CALCAREOUS MICROBIALITES AND ASSOCIATED BIOTA IN THE MEDITERRANEAN COASTAL LAGOONS AND PONDS OF SOUTHERN FRANCE: A KEY FOR ANCIENT BIOCONSTRUCTIONS?

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Abstract. Microbialites were found in various lagoons of the Mediterranean coast of southern France. They preferentially develop on pebbles on the edge of semi-enclosed water bodies, which can be temporarily dwatered and desiccated. The microbialitic building is achieved episodically with variations in the characteristics of the different layers. Its cerebroid and pustular appearance is the most remarkable macroscopic feature. Occasionally, other organisms may be involved in the construction, such as bryozoans and serpulid worms that form also in other places small unconsolidated bioconstructions. The analogy of these phenomena with bryozoans-serpulid-microbialite reefs occurring during Miocene in Paratethys, like those noticed in Romania, proposed by some authors, is considered unlikely, despite some common features.

Key words: mediterranean, coastal lagoons, microbial mats, biomineralization, serpulids, bryozoans, Miocene.

1. INTRODUCTION

The Mediterranean coast of Languedoc and Roussillon comprises numerous coastal lagoons that differ very much in size (Fig. 1). Many studies looked into the environmental aspect, the biodiversity, fisheries and shellfish farming, subject to the conditions of an unstable complex system, especially, regarding the fluctuating salinity. Nevertheless, some works based on observations made in the French Mediterranean coastal lagoons proved that salinity is not the only factor of life distribution and led to the biological concept of confinement (Guélorget et al., 1983) and a better understanding of the paralic realm. The confinement corresponds to the water renewal time and/or intake of marine-origin elements at a given point. Thus, the settlements of lagoons are organized from the sea to continental borders of lagoons according to confinement gradient. The confinement at a given point is characterised, in fact, by the lack of an element (or more) of marine origin at this point. Bioconstructions considered as reefs (Lepareur et al., 2013) were reported in such paralic settings, including worms, bryozoans and more or less associated barnacles.

Following the discovery of well developed microbialites in coastal ponds in Sardinia (Saint Martin and Saint Martin, 2015), it was interesting to know whether this phenomenon is common to other coastal areas of the Mediterranean. The Mediterranean coast of Languedoc and Roussillon may, thus, have similar physical-geographic conditions, with confined and/or isolated environments and temporary communications with the sea. A verification of this assertion, therefore, needed to be performed. So, using satellite images, we focused our investigation in similar sites with the potential to support microbialite development. Bibliographic resource about this topic is very scarce, with only a unique short communication without any image support of Boekshoten (1995), during the 2nd European Regional Meeting of ISRS (International Socie-
ty for Reef Studies). This author reported the presence of small mineralized bioconstructions, with presumption of microbial intervention in a brackish pond, named by him "Lac d’Evêque" (SE of Gruissan), but, formally unknown under this name in the local toponymy. He evokes stromatolite limestone crusts developed underneath algal mats on pebbles and small bioherms made of serpulids and the bryozoan *Electra* and compares these structures to Mio-cene buildups of Paratethys.

The first objective of this study was to investigate the whole region housing potentially microbialites sites, including the one mentioned by Boekschoten (1995), in order to assess their extensive distribution, to observe in situ the possible bioconstructions and to take samples for further studies. Consecutively, our explorations finally led to the discovery of microbialites in various lagoons that are distributed from south to north: la Palme Lagoon, Ayrolle Lagoon, and Portiragnes Lagoon, near Agde (Figs. 1, 2).

Fig. 1. A. Location of the studied area in France; B. Map of the Languedoc-Roussillon coast with the numerous coastal lagoons.
The second objective is to compare the expected observations about current lagoonal mineralized bioconstructions with the data acquired during our work on Sarmatian (Middle Mio-
cene) microbial buildups in Romania (Saint Martin and Pestrea,
1999; Saint Martin et al., 2013) in order to determine if they are,
according Boekschoten (1995), really identical phenomena.

2. REGIONAL SETTING

The part of the studied Mediterranean coast extends from
Leucate, south, to Agde, north (Figs. 1, 2). Most lagoons in this
area are in communication with the sea by passages called
“graus” and supplied with fresh water by a set of versants. The
“graus” of studied lagoons have not been artificially modified
and they are the last to operate naturally on the French Medi-
terranean coast. The lagoons are characterised by the presence
of large seagrass areas and salinity gradients corresponding to
diverse environments: reeds, rushes, sansouires, salt steppes…
The lagoons are of recent formation: the coastal plain was
flooded in Flandrian and gradually filled by Quaternary fluvi-
ual inputs. They are bordered by massive limestone reliefs with
karstic networks that play an important role in fresh or brackish
water supply. The lagoons are located in the northern zone of
Mediterranean climate, with mild winters and hot summers,
relatively low annual rainfall and intense summer drought. Signif-
icant seasonal salinity variations occur, up from 10 g/l of salt
in the winter, after heavy rainfall, to more than 40 g/l in sum-
mer, due to the phenomenon of evaporation. The instability
of the environment is also reflected by changes in the trophic
quality with risk of eutrophication related to excess nutrient
inputs, causing blooms of algae macrophytes and, probably,
cyanobacterial mat development. The average water depth is
between 0.5 and 1 m, but the fluctuations in water level can
be significant (± 0.5 m), depending on the speed and wind
direction. In general, the edges of the lagoons were not very
urbanized. Agricultural activity is occasionally operating, with,
mostly, vine crops causing potential supplementary pressure
on water quality of the nearby lagoons.

3. MATERIAL AND METHODS

Microbialites covering pebbles were sampled in the dif-
ferent sites. Petrographic thin sections of standard thickness
of 30 microns, with a cover-slip, were prepared in order to
perform observations under, both optical, and confocal mi-

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Fig. 2. Coastal lagoons in the studied area at the La Palme (1), southwest of Gruissan (2) and Portiragnes coast (3), Google Earth source.
For optical microscopy, the thin sections were studied under a microscope Zeiss Axioscope 40 at x2.5, x10, x40, x63 and x100 (oil immersion Zeiss) magnifications.

Confocal Laser Scanning Microscopy (CLSM) observations (see Saint Martin and Saint Martin, 2015) were performed on a Leica TCS SP5 microscope, at the University Pierre et Marie Curie at x20, x40 and x 60 magnifications.

Examinations and EDS microanalysis in Scanning Electron Microscope (SEM) were performed using a Hitachi SU 3500 hosted in the Muséum National d'Histoire Naturelle de Paris.

4. RESULTS

4.1. DESCRIPTION OF SITES BEARING MICROBIALITES

4.1.1. La Palme lagoon

The La Palme Lagoon (600 ha) communicates with the sea by the natural grau of Franqui (Fig. 2) whose outlines and openings vary over time (Larue et al., 2009). A general synthesis of hydrobiological functioning and eutrophication state (Wilke and Boutière, 1999, 2000; Souchu et al., 2001; Ifremer, 2002-2012) shows that the watershed consists, mainly, of limestone, partially, occupied by vineyards. The freshwater inflows are, mainly, provided by constant flow resurgences located in the northern part. From the west, mainly temporal streams very dependent on rainfall can partially supply the lagoon with fresh water. The monitoring studies found a good general state of the lagoon which shows no signs of eutrophication (Ifremer, 2002-2012).

Two locations have led to the discovery of microbialites (Fig. 2), one, to the south edge of pond and grassy areas, and, the other, northwest, near the La Palme town. At the south, several sub-lagoon expanses are lined by dense pebbles accumulations (Fig. 3A). The pebbles are easily distinguishable by their black surface color. They also have a brain appearance with small meandering ridges and pustules (Fig. 3B-C). These characteristics correspond to more or less developed and thick microbialitic crusts. Some pebbles are largely covered by the encrusting organisms, such as serpulids, bryozoans and balanids (Fig. 3D).

Fig. 3. Site of La Palme. A. View of the lagoon edge with accumulation of pebbles; B. Pebbles with the microbialitic crust; C. Detail of pebble showing the characteristic cerebroid aspect of the crust surface; D. Pebble covered by the serpulid (se)-bryozoan (br)-balanid (ba) consortium.
4.1.2. Ayrolle lagoon

The Ayrolle lagoon covers a vast area (1400 ha), south of the calcareous Clape mountain between Sigean lagoon and the sea, in direct communication with the sea in its southern part by a natural grau (Fig. 2). Freshwater inputs and freshening are, mostly, associated with contributions from the Camponol lagoon located northwest that serves as a buffer zone in case of flooding in the Aude low plain and small resurgences (Souchu et al., 2001). Its perimeter being barely urbanized, this ecosystem is not receiving effluents; when it does, it is on a small part. South of Gruissan, there is Saint-Martin Island, mainly made of Mesozoic carbonates. South of the island, near the Evêque site, occur small ponds, often partially dried, separated from the main Ayrolle lagoon by a cordon of 100-200 m width and surrounded by cultivated fields of vineyards and Mesozoic limestone (Fig. 4A). These small ponds include the one called Gassot occupying the central part, where there were observed, in fact, the bioconstructions cited by Boek-Schot (1995) and studied in detail in this work. On the shore, pebble concentric accumulations crop out (Fig. 4B). The pebbles are characterised by a black cerebrid-pustular aspect, corresponding to the microbacterial coating (Figs. 4C-E). Some pebbles are colonized by various encrusting organisms, such as serpulids, bryozoans and balanids (Fig. 4F). During the dry season, the pebble accumulations are covered by bleached cyanobacterial filaments (Fig. 4F). The pond bottom is full of monospecific shell accumulation of the cardiid bivalve **Teredo glaucum** (Bruguière, 1789) (Fig. 4G).

4.1.3. Lagoon and channels near Portiragnes

South of the Portiragnes town, a water body forms a small lagoon (Fig. 5A), but with varying dimensions, depending on season, and leading on the beach Portiragnes-Plage. This is, actually, the spreading area of the Ancien Grau du Libron, itself formerly involved in the natural outlet of the Libron coastal river. The entire outlet of Libron went, recently, through development management in the region, especially, for the prevention of floods, mitigating the natural settings. So, there remains a grau currently not directly related to the Libron flow.

In the lagoon, only scattered pebbles (Fig. 5B) occur and the samples were collected during a dry period. These rare pebbles, mainly made of volcanic rocks, probably, resulting from the erosion of nearby outcrops of Pliocene volcanoes, are covered by thin microbialitic crust with pustular surface (Fig. 5C). Behind the pond, several channels connected to the Ancien Grau du Libron are noteworthy for the presence of bioconstructions dominated by worm tubes and bryozoans, also observed and sampled (Fig. 5D-E).

4.2. MICROBIALITE FEATURES

Macroscopically, the microbialite crusts are characterised by their black surface and their arrangement on pebbles or rocky substrate, according to a meandering ridges and pustules pattern.

The combination of different observation techniques used in this work allow complementary information and a good understanding of the microbialitic phenomenon in the studied sites (Figs. 6-8). The principal feature is that pustules and ridges are formed by the successive episodes of development of erected filamentous cyanobacteria and laminated or wavy layers accompanied by Mg calcite micritic mineralization. The erected filaments correspond to **Scytonema**-like cyanobacteria exhibiting a thick lamellar sheath and false branching.

Pustules match digitations not exceeding the length of a few centimeters. Depending on the site, the age of the microbialite and the density of the mineralization, carpet filamentous cyanobacteria appear clearly, sometimes, revealing the preserved remains of sheaths (Fig. 6A-B, 7A-D). Several stages in the mineralization process are distinguished from poorly cemented layers where micrite is relatively sparse (Fig. 8A-C) to dense micrite layers corresponding to a very consistent and solid crust (Fig. 8E-F). When the mineralization is more consistent, the location of cyanobacterial filaments is marked by elongate voids (Figs. 6C-D, 8E-F). Both ways can follow one another in the thickness of the microbialite (Figs. 6A-D, 7A-C, 7E, 8E). In some cases, we clearly noticed that the poorly cemented external part consists of cyanobacterial filaments gathered in tufts explaining, probably, the pustular appearance (Figs. 6E, 7E, 8D). Such a feature is known from **Scytonema** mats (Dupraz et al., 2006; Dupraz et al., 2013). The direct encrustation of pebble substrates may occur through erected filamentous carpets (Fig. 7D), or dense micritic crusts (Figs. 6E, 7E).

Other encrusting organisms providing some cohesion to mineralized microbial mats are present, either on the surface, sealing the construction, in the case of bryozoans (Fig. 6A-B), or incorporated in the microbialite, as noticed in Portiragnes, in the case of serpulids (Figs. 6F, 7C).

These microscopical observations allow to highlight the sequence of events involving settlement phenomena on substrates, development of microbial mats and mineralization. These various events are marked by lines of growth stops, a more or less density of produced micrite and variations in color.

4.3. ASSOCIATED BIOTA

4.3.1. Invertebrates

Three main types of invertebrates with mineralized parts (skeleton or protecting tube) are involved in the constitution of bioconstructions, in various crusts on the pebbles and rocky substrates, or in the formation of microbialite: serpulids, bryozoans and barnacles.

The main contributor to bioconstructions found in the studied coastal area is the polychaete serpulid **Ficopomatus enigmaticus** Fauvel, 1923 (Figs. 3, 4). This invasive species is considered a, reef-builder in estuaries or brackish temperate environments and colonizes many areas in the world in...
Fig. 4. Site of Gassot ponds, southwest of Gruissan. A. General view showing the examined ponds situated between the main lagoon and the land characterised by cultivated fields of vineyards and Mesozoic limestone outcrops; B. View of the pond river with pebble accumulations; C-D. Pebbles with microbialitic coating exhibiting the characteristic cerebroid aspect; E. Detail of pustular cerebroid surface of microbialitic crust; F. Pebble colonized by the serpulid (se) - bryozoan (br) - balanid (ba) consortium and covered by cyanobacterial filaments; G. Monospecific shell accumulation on the pond bottom of the cardiid bivalve Cerastoderma glaucum.
Fig. 5. Site of Portiragnes. A. General view of the pond with few scattered pebbles (arrow); B. Pebble covered by thin microbialitic crust; C. Detail of the pustular crust surface (arrow); D. Internal channels invaded by constructions with serpulid worms and bryozoans; E. Individual construction with serpulid worms and bryozoans; F. Detail of construction with serpulid worms (se) and bryozoans (br).
Fig. 6. Microbialite petrographic thin sections. A-B. Microbialitic digitations of successive filamentous poorly mineralized layers and covered by bryozoans colonies (br), Évêque; C-D. Microbialite pustules on pebble substrate made of alternation of filamentous layers (arrows) and micritic layers, La Palme; E. Crust on pebble substrate (sub) exhibiting a dense wavy micritic layer (m) at the base, followed by a poorly cemented layer of cyanobacterial filaments forming terminal tufts, La Palme. F. Filamentous crust on pebble substrate (sub) encompassing serpulid tubes (se), Portiragnes.
Fig. 7. Petrographic thin sections studied under CLSM (Confocal Laser Scanning Microscopy) technique restituting the autofluorescence signal inside the microbialite. A. Transect along a digitation exhibiting the alternation of finely laminated, wavy and filamentous mineralized layers, (sub: rocky substrate), La Palme; B. Transect along a digitation with partially preserved cyanobacterial sheath (arrow), Evêque; C. Microbialite with successive filamentous crusts encompassing serpulid tubes (se), (sub: rocky substrate), La Palme; D. Thin filamentous crust on rocky substrate (sub), La Palme; E. Filamentous crust ending by tufts (arrow) of cyanobacterial filaments, (sub: rocky substrate), Pradel.
comparable environments with the Mediterranean coastal lagoons (Schwendt et al., 2001; Luppi and Bas, 2002; Schwindt et al., 2004; Nonnis Marzano et al., 2007; Dittmann et al., 2009; Baztérca et al., 2012). The main constructions of *Ficopomatus*, known here under the local name of „cascails“, are especially identified in the channels, but also within the lagoons. The abundant development of the serpulids trigger the repeated destroying of the reefs that threatened to obstruct or fill passage ways or water bodies making difficult the navigation and the hydraulic circulation (Lepareur et al., 2013). However, these reefs, even though they are reinforced by the presence of bryozoan colonies, are not cemented and not directly associated with the microbialite formation. The *Ficopomatus* tubes can only form limited encrustations on the microbialite surface and, more rarely, may be incorporated into the microbialite development (Figs. 6F, 7C).

The role of the serpulid *Serpula vermicularis* Linnaeus, 1767 is reduced to a few encrusting tubes on the pebbles or microbialite surface.

Bryozoans are represented by the species *Conopeum seurati* (Canu, 1928) (determination by P. Moissette, personal communication). The identification as *Electra* species proposed by Boekschoten (1995) needs, thus, to be rejected. Lightly calcified colonies of *Conopeum seurati* are ubiquitous in recoveries on microbialites (Fig. 6A-B) and pebbles, and in the “worm reefs”, but never seen inside the microbial crusts. On even substrata, colonies form regular lacy sheets, while on uneven surfaces colonies become irregular, and the zood size and shape can vary substantially (O’Dea and Okamura, 1999). *Conopeum seurati* is one of the few euryhaline bryozoan species that live, mostly, in desalinated water (estuaries, lagoons, etc), especially found in the Mediterranean coastal lagoons (Maluquer and Barangé, 1987).

Barnacles play an insignificant role in crusts on pebbles and microbialite (Figs. 3, 4). We identified (determination by R.P. Carriol, personal communication) *Amphibalanus amphitrite amphitrite* (Darwin, 1854). It is a cosmopolitan euryhaline species of tropical to warm temperate waters that tolerates brackish water (Kerckhof, 2002), but does not survive for a very long time in desalinated water (Newman and Abbott 1980). Species usually inhabit the intertidal zone, often, in brackish water on rocks, shells, in the mangroves, and, also, on vessels (Henry and McLaughlin, 1975). This species is also very common in polluted areas (Kerckhof and Cattrijse, 2001).

4.3.2. Diatoms

The importance of diatoms associated with microbial mats and microbialite is now well proven and supported by numerous recent works (for more details, see Saint Martin and Saint Martin, 2015). Because of the fragility of their frustules, conservation is often poor and determination can be difficult. Moreover, diatoms are partly represented by species of very small size, also, making them difficult to specific determination.

The examination under optical microscopy allowed the observation of diatom frustules that are uniformly distributed on the microbialite surface. There is a monospecific population of *Rhopalodia gibberula* var. *musculus* (Kützing) Cleve-Euler 1952. Some frustules seem to be attached to the “substrate” (Fig. 9A). Indeed *Rhopalodia* is known as epilithic or epiphytic genus (Round, 1984).

Inside the microbialite, the dense material hides information concerning the presumed presence of diatoms. So, in order to access these data, microbialite broken samples were investigated under SEM. Effectively, this inside observation revealed numerous diatom frustules belonging to genera *Diploneis*, *Navicula* and *Nitzschia*, the latest being the most frequent (Figs. 9B-F). Attentive examination depicts the relation between diatoms and cyanobacteria. Very minute frustules (5 μm) of *Nitzschia* cf. *frustulum* (Kützing) Groun in Cleve & Groun 1880 and of *Navicula* cf. *cryptodepha* var. *veneta* (Kützing) Rebenhorst 1884 seem to be attached to the intertwined cyanobacteria filaments (Fig. 9B). The frustules of *Diploneis* cf. *bombus* (Ehr.) Ehrenberg 1853 and *Nitzschia* sp. are progressively coated by EPS (Extra Polymeric Substances) cyanobacterial biofilm (Fig. 9B, E) going up to the complete embedding. The electron micrographs document the epiphytic and interstitial lifestyle of the entrapped diatoms into the microbialite and provide, also, information on different stages of diatom preservation. Some frustules are still well to moderate preserved, such as *Diploneis* cf. *bombus* (Fig. 9B) and *Nitzschia* cf. *frustulum* (Fig. 9D), respectively. Other frustules, like *Diploneis* sp., appear to be corroded (Fig. 9C). We also observed imprints of frustules preserved in a more calcified material; morphological details are inscribed with more or less fidelity (Fig. 9F).

The recorded diatom community is typical for the microphytobenthos in shallow coastal waters that may be subject to salinity variations (Witkowski, 1990, 1991; Sundbäck and Snoeijis, 1991; Vilbaste et al., 2000).

5. DISCUSSION

5.1. MICROBIALITE FORMATION AND ENVIRONMENTAL VARIATIONS

The microbialites covering the pebbles or the rare *in situ* bedrock in the studied area show an aspect and formation processes exactly like those highlighted in Sardinia by Saint Martin and Saint Martin (2015). However, the microbialites described in the present paper are less developed and remain limited to a few centimeter thick, at most. So, according to Saint Martin and Saint Martin (2015), the observed lamination more or less clearly expressed in the French microbialite reflects the irregular temporal sequence of environmental changes, such as impoundment or, at the contrary aerial exposure, the season succession (water temperature, wetting and drying), the nutrient supplies, the salinity fluctuations,
Fig. 8. SEM views of the microbialite. **A.** General view of relatively poorly mineralized microbialite constituted by dense erected filaments, Portiragnes; **B.** Detail of filamentous microbialite with micrite cement (arrow) between the filaments, Portiragnes; **C.** Detail of filamentous microbialite with micrite cement (arrow) between the filaments, Evêque; **D.** Cyanobacterial erected filaments forming tufts at the crust surface, La Palme; **E-F.** Filamentous crust with elongate voids indicating the original presence of cyanobacterial filaments, La Palme.
Fig. 9. Diatoms associated with the microbialite. A. Thin section showing frustules of the diatom Rhopalodia gibberula var. musculus (Kützing) Cleve-Euler 1952 colonizing the surface of microbial mat (arrows); B. Frustules of Diploneis cf. bombus (Ehr.) Ehrenberg 1853 between intertwined cyanobacterial filaments and minute frustules of Nitzschia cf. frustulum Kützing) Grunow in Cleve & Grunow 1880 (white arrows) and Navicula cryptcephala var. veneta (Kützing) Rebenthor 1864 (black arrow); C. Corroded frustule of Diploneis sp.; D. Frustules of Nitzschia cf. frustulum; E. Partially dissolved frustules of Nitzschia sp. entrapped in degraded Extra Polymer Substances (EPS); F. Casts of diatom frustules (arrows) in micrite.
favoring the development of successive specific microbial mat layers and, then, their mineralization. The thinness of the French microbialites shows that it is a limited phenomenon in time, in this region. This led to question about the mineralization processes in action (Saint Martin and Saint Martin, 2015) and, also, their initiation. We can remark that there are no microbialites in the nearly coastal open marine conditions or in the lagoons center. Monitoring studies are needed to determine the exact time of microbialite formation regarding the lagoon water level, the rainfalls impact and the influence of marine inputs through the passage ways (“graus”). But, unfortunately, we cannot exclude that the relevant phenomena are very discontinuous in time and, therefore, difficult to define precisely. Other parameters are also considered by Boekschoten (1995), such as the very low relief of the French coastal region and the faint ecological gradient keeping the potential reef-builders spatially isolated. Under these conditions, he suggested that the stromatolite/bryozoan/barnacles/brachiozoans “reefs” would develop along Mediterranean coasts as soon as favourable conditions (varied lagoon bottom topography, rise of brackish water level, and decrease of clastic influx) arise. However, our field investigations show that the stromatolite/bryozoans/brachiozoans “reefs” do not really exist, but, on the one hand, the microbialitic crust with some limited serpulids/bryozoans/barnacles encrusting setup and, on the other hand, the serpulids/bryozoans “reefs” without microbialites!

An additional question could be related to the location of microbialites relative to confinement. We can observe that the Gassot site, but, also, to a lesser extent, that of La Palme correspond to a shift away from direct exchange areas with the marine environment. Given an identical situation in Sardinia, where direct interactions with the marine environment are only temporary or occasional (Saint Martin and Saint Martin, 2015), we can ask whether the concept of confinement in paralic realm may be applied, not only to common biota (invertebrates, fishes…), but also to the consortium of microbial organisms needed for the microbialite formation.

5.2. A MODERN ANALOGUE TO ANCIENT BIOCONSTRUCTIONS?

According to Boekschoten (1995) the stromatolite-serpulids-bryozoans consortium, like those observed in the Gruissan area, have built reefs from the Carboniferous times, always in marginally marine environments under shallow hypo- or hypersaline water, particularly with the massive bi-oherms along the Miocene coast of the Paratethys in Poland and Russia. Similarly, Bijma and Boekschoten (1985) consider that the bryozoan reefs and stromatolite observed on the Dutch coast in an artificial men-made environment have fossil counterparts with the Miocene bryozoan/stromatolite reefs in the Paratethys. These reefs were specifically established during the Sarmatian (Middle Miocene), throughout the whole Paratethys. They are well represented in Romania, where we could perform observations useful for comparisons (Saint Martin and Pestrea, 1999; Saint Martin et al., 2013). Indeed, in the present lagoons and in the Miocene bioconstructions, we generally found partly similar components, especially with brackish affinities, but this direct comparison requires a closer examination of the specific characteristics of each situation. In addition, the Sarmatian is, in fact, characterised by two successive periods, each with specific types of constructions and contents. In the lower part of the Sarmatian, during Volynian, thick buildings containing interlinked serpulid colonies and abundant micrite are mostly developed. Later, during Bessarabian, there are less voluminous bioconstructions consisting of small reefs made by dominant bryozoans associated with foraminifers (nubecularids) and calcareous algae, the microbialite intervention being restricted. Table 1 summarizes the key features and shows disparities that need to be discussed.

The first observation concerns the microbialite size. Thus, macro-oncoids observed in Sardinia and the microbial crusts from the lagoons of France remain quite small, compared to the, sometimes, very important, Sarmatian reefs (Fig. 10A). This is due, of course, primarily to the fact that current microbialite develop over a short period of time in a very shallow environment (few centimeters to decimeters deep). The distribution of current microbialite is very punctual and their extension is limited. Yet, all ponds laced in identical conditions do not necessarily host microbialite formations. Sarmatian reefs show, on the contrary, a much more significant scope and a very broad distribution (Bucur et al., 1992; Pisera, 1996; Jasionowski, 1996; Jasionowski et al., 2002; Jasionowski et al., 2003; Jasionowski, 2006), sometimes almost continuous in the case of the Moldavian platform (Saint Martin and Pestrea, 1999; Saulea, 1995). Another notable difference is the absence in the current constructions of calcareous algae and encrusting foraminifera as nubecularids, the microbialitic features and the associated biota (Tabl. 1). Finally, the microbialitic fabric of Sarmatian reefs is significantly different exhibiting mainly clotted and/or peloidal micrite (Fig. 10B, C) and, also, fibrous calcite rims around the serpulid tubes (Bucur et al., 1992; Saint Martin and Pestrea, 1999) (Fig. 10D).

In summary, despite some similarities, it is difficult to establish strict analogies between biosedimentological events, separated in time and space, as suggested by Bijma and Boekschoten (1985) and Boekschoten (1995). Time, space and bio-physics-chemical features have to be considered. The current microbial structures were developed in limited time and particularly changing environmental conditions, in small water bodies with enormously variable hydrology, determined, primarily, by floods, caused by sea storms and/ or heavy rains and subsequent summer desiccation. On the other side, the Miocene reef structures are part of a longer biological and paleogeographic history marked by specific physico-chemical conditions, including alkalinity which seems to have played a significant role, ruling in a sea of great extension (Harzhauser and Piller, 2004; Piller and Harzhauser, 2005; Jasionowski, 2006; Harzhauser and Piller, 2007; Piller et al., 2007).
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<th>MAIN FEATURES</th>
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<th>MIocene (Sarmatian) MICROBIALITES FROM ROMANIA</th>
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<td>Within a vast intern sea inherited of ancient Tethys</td>
</tr>
<tr>
<td>Main distinguishable microbial composition</td>
<td>Filamentous (Scytonema) and coccoid cyanobacteria</td>
<td>Filamentous (Scytonema) and coccoid cyanobacteria</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Diatoms</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Diatoms</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No clearly identifiable microbial remains</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No clearly identifiable microbial remains</td>
</tr>
<tr>
<td>Macroscopic features</td>
<td>Oncolite or crusts around pebbles</td>
<td>Masses forming variously sized reliefs (from limited patches to bioherms)</td>
</tr>
<tr>
<td></td>
<td>Digitations</td>
<td>Digitations</td>
</tr>
<tr>
<td></td>
<td>Laminations</td>
<td>Laminations</td>
</tr>
<tr>
<td></td>
<td>Surface with a pustular cerebroid aspect</td>
<td>Surface with a pustular cerebroid aspect</td>
</tr>
<tr>
<td>Associated biota</td>
<td>Serpulids and bryozoans only as encrusters</td>
<td>Serpulids and bryozoans incorporated in the buildup</td>
</tr>
<tr>
<td></td>
<td>Few serpulid tubes incorporated in the microbialite</td>
<td>sparse nubecularids and calcareous algae</td>
</tr>
<tr>
<td></td>
<td>Consortium of bryozoans, foraminifers (nubecularids), serpulids and calcareous algae</td>
<td></td>
</tr>
<tr>
<td>Other features</td>
<td>Presence of small reefs with serpulids and bryozoans without microbialite in confined lagoons and channels</td>
<td>Oolitic deposits</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Abundant oolitic deposits</td>
</tr>
<tr>
<td>Main surrounding biota</td>
<td>Bivalves (cardiids)</td>
<td>Bivalves (cardiids, mytilids) . . .</td>
</tr>
<tr>
<td></td>
<td>barnacles</td>
<td>Gastropods (riossids, potamids)</td>
</tr>
<tr>
<td></td>
<td>crabs</td>
<td>Diversified bivalve fauna</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Diversified gastropod fauna</td>
</tr>
<tr>
<td>Bioesdimentology</td>
<td>Dense micritic product organized in fine laminations or wavy layers</td>
<td>Dense micritic product organized in fine laminations or wavy layers</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Clotted, peloidal micrite.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Few small and isolated stromatalites</td>
</tr>
<tr>
<td></td>
<td></td>
<td>laminated microbial mats</td>
</tr>
<tr>
<td></td>
<td></td>
<td>composite crusts</td>
</tr>
</tbody>
</table>
6. CONCLUSION

We have shown that the phenomenon of microbialite formation in the context of lagoons or smaller ponds was quite widespread in the Mediterranean, at least, in the state of current knowledge in Sardinia and France. It is likely that many other lagoons show similar features that can, therefore, be regarded as specific in their composition and their appearance. Systematic research must, now, be undertaken in all Mediterranean areas susceptible to host this type of microbialite.

Contrary to common ideas, current microbialites constructions associated or not with different metazoans cannot always be simply transposed to explain microbialitic old buildups, even for fairly similar time as the Miocene. It is absolutely necessary to take into account, not only the components of these microbialites, but all the parameters that govern aquatic environments and, also, the environments characterised by major rapid fluctuating conditions.

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