are filled with equant calcite spars forming granular and drusy mosaics without original pore space and obvious geopetal structures. Similar cavities have been observed in the cortex of Cambrian oncoids with abundant preserved fossils of filamentous cyanobacteria like *Girvanella* (Wilmeth et al. 2015; Riaz et al. 2020). These mm-
scale fenestral cavities (“F” in Fig. 5c, d) may provide some useful information for further understanding of the forming process, especially the calcification of dendrolites.

Fossils of the possible calcimicrobes belong to the Epi-
phyton group in large-scale shrub-like thalli (Fig. 6), and can be further subdivided into three members (Riding 1991a, 1991b), that is: (1) the typical Epiphyton (solid micritic, dendritic) (“E” in Fig. 6b) consisting of bifurcating branches of dense micritic rods, 50–70 μm in diameter, which has been described as type I Epiphyton (Woo et al. 2008); (2) the Gordonophyton with segmentation (Fig. 6a, c), which is similar to typical Epiphyton in terms of outline thalli and branch dimension, and has been described as type II Epiphyton (Woo et al. 2008); and (3) the Korilophyton (“K” in Fig. 6b, d) with short branches and a large length-diameter ratio, which has been described as type III Epiphyton (Woo et al. 2008). Moreover, the special interstices within these large shrub-like thalli of Epiphyton are composed of carbonate spars that are different from the microscopic fabrics of possible Lithocodium (“B” in Fig. 5c, d) (Schlagintweit and Bover-
However, shrub-like thalli of *Epiphyton* are similar to filling carbonate spars of mm-scale fenestral cavities (“F” in Fig. 5c, d). Further, both *Epiphyton* and *Korilophyton* can develop the corresponding large-scale fan-shaped shrub-like thalli (Fig. 6b).

Moreover, possible calcimicrobe fossils of typical *Epiphyton* shape, with relatively thick bifurcating branches of dense micritic filaments, frequently vegetated into a cm-sized fan-shaped shrub-like thallus in the matrix represented by the *Bacinella*-like fabric (Fig. 6b). Photomicrographs were taken under PPL.

**Fig. 7** Photomicrographs portray large-scale fan-like thalli of typical *Epiphyton* in the dendrolites in top part of the subsequence III of the Zhangxia Formation at the Anjiazhuang section, Shandong Province. **a** A cm-sized fan-shaped shrub-like thallus vegetates within the matrix represented by the *Bacinella*-like fabric; **b** Two cm-sized fan-shaped shrub-like thalli make an upgrowth in the transitional boundary between the matrix represented by the *Bacinella*-like fabric and the matrix composed of dense micrites with development of small-scale shrub-like thalli.

Arnal 2013). However, shrub-like thalli of *Epiphyton* are similar to filling carbonate spars of mm-scale fenestral cavities (“F” in Fig. 5c, d). Further, both *Epiphyton* and *Korilophyton* can develop the corresponding large-scale fan-shaped shrub-like thalli (Fig. 6b).

Moreover, possible calcimicrobe fossils of typical *Epiphyton* shape, with relatively thick bifurcating branches of dense micritic filaments, frequently vegetated into a cm-sized fan-shaped shrub-like thallus in the matrix represented by the “*Bacinella*-like” fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013) and dense micrites. The small shrub-like thalli can grow within dense micritic matrix (Fig. 7b). Furthermore, typical *Epiphyton* form large-scale (cm-sized) chamber-shaped colony (Fig. 8a, b) in the matrix composed of “*Bacinella*-like” fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013). Several small-scale (< 1 mm) chamber-shaped colonies also vegetate in the matrix of the “*Bacinella*-like” fabric (Fig. 8c). These
fossils of possible calcimicrobes belonging to *Korilophyton* with short branches and intergrowth of typical *Epiphyton* can also form large-scale bush-like colonies in the matrix of the “Bacinella-like” fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013) (Fig. 9).

In the *Epiphyton* group, besides the above three members of typical *Epiphyton* (Figs. 5a, b, 6b, d, 7, 8), *Gordonophyton* (Fig. 6a, c), and *Korilophyton* (Figs. 6a, d, 9), *Hedstroemia* (Fig. 10), another typical fossil of calcified cyanobacteria is observed within the clotted clusters as shown in Fig. 4, which has been described as type IV *Epiphyton* (Woo et al. 2008). *Hedstroemia* is comprised of delicate tubular branched filaments characteristically arranged in fan-like radiating clusters, and is subdivided into two varieties: (1) typical *Hedstroemia* (Riding 1991b, 1991c; Woo et al. 2008; Yan et al. 2017) marked by a fan-like radiating cluster and is composed of relatively thick filaments (about 10 μm) (“H-1” in Fig. 10a); (2) the *Cayeuxia* (Riding 1991b, 1991c; Woo et al. 2008; Liu et al. 2016) that is characterized by a hemisphere-like radiating cluster, and is composed of relatively thin elongated filaments (<5 μm) radiating from a base point (“H-2” in Fig. 10a, b, c). The colonies of typical *Hedstroemia* and *Cayeuxia* intergrow with the colony of *Epiphyton* (Fig. 10b). These colonies develop a microspar matrix composed of *Bacinella*-like fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013), which further share following features: (a) branching clusters of tubes or filaments, (b) clusters with fan-like longitudinal section, (c) radial erect growth, and (d) adjacent tubes or filaments which either share common or separate walls.
4.2 Fundamental features of leiolite intergrown with dendrolite

4.2.1 Macroscopic feature

As mentioned above that isolated dendritic clusters of calcimicrobes are composed of mm- or cm-sized dark dense micrite (Fig. 4; e.g. Riding 1991a, 2000; Howell et al. 2011) which are similar to mesoclots in thrombolites (Aitken 1967), it led to the situation that dendrolite is commonly described as thrombolite (Kennard and James 1986; Woo and Chough 2010; Qi et al. 2014; Ezaki et al. 2017; Yan et al. 2017). In contrast, leiolite was regarded as a type of microbial carbonate (Braga et al. 1995) that is relatively structureless (aphanitic) with mesostructure, and lacking laminations and clots. It could be used as a synonym for “cryptomicrobialite” and could be grouped into thrombolite (Kennard and James 1986). No example of modern leiolite has been published (Dupraz et al. 2011), and only a few particular examples of ancient leiolites have been described in the stratigraphic record.

Some leiolites with intergrowth of dendrolites form massive limestones (Fig. 3d). These leiolites (Fig. 11) have no structure which fit to the definition of leiolite as described by Braga et al. (1995), Riding (1991a, 2000) and Dupraz et al. (2011). The macroscopic features of these leiolites are: (1) an abrupt boundary with dendrolites, which is a possible hardground; (2) the primary composition of dense micrites with a few small lenses or stripes.
of cherts (Fig. 11a, b); and (3) the possible heterogeneous sedimentary fabric comprising some irregular-shaped dark microclots (organic substances) and lenses of dolomite (Fig. 11c).

4.2.2 Microscopic feature
The leiolite of the Miaolingian Zhangxia Formation at the Anjiazhuang section provides distinctive characteristics of the heterogeneous sedimentary fabric (Fig. 12). As a fundamental feature of leiolite, the dense micrites (Fig. 12a) are partly rich in trilobite fossils (Fig. 12b). There are several other features of leiolite as below. (1) It shows a possible *Bacinella*-like fabric (Fig. 12c) which is similar to the fabric described in dendrolites (Figs. 5, 6, 7, 8, 9). (2) It contains occasional sponge fossils (Fig. 12d, e) with the outline of 6 mm in diameter and obvious spongoceol of about 2 mm in diameter. In term of the clear channel system and contrast to the *Lithocodium*-like fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013) (Fig. 5c, d), the skeleton texture represented by regularly- and morphologically-repeated microspars in dense micrites, was grouped into the fossils of lithified sponge (Adachi et al. 2015; Lee et al. 2016). (3) There are small-scale (< 0.5 mm) colonies of *Epiphyton* within dense micritic fabrics (Fig. 12e).

The detailed observations show that microscopic fabrics of leiolites can be grouped into the special microspar fabric that is similar to *Bacinella* or *Bacinella*-like fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013), and the dense micrites, which are associated with...
many small-scale colonies of mm-sized *Epiphyton* as well as other fossils of calcimicrobes (Fig. 13). Some possible calcimicrobes observed in dense micrites belong to *Renalcis* (Fig. 13c), which have clusters of hollow reniform bodies with a diameter of 50–100 μm, and have thick walls of micrite. Some ambiguous and non-tangled fossils of calcified filamentous cyanobacteria which have a length of several hundreds micrometers and a diameter of 10–20 μm are identified in transitional boundary between the dense micritic fabric and the “*Bacinella*-like” fabric (Fig. 13d, e). The slightly even walls (1–2 μm in thickness) of calcified fossils of filamentous cyanobacteria are composed of dark micrites and are comparable to modern *Tychonema* (De los Ríos et al. 2015). It can be grouped into a new type of fossil, i.e. the *Girvanella* group (Riding 1991b, 1991c).

Typical and tangled filamentous fossils of *Girvanella* intergrowing with small-scale (about 0.1 mm) colonies of *Epiphyton* were observed inside *Bacinella*-like fabric (Fig. 14). These filamentous fossils are several hundreds of micrometers in length, about 10 μm in diameter, and with a dark micritic wall in thickness of 1–2 μm (Fig. 14b, c). Like non-tangled fossils of calcified filamentous cyanobacteria (Fig. 13d, e), the relatively even walls of
tangled fossils may be products of in vivo calcification, which is genetically related to ecophysiological concentration of atmospheric CO₂ (Kah and Riding 2007) and is beyond the scope of present study. Furthermore, they are grouped into typical fossils of *Girvanella* and are similar to modern *Plectonema* (Riding 1991b, 1991c).

Further observation indicates that the *Lithocodium-* and *Bacinella*-like fabrics are the fundamental fabrics for the leiolite (transitional type between leiolite and dendrolite) (Fig. 15a, b). The *Lithocodium*-like fabric may intergrow with the *Bacinella*-like fabric (Fig. 15a), or isolatedly occur within the *Bacinella*-like fabric (Fig. 15b). Importantly, small (200–400 μm) chamber-like or irregular-shaped colonies of *Epiphyton* not only overgrow within the *Lithocodium*-like fabric (Fig. 15a–d) but also occur in the *Bacinella*-like fabric (Figs. 15a and 16). The phenomenon further demonstrates that the *Lithocodium-* and *Bacinella*-like fabrics can be used as growth matrix of small colonies of *Epiphyton*, which is similar to the phenomenon of the Late Jurassic *Epiphyton*-like cyanobacteria described by Săsăran et al. (2014).

5 Discussion
Microbial reefs with a thickness of more than 100 m are observed in upper part of subsequence III of the Miao-lingian Zhangxia Formation at the Anjiazhuang section.
in Feicheng city of Shandong Province (Figs. 1, 2b and 3). They are dominated by dendrolites with intergrowth of leiolites, and were formed during the forced regression period that fulfills the model of Hunt and Tucker (1992), Mei and Yang (2000), and Schlager and Warrlich (2009). The Zhangxia Formation is a third-order sequence bounded by “drowning unconformities” (e.g. Schlager 1989, 1998, 1999), which represents a macroscopically remarkable sedimentary deposit and conforms to the concept of microbial reef that is formed in a relatively high-energy marine environment (Riding 2002a; Rowland and Shapiro 2002; Kiessling 2009; Lee et al. 2014a, 2014b, 2015, 2019; Lee and Riding 2018; Cordie et al. 2019).

The dendrolites are made up of cm- and mm-sized dark dense micritic clots which are bush-like (or shrubs) (Fig. 4). These clots are macroscopically similar to the mesoclots inside the thrombolites (Aitken 1967; Kennard and James 1986). However, they have different possible calcimicrobes, i.e. the Epiphyton group, including typical Epiphyton (Figs. 5a, b, 6b, d, 7 and 8), the Gordonophyton (Fig. 6a, c), and the Korilophyton (Figs. 6b, d and 9), which make the Miaolingian dendrolites structurally distinct from the cotted thrombolites (Riding 1991a, 2000; Howell et al. 2011). Therefore, diversified calcimicrobes and complicated sedimentary fabrics of Miaolingian dendrolites at a microscopic level, are relatively more sophisticated than previously visualized.

The typical inference of calcified fossils of the Epiphyton group (Figs. 5, 6, 7, 8, 9) in cotted clusters (Fig. 4) can be an analogue to modern cyanobacteria such as Stigonema (Riding 1991b, 1991c; Laval et al. 2000; Woo et al. 2008; Woo and Chough 2010; Ezaki et al. 2017; Mei et al. 2017, 2020), which further reflect the following phenomena: (1) possible modern analogue of shrub fabric in the Miaolingian dendrolites was formed by typical filamentous cyanobacteria (Laval et al. 2000; Bradley et al. 2017); and (2) possible calcified fossils, i.e. the Gordonophyton (Fig. 6a, c), are similar to modern cyanobacteria such as Lyngbya or Tolypothrix that form hormogonia (Mohr et al. 2011; Suosaari et al. 2018). The present paper follows this typical idea. There are different thoughts for the biological affinity of fossils belonging to Epiphyton group (Figs. 5, 6, 7, 8, 9, 10, 12, 13, 14), such as: (1) evolution in life cycle similar to calcareous red algae (Luchinina and Terleev 2008, 2014; Luchinina 2009); (2) precipitation via bacteria rather than photosynthetic cyanobacteria (Chafetz and Guidry 1999); (3) a
type of microbial fossil that can not be grouped into calcified cyanobacteria like *Renalcis* (Adachi et al. 2014); and (4) calcified microproblematica (Liu et al. 2016). Furthermore, phototropism of *Epiphyton* (Woo and Chough 2010) provides relatively more reasonable explanation for their recognition through following phenomena: (a) the intergrowth of *Epiphyton* and *Hedstroemia* within the dendrolites is shown in Fig. 11, with the *Hedstroemia* analogue to modern cyanobacteria *Rivularia* (Whitton and Mateo 2012; Liu et al. 2016; Mlewski et al. 2018); and (b) the intergrowth of *Epiphyton* and *Girvanella* in leiolites is shown in Figs. 13e and 14, with *Girvanella* similar to modern cyanobacteria *Plectonema* or *Tychonema* (Riding 1991b, 1991c; De los Ríos et al. 2015). Therefore, based on cyanobacterial affinity of *Epiphyton*, it is assumed that shrubs in the Miaolingian dendrolites are constructed by filamentous cyanobacteria like the modern examples (Bradley et al. 2017; Suosaari et al. 2018).

However, two fabrics of possible calcimicrobes increase complexity for the dendrolites in this study, i.e., (1) the possible *Lithocodium* or *Lithocodium*-like fabric (Cherchi and Schroeder 2006; Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013) which comprise micritic aggregation with filamentous networks or aphanitic–microtubular fabric within dark dense micrites of the dendrolites (“L” in Fig. 5c, d) and the leiolites (“L” in Fig. 15); (2) *Bacinella* or *Bacinella*-like fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013) in the dendrolites (Figs. 5, 6, 7, 8, 9, 10) and the leiolites (Figs. 11 and 13 to 14, 15a, b and 16) which is marked by microscopic fabrics that are separated by thin films of dark micrites. Although they make up a possible group of calcified cyanobacteria, i.e. the *Lithocodium*–*Bacinella* group (Rameil et al. 2010) has been a form of codiacean algae by nomenclature (Elliott 1956) and was interpreted as a type of foraminifer (Schmid and Leinfelder 1996). Therefore, it is suggested that this fabric should be grouped into possible residue of colonies of calcified cyanobacteria (Cherchi and Schroeder 2006) or micro-encrusters.
that may suggest a phototrophic metabolism (Rameil et al. 2010). Although the Bacinella-like fabric (Schlagintweit and Bover-Arnal 2013) was interpreted as a type of algae in nomenclature (Radoičić 1959) and later interpreted as a type of green algae (Schlagintweit et al. 2010), further research suggested that the Bacinella-like fabric should be grouped into a feature of modern microbialites (e.g., microbial mats) (Schlagintweit and Bover-Arnal 2013). As a possible group of calcified cyanobacteria, the Lithocodium–Bacinella group (Riding 1991b, 1991c) has been widely described in microbial carbonates of the Mesozoic (Cherchi and Schroeder 2006; Védrine et al. 2007; Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013). Furthermore, the Lithocodium or the Lithocodium-like fabric is enigmatically similar to sedimentary fabric (Figs. 5c and 15), which has been frequently interpreted as sponge cast in recent years, i.e., (1) the network of spicules of silicious sponge (Kwon et al. 2012; Chen et al. 2014a, 2014b; Lee et al. 2014a, 2014b, 2015; Li et al. 2015; Park et al. 2015; Coulson and Brand 2016; Shen and Neuweiler 2018), and (2) the sponge fibre of non-spicular keratose demosponges (Luo and Reitner 2014, 2016; Larmagnat and Neuweiler 2015; Lee and Riding 2018; Shen and Neuweiler 2018; Lee et al. 2019). All of these increase complexity for biological affinity of the “Lithocodium-like” fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013). However, true sponge fossils composed of spicules (Fig. 12d, e) are distinctive from the “Lithocodium-like” fabric represented by microsparic microcanaliculus in the dense dark micrites (Figs. 5c, d and 15) (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013). Furthermore, it provides some useful information to interpret the Lithocodium or the Lithocodium-like fabric as a colony of calcified cyanobacteria (Cherchi and Schroeder 2006) and a useful mirror image for studies of many researchers who always interpreted it as a spongy cast in recent years (Kwon et al. 2012; Chen et al. 2014a, 2014b; Lee et al. 2014a, 2014b, 2015; Li et al. 2015; Luo and Reitner 2014, 2016; Larmagnat and Neuweiler 2015;
Park et al. 2015; Coulson and Brand 2016; Lee and Riding 2018; Shen and Neuweiler 2018; Lee et al. 2019). Furthermore, the establishment of the *Lithocodium* or *Lithocodium*-like fabric within Miaolingian dendrolites and leiolites may provide some useful thinking approaches and a research clue for further research of the history of Early Paleozoic skeletal reefs (Rowland and Shapiro 2002; Kiessling 2009; Lee et al. 2015, 2019; Ezaki et al. 2017; Lee and Riding 2018; Cordie et al. 2019), microbe-metazoan transitions of the Cambrian (Chen et al. 2019) and their driving factors (Rowland and Shapiro 2002; Kiessling 2009; Lee and Riding 2018; Riding et al. 2019). For example, the Cambrian/mid-Ordovician (541–465 Ma) is a period marked by abundance of microbial carbonate (MC) (Riding et al. 2019) and overall lack of metazoan reef-framework builders (Rowland and Shapiro 2002; Kiessling 2009; McKenzie et al. 2014; Ezaki et al. 2017; Lee and Riding 2018; Cordie et al. 2019). Further dominant lithisted sponge–microbial reefs are documented during the Miaolingian to the Furongian (Chen et al. 2014b; Lee et al. 2015, 2019; Lee and Riding 2018; Cordie et al. 2019) and the peak of metazoa are marked by maceriate (maze-like) sponge reef during the late Furongian (Chen et al. 2019), which can not support actual microbial components like *Lithocodium* or *Lithocodium*-like fabric during the Cambrian (Figs. 5c, d and 15).

On the basis of two important cognitions, i.e. the *Lithocodium* or *Lithocodium*-like fabric should be grouped into a colony of calcified cyanobacteria (Cherchi and Schroeder 2006), and a micro-encruster might suggest a phototrophic metabolism (Rameil et al. 2010), it is more reasonable to imagine that *Lithocodium*-like fabric (Figs. 5c, d and 15) should be resulted from the calcification of one type of filamentous cyanobacteria with common sheaths, i.e. the *Nostoc* (Potts 1997; Wright et al. 2005; Soule et al. 2009; Helm and Potts 2012). Although this idea is a bold interpretation that needs further investigation, some of their evidence is as follows. (1) Similar to the dark micrites forming wall of calcified cyanobacteria like the *Girvanella* (Figs. 13e and 14), it might be resulted from in vivo calcification of structured sheaths (Riding 1991b, 1991c, 2011b; Kah and Riding 2007) composed of extracellular polymeric substances (EPSs; Dupraz et al. 2009, 2011; Decho 2010, 2011; Stal 2012; Tourney and Ngwenya 2014; Decho and Gutierrez 2017) with special protein of extracellular sunscreen scytonemins. Similarly, dark dense micritic matrix which hosted the aphanitic–microtubular fabric of the *Lithocodium*-like fabric (Figs. 5c, d and 15) might result from in vivo calcification of structured common sheath of *Nostoc* that is made up of EPS along with extracellular sunscreen scytonemin (Potts 1997; Wright et al. 2005; Soule et al. 2009; Helm and Potts 2012). (2) Irregular and snaky filaments of aphanitic–microtubulars with diameter of 10–20 μm are filled by microspars within dark dense micrites (“L” in Figs. 5c, d and 15). They might be a decomposition residue of the filaments (trichomes) of *Nostoc*, which are similar to the calcified...
residue of filamentous cyanobacteria like the *Girvanella* that are mimic to the modern cyanobacteria *Plectonema* or *Tychonema* (Riding 1991b, 1991c; De los Ríos et al. 2015) (Figs. 13e and 14).

The intergrowth among the *Bacinella*-like fabric (Fig. 6c, d) (Riding 1991b, 1991c; Cherchi and Schroeder 2006; Védrine et al. 2007; Schlagintweit and Bover-Arnal 2013), *Lithocodium*-like fabric (Rameil et al. 2010; Schlagintweit and Bover-Arnal 2013), and *Epiphyton* in the dendrolites (Fig. 6c, d) and the leiolites (Figs. 15a, b and 16) clearly demonstrates that the *Lithocodium*-like fabric should be grouped into a type of microbial crusts. Finally, the *Lithocodium*-like fabric (Figs. 6c, d, 15 and 16) can be an analogue to modern *Nostoc* colonies with grape-shape or irregular morphology, and the size ranges from several tens to hundreds micrometers within microbial mats dominated by filamentous cyanobacteria (De los Ríos et al. 2015). However, there are lots of detailed problems that need further research.

Microbial reefs dominated by dendrolites with intergrowth of leiolites in the Miaolingian Zhangxia Formation at the Anjiazhuang section (Fig. 3) depicted sophisticated calcification on microbial mats comprised of cyanobacteria (Stal 2012), including: (1) the intergrowth of multiple cyanobacterial colonies in the dendrolites such as possible *Nostoc* (Potts 1997; Wright et al. 2005; Soule et al. 2009; Helm and Potts 2012) (Fig. 5c, d), the *Epiphyton* (Figs. 5, 6, 7, 8, 9, 10), and *Hedstroemia* (Fig. 10); (2) sheath of filamentous calcified cyanobacteria such as *Girvanella* within the leiolites (Figs. 13e, 14 and 16); 3) widespread *Bacinella*-like fabric in the dendrolites (Figs. 5, 6, 7, 8, 9, 10, 11) and the leiolites (Figs. 14, 15, 16) that forms a special matrix for the growth of multiple colonies of cyanobacteria. Like the modern analogies of dendrolites, which are dominated by filamentous cyanobacteria that forming dendritic microstructure (Laval et al. 2000), dendritic cone (Bradley et al. 2017), and dendrolitic shrub (Suosaari et al. 2018), the clotted clusters are composed of dark dense micrites (Fig. 4) which can be interpreted as shubs that are chiefly dominated by *Epiphyton* (Figs. 5, 6, 7, 8, 9) and *Hedstroemia* (Fig. 10). They occurred on an underlying microbial mat (Figs. 5, 6, 7, 8, 9) that is represented by *Bacinella*-like fabric, which can be grouped into a feature of modern microbialites (e.g. microbial mats) (Schlagintweit and Bover-Arnal 2013). The dense micrite may be understood as the result of precipitation of CaCO₃ morphs occurred simultaneously in metabolically active cyanobacterial layers (Kaźmierczak et al. 2015). It should be noted that widespread *Bacinella*-like fabric in the dendrolites (Figs. 5, 6, 7, 8, 9, 10, 11) and the leiolites (Figs. 14, 15, 16) are not completely similar to the typical *Bacinella*-like fabric of the Lower Aptian (Schlagintweit and Bover-Arnal 2013). However, following features can still characterize this type of fabric as an irregular vesicular microcrust that follows some of the formative characteristics of microbialites: (1) a distinctive growth matrix of multiple colonies of cyanobacteria from the dense micrites; (2) a special composition of microspars that are separated by thin films (~1 μm) of dark micrites enriched with relatively more organic substances that are distinctive from common inorganic micrites or microspars; and (3) the diversified calcified microbes especially for those fossils of calcified sheath of cyanobacteria represent by *Girvanella*, possible *Nostoc* and *Epiphyton* within this type of fabric. Further investigations required on their ultramicrofabric characteristics and microorganisms that involved in their formation, can lead to a result of true sedimentary and microbial nature of this type of enigmatic fabric.

Of course, it can not simply think that the microbial reefs in the Miaolingian Zhangxia Formation (Fig. 3) are the products directly built up by cyanobacteria. It is definitely a result of complicated microbial precipitation (Riding 2000, 2011b) or the early lithification (Dupraz et al. 2009, 2011) in the microbial mats dominated by cyanobacteria. However, a major problem is the fact that majority of present day microbial mats dominated by cyanobacteria do not form consolidated rocks (Stal 2012). Further, these dendrolites (Figs. 3, 4, 5, 6, 7, 8, 9) with intergrowth of leiolites (Figs. 11, 12, 13, 14) may be products of sophisticated calcification of EPS (Dupraz et al. 2009, 2011; Decho 2010, 2011; Stal 2012; Tourney and Ngwenya 2014; De los Ríos et al. 2015; Decho and Gutierrez 2017) that form microbial biofilms (Riding 2000, 2002b; Flemming and Wingender 2010; Reitner 2011a; Flemming et al. 2016) in microbial mats (Gerdes 2010; Mei 2011b, 2014; Reitner 2011b) dominated by cyanobacteria. Although there are lots of problems needing future investigation, these dendrolites (Figs. 3, 4, 5, 6, 7, 8, 9) with the intergrowth of leiolites (Figs. 11, 12, 13, 14, 15, 16) occur as consolidated rocks, and show a complex mineral composition, including: (1) chert lenses in the leiolites (Fig. 11b), (2) special dolomitic cement in the dendrolites (Fig. 5c), and (3) residue of pyrite inside the dendrolites (Figs. 6, 7, 8, 9) and the leiolites (Fig. 14).

The dolomitic cement (e.g. Choquette and Hiatt 2008) in the dendrolites (Fig. 5c) may be classified into a dolomitic precipitate formed in the microenvironment enriched with organic substances in microbial mat (or biofilm) (Roberts et al. 2013; Gregg et al. 2015). In addition, it might represent the residue resulting from organic mineralization of amorphous magnesium silicate before the calcification of EPS making up microbial mats (or biofilms) (Miller and James 2012; Pacton et al. 2012; Burne et al. 2014; Perri et al. 2018). Further, this type of residue not only represents a particular organic dolomite (Mazzullo 2000; Mei 2012) but also forms a special
dolomitic residue (including the chert) in microbial carbonates (Kruse and Reitner 2014) like dendrolite as shown in Fig. 5c. Thus, the small-scale chert lenses (Fig. 11b) might be products of similar process.

The possible pyrite residues in dendrolites (Figs. 6, 7, 8) and leiolites (Fig. 14) might be a result of microbial precipitation (Riding 2000, 2011b) or early lithification (Dupraz et al. 2009, 2011), genetically related to sulfate reduction (Rickard et al. 2017; Lin et al. 2018; Bostick 2019). It also demonstrates two key components i.e. the alkalinity engine (Gallagher et al. 2012) and the organic EPS matrix (Decho 2010; Flemming and Wingender 2010; Tourney and Ngwenya 2014; Flemming et al. 2016; Decho and Gutierrez 2017) involved in carbonate precipitation inside the microbial mats to form dendrolites and leiolites. Briefly, the dendrolites with intergrowth of leiolites mostly dominated by filamentous cyanobacteria represented by Epiphyton (Figs. 5, 6, 7, 8, 9) and Hedstroemia (Fig. 10) in upper part of the Miaoliongian Zhangxia Formation at the Anjiazhuang section in Feicheng city of Shandong Province, grew on a high-energy oolitic-grain bank. It developed in a forced regressive systems tract of third-order depositional sequence that is bounded by drowning unconformities followed by sea-level rise.

Although the fossils of Epiphyton group (Figs. 5, 6, 7, 8, 9, 13, 14, 15, 16) marked by bifurcating branches of dense micrites have been thought as red algal affinity (Luchinina and Terleev 2008, 2014; Luchinina 2009), their dense dark micritic composition clarifies that they have a similar origin to the Girvanella (Figs. 13e, 14 and 16) and the Hedstroemia (Fig. 10), i.e. in vivo calcification (Riding 1991b, 1991c, 2011b; Kah and Riding 2007) of structured sheaths that composed of special extracellular polymeric substances (EPS; Dupraz et al. 2009, 2011; Decho 2010, 2011; Stal 2012; Tourney and Ngwenya 2014; Decho and Gutierrez 2017) with special protein of extracellular sunscreen scytonemins. Therefore, it is more reasonable to assign the fossils of the Epiphyton group into filamentous cyanobacteria. The distinctive bifurcating branches of the Epiphyton group forming shrubs in the Miaoliongian dendrolites (Figs. 5, 6, 7, 8, 9) and leiolites (Figs. 12e, 13a, b, d, 14, 15, 16) may represent early examples of modularity (Suosaari et al. 2018), which further demonstrate that the microbial communities (mainly composed of filamentous cyanobacteria) are able to generate specific morphologies to increase the overall efficiency of network activity during the high-energy marine environment like the modern shrubs in living dendrolitic microbial mats in Hamelin Pool (Suosaari et al. 2018). Moreover, similar bifurcating branches formed by large Lyngbya aestuarii filaments in modern shrubs of living dendrolitic microbial mats that provide a modern analogue for the segmented Gordonophyton of the Epiphyton group (Fig. 6a, c). Therefore, the example of shrubs in Miaoliongian dendrolites also reflects biological habitat of microbial communities that build them, and further provides insight into complex interplay of microbial communities and environment that not only broadens our understanding of dendrolite formation but also provides insight into detailed history of Early Paleozoic skeletal and microbial reefs (Rowland and Shapiro 2002; Kiessling 2009; McKenzie et al. 2014; Ezaki et al. 2017; Lee and Riding 2018; Cordie et al. 2019; Lee et al. 2019; Riding et al. 2019), and microbe–metazoan transitions of the Cambrian (Chen et al. 2019).

6 Conclusions

1) Microbial reefs with a thickness of more than 100 m are composed of dendrolites with the intergrowth of leiolites in upper part of the Miaoliongian Zhangxia Formation at the Anjiazhuang section in Feicheng city of Shandong Province, grew on a high-energy oolitic-grain bank. It developed in a forced regressive systems tract of a third-order depositional sequence that is bounded by drowning unconformities followed by sea-level rise.
These remarkable sedimentary deposits along with sophisticated microscopic features are reflected by diversified calcified cyanobacteria and the heterogeneous sedimentary fabrics.

2) The features of these rocks not only characterize the complexity of microbial reefs but also demonstrate the sophisticated mechanism of microbially-induced in situ carbonate precipitation in multiple and relatively thick non-laminated microbial mat dominated by filamentous cyanobacteria. These deposits reflected several features such as: (i) shrub fabrics make up dendrolites chiefly dominated by filamentous cyanobacteria which are represented by *Epiphyton* and *Hedstroemia*; (ii) the special growing matrix for shrubs of dendrolites is marked by *Bacinella*-like fabric that can be grouped into a feature of modern microbialites (e.g. microbial mats), and the dense micrites may be considered as a result of CaCO$_3$ precipitation occurred simultaneously in metabolically active cyanobacterial layers; (iii) a possible colony of *Nostoc* is represented by the *Lithocodium*-like fabric within the dendrolites; (iv) *Epiphyton* and *Girvanella* intergrew within the leiolites; (v) the organic-mineralization of microbial mats is reflected by dolomitmic cements within dendrolites and the chalk lens inside the leiolites; and (vi) the sulfate reduction involved in carbonate precipitation within microbial mats forming these dendrolites and leiolites is reflected by pyritic residues.

3) Although many problems dealing with the forming mechanism and growth pattern for microbial reef complex dominated by dendrolites with intergrowth of leiolites in this study need further investigation, it provides a suitable example for further understanding of dendrolites with the intergrowth of leiolites built up by non-laminated cyanobacteria microbial mats. We hope that our observations and studies especially for the establishment of *Lithocodium* or the *Lithocodium*-like fabric during the Cambrian can provide some useful approaches and research clues for further research and refine the history of Early Paleozoic skeletal reefs and the microbe-metazoan transitions of the Cambrian as well as further understanding of their driving factors in the future.

**Authors’ contributions**
All the authors have actively participated in the preparation of this manuscript. MMX proposed the main concept and MR wrote the manuscript. These two researchers also collected field data. ZWZ, YH, and QFM assisted in establishing microfabrics, review and proofreading of the manuscript. The authors read and approved the final manuscript.

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**Availability of data and materials**
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**Declarations**

**Competing interests**
The authors declare that they have no competing interests.

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