Micro-framework reconstruction from peloidal-dominated mud mounds (Viséan, SW Spain)

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Abstract Currently, the different roles of peloidal micrites are not well understood in matrix-supported bioconstructions such as mud mounds. Upper Viséan outcrops from Guadiato Valley, Cordoba Province, SW Spain provide well-preserved peloidal-dominated mud mounds. Microstructural analysis of these mud mounds has allowed us to establish 13 microfabrics. Although peloids are conspicuous constituents in the microfabrics, they have been differentiated into seven peloidal-dominated types, grouped into three categories: (a) homogeneous fine peloidal types (3 subtypes); (b) homogeneous coarse peloidal types (2 subtypes); and (c) heterogeneous coarse peloidal types (two subtypes). Fast cm-scale microfabric changes were mapped directly onto thin sections. This methodology allows the researcher to recognize the micro-framework of these mud mounds as a complex, multiepisodic organization of microframes and intermicroframes that are formed by primary and secondary autoomicrites (non-reworked and reworked autoomicrites, respectively), autochthonous internal sediment, and allomicrite and marine fibrous cements.

The thin-section mapping was also used to estimate the autoemicrite-allomicrite relationship, which is crucial in separating the microbial and biodetrital mud mounds.

Keywords Mud mound · Peloids · Autoamicrites · Micro-framework · Carboniferous · Spain

Introduction Mississippian bioconstructions have a wide compositional spectrum as well as broad tectonostratigraphic distributions and varied temporal patterns (Webb 1994, 2002; Bridges et al. 1995; Somerville 2003; Aretz and Chevallier 2007). Their study and classification system have been mainly focused on skeletal and non-skeletal components, cavity system development, and diagenetic features (Lees and Miller 1995; Ahr and Stanton 1996; Jeffery and Stanton 1996; Madi et al. 1996).

Bioconstruction is used here in a global sense to include the whole reef-mud mound spectrum. In this report, we take a broad, non-genetic, non-morphological, and descriptive definition of mud mound at the macroscopic scale, as a bioconstruction composed of more than 50% of rock volume by fine-grained ‘carbonate matrix’ that can be texturally and genetically varied (autoamicrites and allomicrites occur). In volume terms, mud mounds can be considered matrix-supported bioconstructions. Genetically, however, their carbonate matrix can represent a volume ranging from 10 to 100% of recognizable biologically induced framework, or, on the contrary, it can represent an allochthonous carbonate matrix externally produced and later deposited. Despite the current body of knowledge, some questions still remain. How much volume percentage is necessary in order to consider them ‘microbial mounds’ or ‘biodetrital mounds’?
How is it possible to recognize their internal organization or accretional mechanisms?

The isolated Mississippian outcrops from Guadiato Valley (Córdoba, Spain) are composed mainly of siliciclastic deposits of a narrow foreland basin sensu lato developed during the Variscan continental collision between the Ossa Morena and the Central Iberian terrains (Fig. 1a). In this tectonic context, the mud mound factory was very important and well developed in narrow and tectonically controlled blocks. In fact, the Upper Viséan strata provide abundant and excellent examples of mud mounds (Rodríguez-Martínez 2005; Rodríguez-Martínez et al. 2003). Their macrofabric is massive, cryptalgal or leiolitic, but, microscopically, they comprise complex microfabrics, most of them peloidal-rich ones.

In regards to the study of mud mounds, general related questions are (1) the origin and/or source of carbonate mud, (2) the existence or lack of internal organization, and (3) the ability of the produced carbonate mud to form rigid, synsedimentary accretional structures. This study is focused on the last two questions.

Since the 1980s, mud mounds have been seen as self-sufficient carbonate factories where the autochthonous production of carbonate mud occurs. In Mississippian mud mounds, the ‘multi-component mudstone’ of Lees (1964) and the later ‘polymuds’ fabrics (Lees and Miller 1985) are...
reference terms or concepts in the mesostructural study of their internal organization. However, the accretion mechanisms are not so clear when both massive or leiolitic fabrics and aphanitic micritic microfabrics occur simultaneously and are dominant. The presence of peloidal micrites is then argued as microbial-mediation activity, and the possibility of the abiogenic character of the mud mound disappears. However, the focus should be whether peloidal microfabrics were able to produce rigid, accretionary microstructures and how to recognize and quantify these microstructures.

Peloidal matrices have been described both in Recent and fossil bioconstructions (see Riding 2000; Reitner and Neuweiler 1995; Reitner et al. 2000; Folk and Chafetz 2000; Pratt 1995), and their origins have also been discussed for a long time (Monty 1965; Macintyre 1985; Chafetz 1986; Reitner 1993; Kazmierczak et al. 1996). However, there are few microstructural classification schemes based on descriptive parameters in which peloids are the dominant elements (Permian-Triassic: Reid 1987; Adachi et al. 2004; Jurassic: Sun and Wright 1989; Schmid 1996). This would be irrelevant in those bioconstructions dominated by skeletal-frameworks where peloids have a secondary importance and have been interpreted in several ways (peloidal cement and/or peloidal internal sediment in cavities/porosity, peloidal interframe, peloidal microbacterial crust). What were the roles played by the peloidal micrites, in the apparently frameless, peloidal-dominated mud mounds of the Guadiato Valley?

The main objectives of this paper are to show how the classification and detailed mapping of mud mound microfabrics allow us to: (1) recognize the different roles of peloidal micrites, (2) estimate the volume relationship between autochthonous and allochthonous mound carbonate production, and (3) reconstruct the micro-framework of these mud mounds.

**Geological setting**

We studied the mud mounds of the Guadiato Valley, which are close to the villages of Espiel and Belmez at NW Cordoba province, in the southern part of the Iberian Massif (Fig. 1a). The Iberian Massif corresponds to the SW part of the European Variscan Orogen, formed as a consequence of the collision between Laurentia and Gondwana during Devonian-Permian times (Matte 1991). The analyzed sedimentary record is located in a narrow band of Precambrian-Carboniferous outcrops, tectonically bounded by the Hornachos and Peraleda faults, which are part of the BCSZ (Badajoz-Córdoba Shear Zone, Fig. 1). The BCSZ is a NW-SE structural band 400 km in length between Tomar (Portugal) and Córdoba (Spain). This structure is an important and complex cortical accident and has been, in general, accepted as the limit between the Ossa Morena Zone (OMZ) and the Central Iberian Zone (CIZ) -Iberian Massif divisions (works from Iberian Massif in Ábalos et al. 2002; Pérez-Estaún and Bea 2004). However, the boundaries between OMZ and CIZ have been a source of controversy (San José et al. 2004). The BCSZ has been interpreted as a suture, part of the Cadomian subduction complex, which will later reactivate during the Variscan Orogeny (Quesada 1992) as an intracontinental shear zone (Eguíluz et al. 2004). It has also been interpreted as just a Variscan suture (Azor et al. 2004). CIZ would be interpreted as part of the Gondwana margin whereas the OMZ is viewed as a terrain related to Gondwana (Simancas et al. 2004).

In the Guadiato Valley, the dispersed and isolated Carboniferous outcrops were first mapped and informally divided into three bands by Pérez-Lorente (1979), probably based on the earlier work by Ortuño (1971). Mississippian outcrops (Fig. 1c) are situated along the southwest and central bands, whereas the northeast band corresponds to the Pennsylvanian ones. Different paleogeographic models have been proposed for the Mississippian outcrops (Gabaldón et al. 1985; Martínez Poyatos et al. 1998; Martínez Poyatos 2002; Cózar and Rodríguez 2003). However, the relationship between the Mississippian central band and its pre-Carboniferous basement is unknown; and there are no lithostratigraphic correlations between both of the Mississippian bands.

The studied peloidal-dominated mud mounds are located in two sectors, both placed in the central band (1 and 2 in Fig. 1c): Sierra Boyera (SB) and Sierra del Castillo-Sierra de la Estrella (SCSE). Most studies carried out in the central band have been focused on paleontology (corals, algae, foraminifers, conodonts), biostratigraphy (mainly foraminifers and corals) and the stratigraphy of the calcareous deposits (Mamet and Martínez-Díaz 1981; Cózar and Rodríguez 1999, 2003, 2004; Cózar and Mamet 2001; Cózar 2003a, 2003b; Cózar and Vachard 2004a, 2004b; Bermúdez-Rochas et al. 2004; Rodríguez et al. 2004). Detailed cartographic, stratigraphic, and sedimentological studies allowed the recognition of five mud mound stages within the Upper Viséan succession of up to 900 m in thickness (Rodríguez-Martínez 2005).

**Methodology**

The mapping of the outcrops was done with enlarged aerial photographs (original scale 1:18,000) to delineate both the mud mound and off-mound facies distribution. Fourteen stratigraphic sections were measured. In total, 1,013 samples were collected (318 from the SB sector and 697 from the SCSE sector). An identical number of thin sections
were prepared: large thin sections (5 × 8 cm in size) from mud mound samples and standard sizes (4.6 × 2.5 cm) from the associated facies. The sampling from the most continuous and complete sections was selected for detailed microfacies analysis (698 thin sections). The analyzed features from both mud mound and off-mound thin sections were: (a) a semi-quantitative estimation percentage of 34 components (visual estimation charts from Flügel 2004); (b) counting the number of specimens per thin section of components at different taxonomic levels (red algae as unguellarellaceans and *Fasciella*, green algae as *Koninkkpora*, and *Incertae sedis* as *Saccamminopsis*); (c) qualitative estimation of small and rare calcimicrobes (*Girvanella*, *Renalcis*) and problematic taxon such as *Aphralysia*.

The maps of mud mound microfabrics were drawn directly over the thin sections under the stereomicroscope (more than 200 thin-section maps). Each type of microfabric was codified and numbered, and the resulting mapped thin sections with their codes were photocopied. The photocopies were digitalized and re-drawn with CorelDRAW software.

**Mud mound stratigraphic record**

The Mississippian succession from the SB and SCSE sectors has been informally divided into three groups (1, 2, and 3 in Fig. 2): (1) lower heterolithic units, (2) middle carbonate units, and (3) upper heterolithic units. Mud mound facies occur in the lower and middle units and comprise five megastructural stages, from the base to the top (A–E in Fig. 2): (A) mud mound-derived boulders, (B) tabular banks and associated breccias, (C) isolated-pinnacles belt, (D) amalgamated-bioherms belt, and (E) isolated-bioherms belt. The first two mud mound records (A, B in Fig. 2) occur in both sectors whereas the rest of the stages (C–E in Fig. 2) were only developed in the SCSE sector.

Lower heterolithic units (up to 180 m in thickness) start with moderately channelized to sheet-like bodies of internally massive poorly sorted, pebble-cobble, mainly clast-supported polymictic conglomerates. The conglomerates grade into centimeter-scale tabular beds of calcarenites, lenticular beds of granule-pebble polymictic conglomerates, and interbedded marls, forming a fining upwards interval. It is followed by slumped levels and massive cobble-boulder polymictic megabreccias. The size and composition of clasts in the megabreccias vary from angulose to well-rounded sandstone and calcarenites pebbles, quartzite pebble-cobbles, and large metric mud mound-derived boulders (A in Fig. 2). The top of the lower heterolithic units corresponds to massive oolitic limestones with sandstone pebbles, which overlie unconformity the previous megabreccias.

The middle carbonate units are always tectonically bounded. Thickness ranges from 60 m in the SB sector to up to 700 m in the SCSE sector. Middle carbonate units have four mud mound phases (B–E in Fig. 2). Stage B has 130 m of maximum thickness and from 500 to 2,340 m in width in the SB and SCSE sectors, respectively. Basal beds correspond to black, well-bedded wackestones to packstones, which grade into the mud mound facies. The last ones are formed by massive bank intervals 5–20 m thick, interbedded with mud mound-derived monomictic breccia.
intervals of up to 5 m thick. Because stage B is always tectonically bounded, the stratigraphic relationship with the coeval off-mound record is unknown.

The base of stage C (Fig. 2) begins with pelmatozoan-rich deposits. Stage C is formed by four isolated pinnacle mud mounds, close intermound facies, and off-mound coeval well-bedded limestones. Pinnacles have 110 m of maximum thickness and range from 144 to 555 m in width. At the top, pinnacles show flank facies development, and, in the flank facies, small interbedded satellite mud mounds occur.

The following mud mound megastructural stage (D in Fig. 2) starts after an interval of marls with interbedded black wackestones. Mud mound facies reach 180 m in thickness and extend up to 3,000 m in width. They are organized in lenticular to hemispheroid 1–3 m thick bodies, vertically stacked in 5–60 m thick intervals.

The last mud mound stage, in the middle carbonate units, corresponds to isolated bioherms (E in Fig. 2) and off-mound coeval limestones. Bioherms are up to 35 m in vertical stacking and 250 m in width. The upper part of the middle carbonate units consists of well-bedded limestones, texturally and compositionally varied. The boundary between the middle carbonate units and the upper heterolithic units is covered in the SCSE sector and corresponds to a thrust in the SB sector. The upper heterolithic units (3 in Fig. 2) correspond to sandy calcarenites, limestones, and channel bodies of pebble and cobble siliciclastic conglomerates, sandstones, and siltstones with bedding planes rich in plant remains.

In outcrop scale and hand-polished samples, these mud mounds have a massive or cryptagal mesostructure (scales according Shaphiro 2000). Macrofossil content is really very scarce and dominated by pelmatozoan remains. Only mud mound-derived boulders (stage A in Fig. 2) have a significant amount of sponges, solitary and colony corals, brachiopods, and gastropods, all of them with a patchy cluster distribution that never forms a skeletal framework. The content in skeletal (microfauna and microflora) and non-skeletal grains of Guadiato mud mounds is around 25% of the rock volume average (Fig. 3a). However, in most cases (more than 50%), these grains are just accessory components (<1% of the rock volume), and only some of them have volume averages of 5–10% in the A and B stages (Fig. 3b).

**Mud mound microfabrics**

In general, most of the mud mound microfabrics show an elevated degree of heterogeneity. In a few centimeters, fast microfabric changes occur, such as a patch of fenestellid cementstone surrounded by biomicrites, which grades into a pelsparitic cavity filling, where the roof can be partially supported by skeletal remains (some of them covered by fine peloidal crusts). In this hypothetical case, the mud mound is composed of different carbonate fractions (skeletal grains, peloids, micrite, and cement) where peloids can occur isolated in the biomicrites, as part of the cavity filling as well as forming the crust. In addition, giving a final general microfacies name that responds to this heterogeneity on the microscale can be not only difficult but also unrealistic in descriptive terms.

The spectrum of Guadiato mud mound microfabrics are described below as peloidal-, intraclastic-, bioclastic-dominated, and micritic microfabrics.

**Peloidal-dominated microfabrics**

The peloidal-dominated microfabrics (Figs. 4, 5a–g) are those where ≥50% of the rock volume is formed by the peloidal fraction (fine and/or coarse peloids). The peloidal fraction is composed of spherical to ellipsoidal peloids 10–100 μm in diameter. The distribution of peloidal sizes can be homogeneous or heterogeneous (a mixture of sizes).
Fine peloidal microfabrics

The size of peloids varies from 10 up to 70 μm, with an average of 20 μm. The microfabric subtypes have been divided on the basis of the peloid: cement ratio, the matrix- or cement-supported character. A total of three fine peloidal microfabric subtypes have been recognized from densely packed to cement-supported subtypes (70:30; 50:50; and 20:80 peloid: cement ratios; see Fig. 5a–c). Normally, all of them appear associated and grading into each other.

Fine peloidal microfabrics may form crusts from 1 mm up to 5 mm in thickness around skeletal remains (Fig. 4a) and commonly appear associated with encrusting bryozoans and sponges. In turn, crusts can also be colonized by microencrusters like foraminifers and others of unknown origin. Fine peloidal microfabrics also occur within the cavity system, in anti-gravitational relationship, as the substrate of marine fibrous cements and in a geopetal relationship as a part of the filling sequence. Secondary porosity, such as fractures and interclastic spaces in breccias, can display fillings of these microfabrics in some mud mound stages. Isolated microdomes (3 × 2 cm of maximum size), microcolumns (1 × 2 cm of maximum size), and more complex reticulate forms are built by the homogeneous fine peloidal microfabrics (Fig. 4b, c). Microdomes can be internally laminated or not. Most of the time, the lamination corresponds to an irregular and discontinuous thin lamination. However, laminae, which range from 200 to 400 μm can be recognized. Lamination results from both packed- and size-changes of the peloids. Each lamina
Fig. 5  a–e Peloidal-rich mud mound microfabrics with a homogeneous distribution of the size of peloids: a–c fine peloidal microfabrics from the matrix to the cement-supported varieties (m-s and c-s, respectively); d, e coarse peloidal microfabrics (m-s and c-s, respectively). f–k Heterogeneous mud mound microfabrics: f, g heterogeneous coarse peloidal microfabrics (c-s and m-s subtypes); h, i heterogeneous intraclastic microfabrics (c-s and m-s subtypes); j, k heterogeneous bioclastic microfabrics (c-s and m-s subtypes). Graphic scale bars are 1 mm in the f–k pictures. Numbers in pictures represent the assigned codes during the mapping of the thin sections; see Fig. 6.
starts with a clear, less packed peloidal interval (up to 50 μm for the diameter size of the peloid) and is gradually followed by a dark, densely packed interval with peloids from 10 to 20 μm in size.

Coarse peloidal microfabrics

We use the term coarse peloidal microfabrics when the distribution of the size of peloids ranges in diameters from 30 to 100 μm. The coarse peloidal microfabrics are divided into two subclasses: homogeneous coarse peloidal microfabrics (Fig. 5d, e) and heterogeneous coarse peloidal microfabrics (Fig. 5f, g).

Homogeneous coarse peloidal microfabrics develop the cement as well as the matrix-supported subtypes (c-s and m-s, respectively). The diameter sizes of peloids on the cement-supported type are slightly larger, with an average of 55 μm, in contrast to 40 μm for the matrix-supported type. Enclosed bioclastic content is low and generally formed by ostracod valves. Fecal pellets can be locally abundant and non-skeletal grains like peloidal intraclasts also occur.

Homogeneous coarse peloidal subtypes have always been observed as associated with the filling cavity system and generally in geopetal relationship. The cement-supported type can also appear in anti-gravitational relation to the cavities. Some large reticulate cavities are particularly characterized with the filling of this peloid type that can or not occur banded. Irregular bands are the result of differences in both peloid size and packing, but banding is also a result of the changes in the microfabric type, when interbedded sequences of the c-s and m-s subtypes occur. The contact surface, or transition, in these interbedded sequences of the c-s and m-s subtypes are around 50:25:25 PIB ratio and occur as c-s and m-s subtypes. The coarse peloidal microfabrics are divided into two subclasses: homogeneous coarse peloidal microfabrics (Fig. 5f, g).

Irregular peloidal intraclasts, not angular in shape and with diffuse margins, are the dominant elements in the heterogeneous intraclastic microfabrics that have a 25:50:25 PIB ratio. The irregular intraclasts are randomly distributed and surrounded by fine and coarse peloids. The intraclasts are identical to the surrounding matrix; thus, they can be as large as 2 mm in some cases, but they are normally found in the 500–1,000 μm interval.

The heterogeneous intraclastic microfabrics also display the c-s and m-s subtypes (Fig. 5h, i, respectively). These microfabrics occur with a patchy distribution: (1) as basal geopetal filling in large cavities, (2) associated with fractures and brecciated areas, and (3) grading and interbedded with other microfabrics.

Bioclastic-dominated microfabrics

The most heteromodal distribution of component sizes occurs when the skeletal content is high and the PIB ratio is around 25:25:50. Bioclastic-dominated microfabrics develop c-s and m-s subtypes (Fig. 5j, k) and show a patchy distribution. As previously indicated, the volume contribution of skeletal grains is low (Fig. 3a, b). Crinoids, and occasionally sponges in some megastructural stages, are secondary components (5–10% of rock volume on average). Minor components are fenestellid bryozoans (3–5% of rock volume on average) as well as ostracods, calcareous algae, brachiopods, and burrows (1–3% of rock volume on average). Ramose and encrusting bryozoans, echinoid spines, brachiopod spines, foraminifers, gastropods, trilobite fragments, and solitary and tabulate corals are accessory components (<1% of rock volume on average). They occur with a patchy distribution, and only few components can occasionally reach high concentrations. The particular situation where the skeletal and cement volume ratio is around 1:1 has only been observed with fenestrate bryozoans and the Incertae sedis Saccamminopsis.

Micritic microfabrics

When micrite (<4 μm) and/or microspar (4–31 μm) are the dominant components, the resulting microfabric corresponds
to mudstones or wackestones. Micrite is common as geopetal internal sediment in cavities. It also forms isolated patches grading laterally into other microfabrics, and it is associated with secondary porosity related to fractures. Micrite in cavities can occur as the only geopetal infilling, as successive micritic infillings, or alternating with other microfabrics. It can also occur as mudstone-wackestone fining-upwards sequences. The mud mound micritic fraction shows no optical differences with the micrite observed in the coeval off-mound deposits.

**The micro-framework reconstruction**

As it was shown before, peloids characterized several microfabrics which can grade into one another in a few centimeters. The classification and the detailed mapping of these microfabrics can help to recognize their internal spatial distribution, quantify the volumetric relationship between the automicrite production and the allomicrite input, and reconstruct the micro-framework.

**Classification and mapping**

For the mapping of the thin sections, as it has been described through the previous chapters, the classification of microfabrics was made on the basis of the following categories: (1) homogeneous or heterogeneous distribution of sizes, (2) main carbonate fraction (peloidal, intraclastic, bioclastic, and micritic ones), and (3) type of final ‘support’ (matrix or cement-supported character). The resulting microfabric subtypes were codified with numbers (Figs. 5, 6); however, the numbers are not related to the temporal organization of the microfabrics. A total of 13 microfabrics were separated and mapped (Fig. 6): (1) homogeneous fine peloidal microfabrics (subtypes 7, 8, and 9); (2) homogeneous coarse peloidal microfabrics (subtypes 10 and 11); c) heterogeneous coarse peloidal microfabrics (subtypes 1 and 2); (3) heterogeneous intraclastic microfabrics (subtypes 3 and 4); (4) heterogeneous bioclastic microfabrics (subtypes 5-6); and (5) micritic microfabrics (subtypes 12 and 13, mudstone and wackestone, respectively). Dolomicrite is quite uncommon and was codified as number 14.

Marine fibrous cements associated with the cavity system were also mapped and estimated. They correspond to acicular and fibrous types forming isopachous as well as irregular crusts in cavities, and also as isopachous rims around skeletal components. Radial fibrous cements also appear as thick rims in the cavity system. They are turbid, inclusion-rich crystals and they exhibit undulose extinction. Both radial fibrous and fascicular-optic calcite extinction patterns have been recognized. All of them were codified as 15 (Fig. 6).

**Autochthonous production versus allochthonous sedimentation**

In the Upper Viséan mud mounds from Guadiato Valley, the homogeneous fine peloidal microfabrics (subtypes 7, 8, and 9; Figs. 4, 5a–c) correspond to automicrites or autochthonous microcrystalline carbonates formed in place. Automicrites have been interpreted as the result of different genetic modes of production (see summarized in Flügel 2004, pp. 81–88): (1) physicochemical precipitation (e.g.: Macintyre 1985), (2) in-place mineralization via organic matrices (organomicrites sensu Reitner 1993; see organomineral definition comment in Défarge et al. 2009), and (3) via metabolic processes related to microbial benthic communities causing carbonate precipitation (Castanier et al. 1999; Chafetz 1986; Pomar and Hallock 2008). Petrographically, the possible origins are not possible to recognize because different processes result in similar end products.

The homogenous coarse peloidal fabrics (subtypes 10 and 11—Figs. 5d, e, 6) represent the internal sedimentation and production of automicrites that are exclusively developed in the cavity system. The type of ‘support’ is related to the relationship between the sedimentation, production, and the cementation rate in the cavity: matrix-supported types occur for high and continuous rates of automicrite sedimentation and production, whereas low and discontinuous rates produce the cement-supported type. In the cavity, the transition from the matrix-supported to the cement-supported type could also represent an increase in water pumping that favored marine cementation.

The heterogeneous microfabrics (subtypes 1-2, 3-4, and 5-6—Fig. 5f–k) show evidence of a mixture of processes: (1) autochthonous production of fine peloids in the pores and pockets, (2) early lithification (the development of matrix-supported growth cavities, synsedimentary fractures with marine infillings, brecciation, and early marine cementation), (3) a volumetrically diverse skeletal contribution (both autochthonous and paraautochthonous assemblages occur), (4) an external skeletal contribution (allochthonous assemblages which have loose platform-shed components), and (5) reworking (peloidal intraclast production, burrowing, winnowing into the cavity system). All of these processes reveal an asymmetrical balance between the constructive and destructive processes where the autochthonous processes are dominant. The heterogeneous intraclastic subtypes are common in the stages when vertical accretion is dominant (stage C in Fig. 2), whereas the heterogeneous bioclastic subtypes are more common in stages characterized by lateral accretion (stages D and E in Fig. 2). Thus,
heterogeneous intraclastic microfabrics could indicate major gravity and/or hydrodynamic reworking rates, whereas the heterogeneous bioclastic subtype could reflect more suitable conditions to colonize the substrate by non-microbial benthic communities as well as a greater input of shallow-water platform-derived grains. Heterogeneous microfabrics are autochthonous microfabrics, with automicrite production, but signals of microfabric reworking (physical, biological, and/or diagenetic processes) also occur.

Finally, the micritic microfabrics (subtypes 12 and 13), where micrite shows no any recognizable accretional forms, most part of the cases records hydrodynamic, gravitational or synsedimentary deformation features (oriented bioclasts, geopetal fillings, angulose intraclasts), and shows no petrographic differences with the off-mound record, are interpreted here as allomicrites as a part of the external sedimentation input recorded in the Guadiato mud mounds.

Table 1  Comparison of homogeneous peloidal microfabrics with some selected Carboniferous and other bioconstructions

| Homogeneous peloidal microfabrics | Carboniferous examples | Other examples |
|----------------------------------|------------------------|---------------|
| **Fine peloids**                 |                        |               |
| Subtype 7                        | Hummocky/crinkled microstructures | Hemispheroids of dense micritic/ fenestral microbialites |
| Microdomes                        | of clotted micrite and fine peloids [1] | Peloidal micrites showing dome-shaped algal microstromatolites [10] |
|                                  | Peloidal-rich sediment within encrustation or microbial incrustation [2] | Accretionary organomicrites [11] |
|                                  | Pseudo-stromatolitic/thrombolitic structures of fine-grained peloidal fabrics [3] |               |
| Crusts                            |                        |               |
| Subtype 8                        | Encrusting microbialites [4] | Dense peloidal crusts [12] |
|                                  | Microbial incrustations [5] | Laminated peloidal microbial crust or non-skeletal stromatolites [13] |
|                                  | Micrite crusts in Type B boundstones [6] |               |
| Subtype 9                        |                        |               |
| **Coarse peloids**               |                        |               |
| Subtypes 10 and 11               | *Chondrostroma problematicum* [7] | Laminated mudstones [15] |
|                                  | *Spongistromata ovuliferum* [7] | Dense micritic/peloidal microbialites [9] |
|                                  | Loose bacterial/algae peloids [1] | Coarse grained peloidal/detritic microbialite or crust type 1 [12] |
|                                  | Internal sediment [2, 5] | Container organomicrites [11] |
|                                  | Microbial? peloids [8] | Peloidal crust-boundstone [16] |

[1] Bridges and Chapman (1988); [2] Somerville et al. (1992); [3] Horbury (1992); [4] Mundy (1994); [5] Pickard (1996); [6] Della Porta et al. (2003); [7] Gürich (1906); [8] Lees and Miller (1995); [9] Neuweiler (1993); [10] García-Mondéjar and Fernández-Mendiola (1995); [11] Reitner et al. (1995); [12] Leinfelder et al. (1994); [13] Nose et al. (2006); [14] Bourque and Gignac (1983); [15] James and Ginsburg (1979); [16] Koch et al. (1994)
pellets present in both species with stercomes, the waste products derived from the xenophyophores. They are giant, multinucleate, agglutinated rhizopods that excrete a slimy substance. They are typical benthic deposit feeders in the deep sea. They are commonly found associated with seamounts, ridges, canyons, topographic highs in general, and are very abundant in the bathyal north east Atlantic Darwin Mounds today (Hughes and Gooday 2004).

Heterogeneous microfabrics are characteristic mud mound microfabrics and have been referred to in various ways, mainly using Dunham’s and Folk’s classification terms, such as peloidal mudstone/wackestone, pelsparite, and biomicrite. The presence of cavities and pores has often been a noteworthy characteristic in the final name, for example, stromatoids-bearing wackestone, fenestral bimicrite. Some of these textural terms are used as prefixes of biolithite (Pratt 1995) as well as boundstone terms (boundstone type A in Della Porta et al. 2003), and they reflect the author’s conception of the mud mound. When the heterogeneous bioclastic microfabrics are formed by about a 1:1 skeletal/cement volume relationship, they are called skeletal/ non-skeletal organism-type name + cementstone (Fagerstrom 1987) or biocementstone (Tsien 1981; Webb 1996). In Guadiato mud mounds, only Saccamminopsis and fenestrate bryozoans form cementstones where they play a scaffold constructor role (Rodríguez-Martínez et al. 2003). Other delicate organisms have played similar role in the so-called biocementstone frameworks (Webb 1996), common framework in mud mounds.

**Micro-framework contribution: the roles of the peloidal matrix**

The mud mounds comprise both soft (allomicrites and automicrites) as well as early indurate carbonate muds (automicrites), which can be reworked after their formation, resulting in paraautochthonous muds. Homogeneous fine peloidal types show no evidence of bio and/or physical reworking, so they are primary or original microfabrics, considered here to be primary automicrites. However, heterogeneous types show signs of biological and physical reworking, clusters of fine peloidal production in pores, evidence of colonization by non-microbial communities, and/or external skeletal input. Thus, they are secondary, or modified, reworked or colonized microfabrics and are therefore considered herein as secondary automicrites (a mixture of the previous ones, paraautochthonous automicrites). They represent soft substrates that were quickly lithified as the development of matrix-supported cavities makes clear.

The framework scenario (Fig. 7) is built by successive production of firm, primary automicrites, with different accretional morphologies (clusters, crusts, microcolumns, and microdomes), which alternate and develop between and within the secondary automicrites and into the growth cavity system. In the cavities, autochthonous internal sediment (homogeneous coarse peloidal microfabrics) and marine fibrous cements also contribute to the rigidity of the final meshwork. In fact, the cavity filling records many of the last synsedimentary episodes that occurred in the mud mound: the (1) production of cryptic automicrites, (2) colonization by cryptic biota, (3) reworking processes, (4) entry of allomicrite by gravity or currents, and (5) early marine cementation stages and so forth.

Peloidal-dominated mud mounds from the Guadiato Valley are internally highly organized in multiply episodic relative microframes and inter-microframes (Fig. 7). An initial soft intermicroframe turns into microframe once it has been lithified, and the automicrite production continues around and within it.

In Guadiato, the ‘fine-grained carbonate matrix’ and cavities represent about 60–79% of the rock volume, whereas the automicrite production volume is higher than the external allomicrite input (which is less than 13% of the rock volume, on average). The mapping shows that the record of secondary automicrites is dominant (54% of the rock volume, on average), whereas the primary automicrites average is around 16% of the rock volume.

The autochthonous internal sediment and the marine fibrous cement in the cavity system are, on average, 8 and 9% of the rock volume, respectively. Therefore, peloidal-dominated mud mounds from the Guadiato Valley are clearly dominated by automicrites (primary and secondary). The mud mounds are seen here as a continuous series of processes of automicrite production, allomicrite input, and internal reworking (chemical, physical, and/or biological processes).

**Primary automicrite distribution model**

The primary automicrites have a similar distribution (13–19% rock volume) throughout the different mega-structural mud mound stages (A–E in Fig. 2) in the Guadiato Valley. However, their growth morphologies (crusts, microcolumns, and microdomes) show a shallowing-upward trend.

A shallowing-upward mud mound phases model is defined herein (Fig. 8) combining (1) the total number and (2) presence of selected components per thin section. Calcareous algae and other components have been used as a light/energy index on the basis of both mud mound and off-mound facies analysis, which are characterized by distinctive biotic assemblages (Rodríguez-Martínez 2005), and following the previous works from the Béchar Basin, Algeria (Bourque et al. 1995; Madi et al. 1996).

Calcareaous algae are accessory and minor components in these mud mounds (on average, <1% and between 1 and
3% of the rock volume, respectively) whereas they can appear in the off-mound facies as main and secondary components (>10% and between 5 and 10%, respectively). In fact, the presence of different calcareous algae was taken as a good light/energy indicator, independent of the total number of algae, because trends are similar in both records of the mud mound and off-mound facies (Rodríguez-Martínez 2005).

The facies analysis of the coeval off-mound record shows the following features: (a) *Fasciella* has the most conspicuous distribution along the outer and inner platform, with a maximum abundance above the storm weather base; (b) ungdarellaceans are much more restricted to intermound and flank areas, except for the upper part of the succession; (c) calcareous algae meadow deposits occur along the middle and inner platform as *Fasciella*, paleoberesellids, and *Koninckopora*-dominated belts from the deepest to shallow waters, respectively (Fig. 8a). *Fasciella* is generally accepted as red algae (Bourque et al. 1995; Madi et al. 1996; Gallagher 1998; Groves et al. 2003). The presence of paleoberesellids is common in shallow subtidal environments (Adams et al. 1992) at a depth of 10 m close to the fair weather base (Horbury and Adams 1998), and the cosmopolitan dasyclad *Koninckopora* is common in high-energy shallow-water
deposits such as shoals (Mamet 1991). It forms true meadows (Wood 1940) close to and above the fair weather base, between a depth of 5–10 m (Gallagher 1998).

The autochthonous and parautochthonous record of calcareous algae and calcimicrobes in the mud mound is formed by *Fasciella*, *Girvanella*, ungdarellaceans, *Aphralysis*, and *Renalcis*. Moreover, the last two are quite uncommon (a presence of less than 10% in samples with algae), and only *Fasciella* shows a higher frequency (>75% in samples with the presence of algae). Green algae, such as paleoberesellids and *Koninckopora*, are shallow-water-derived allochthonous remains in most of the mud mound mega-structural stages (Fig. 2).

Thus, the idealized shallowing-upwards mud mound sequence is formed by five phases, defined by the component richness and the light/energy component index (phases I–V in Fig. 8b). Component richness is considered here as the total number of components per thin section. It has been grouped in low, middle, and high richness intervals: less than 10, exactly 15, and more than 15 components per thin section, respectively. Calcareous algae such as *Fasciella*, ungdarellaceans, *Koninckopora*, paleoberesellids, and other allochems (such as ooids and oncolites) have been chosen as a qualitative light/energy index.

Phase I is defined by very low component richness and the absence of calcareous algae. We interpret phase I as
deep subtidal, aphytic to dysphotic conditions (Fig. 8b), whereas phase II is defined by the presence of Fasciella in low-richer assemblages. The presence of red algae together with the first occurrence of paleoberesellids and middle component richness, point to the dysphotic to euphotic transition and represent conditions close to the storm weather base (phase III in Fig. 8b). The arrival of the remains of the green algae Koninckopora, later followed by platform-derived grains such as oolites and oncolites, represents mud mound phases in true euphotic shallow-water conditions (phases IV and V, respectively, in Fig. 8b), between the storm and fair weather bases.

The observed frequency of primary automicrites is summarized in Fig. 8c. Crusts around skeletal remains and cavity walls occur without evidence of depth control, from the deepest to the shallow subtidal water mud mound phases (I–V). Massive microdomes are dominant in middle depth mud mound phases (III). The growth morphologies of primary automicrites begin to diversify toward shallow waters. The frequency of microdomes, especially the internally laminated ones, together with the microcolumns and more complex reticulated forms, increase during phases IV and V. Crusts with diverse microencrusters, growing into the remaining cavity system as well as into the secondary porosity, related to breccias and fractures, are common in shallow-water conditions where mud mounds grow close to the FWB. Under these conditions, the cavity system develops more reticulate shapes and is filled by complex sequences dominated by autochthonous internal sediment (homogeneous coarse peloidal microfabrics). Stromatactis-like cavities are common from phases I–III, whereas reticulate cavities are dominant from phases IV–V.

Summary and conclusions

A spectacular succession with five mud mound mega-structural stages characterizes the Visean bioconstruction record from the Guadiato Valley in the southwest region of Spain. These mud mounds have a 60–79% volume average of a fine-grained carbonate matrix, mostly of peloidal character.

Thirteen mud mound microfabrics have been recognized on the following three factors: the composition of the main fraction (peloidal, intraclastic, bioclastic, and micritic fractions), distribution of sizes (homogeneous or heterogeneous), and type of ‘support’ (matrix- or cement-supported character).

The spectrum of microfabrics was analyzed and mapped directly over the thin sections. This methodology has allowed us to recognize the different roles of the peloidal matrix, and determine the best estimation of their volumetric importance and contribution to the mud mound microframework.

The internal organization of mud mounds from the Guadiato Valley shows a continuous series of automicrite production, allomicrite input, and internal reworking processes. The autochthonous production is much higher than the allochthonous input. In fact, these peloidal-dominated mud mounds have a high and complex multiepisodic microframework formed by primary and secondary automicrites (16 and 54% of the rock volume, on average) with the development of growth cavities filled by autochthonous internal sediment (8%), allomicrites (13%), and marine fibrous cements (9%).

Primary automicrites are formed by homogeneous fine peloidal microfabrics (three subtypes) which display different growth morphologies (clusters, crusts, massive and laminated microdomes, microcolumns, and reticulate developments), as well as modes of occurrence (associated with skeletal components, microencrusters, cavities, and fractures). The distribution of the primary automicrite morphologies shows a shallowing-upwards pattern from the simplest (clusters and crusts) and most massive to the complex and internally laminated forms.

Secondary automicrites are formed by heterogeneous microfabrics (six subtypes belonging to the peloidal, intraclastic, and bioclastic-rich subclasses). They show a mixture of both in situ constructive and destructive, or reworking, processes. Thus, they are paraautochthonous automicrites.

The autochthonous internal sedimentation or production of automicrites in the cavity system is formed by homogeneous coarse peloidal microfabrics (two subtypes).

Finally, the microfabrics dominated by the micritic fraction (two subtypes) represent the external allochthonous micrite input into the mud mound, and are considered allomicrites.

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