What makes seep carbonates ignore self-sealing and grow vertically? The role of burrowing decapod crustaceans

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ABSTRACT

The mechanisms that govern the vertical growth of seep carbonates were deciphered by studying the sedimentary architecture of a 15 m-thick, 8 m-wide column of limestone encased in deep-water marl in the Middle Callovian interval of the Terres Noires Formation in the SE France Basin. The limestone body, also called “pseudobioherm” records intense bioturbation, with predominant traces of the Thalassinoides/Spongeliomorpha suite, excavated by decapod crustaceans. Bioturbation was organized in four tiers. The uppermost tier 1 corresponds to shallow homogenization of rather soft sediment, tier 2 to pervasive burrows dominated by large Thalassinoides having walls locally bored by Trypanites representing tier 4. The diagenetic cements filling the tier-3 Thalassinoides are arranged in two phases. The first cement generation constitutes a continuous rim coating the burrow wall having consistent δ13C values near -8‰ to -12‰, indicative of bicarbonate originating from the anaerobic oxidation of methane. In contrast, the second cement generation is dominated by saddle-dolomite precipitated at temperatures >80°C, at a time when the pseudobioherm was deeply buried. The fact that the tubes remained open until deep burial means that vertical fluid communication was possible over the whole vertical extent of the pseudobioherm up to the seafloor during its active development. Therefore, vertical growth was fostered by this open burrow network, providing a high density of localized conduits through the zone of carbonate precipitation, in particular across the sulfate methane transition zone.

Burrows prevented self-sealing from blocking upward methane migration and laterally deflecting fluid flow. One key aspect is geometric complexity of the burrows with numerous subhorizontal segments that could trap sediment shed from above and hence, prevent their passive fill.

Key Words
Seep carbonates; bioturbation; decapod crustaceans; permeability; focused fluid flow; burrow connectivity
1. Introduction

Seep carbonates are produced by the anaerobic oxidation of methane (AOM) or other heavier hydrocarbons coupled with seawater sulfate reduction (Boetius et al., 2000; Orcutt et al., 2010; Zwick et al., 2018). Anaerobic hydrocarbon oxidation generates bicarbonate that may precipitate with seawater cations into carbonates like aragonite, calcite, or dolomite. The other product of this process is (hydrogen) sulfide that combines with available iron to precipitate iron sulfides while the excess remains in solution and is potentially re-oxidized at the seafloor (Jourabchi et al., 2005; Blouet et al., 2021a). In settings with upward advection of hydrocarbons, the depth of the reaction front, the so-called sulfate-methane transition zone (SMTZ), typically lies a few centimeters below the seafloor (Regnier et al., 2011), where a strong redox gradient favors the settlement of chemosymbiotic macrofauna (Kiel, 2010). Seep carbonates typically appear as concretionary bodies exhibiting a wide diversity in shape and size ranging from isolated nodules a few centimeters in diameter (e.g. Haas et al., 2010) to massive mound-shaped structures tens of meters in diameter (e.g. Kauffman et al., 1996). They are commonly associated with other fluid expulsion features such as pockmarks (Ho et al., 2018a, b). Oceanographic observations demonstrated that hydrocarbon seepage at the seafloor is a ubiquitous phenomenon in the world's oceans (Judd and Hovland, 2007), and seep carbonates preserve this elusive phenomenon in the rock record. As such, seep carbonates mark the outlet of hydrocarbon migration pathways underneath, such as permeable layers, faults, fluid chimneys, hydraulic fractures, etc., and they have been used to reconstruct fluid-flow mechanisms in the shallow sub-seafloor sediment (Hovland, 1982; et al., 1985, 1994; Gay et al., 2003; Mazzini et al., 2003; Agirrezabala et al., 2013; Ho et al., 2016, Blouet et al. 2021b). Seep carbonate systems are commonly observed on seismic data as vertical or sub-vertical stacks of lenticular anomalies, which document that seep sites can be long–lived features (Kauffman et al., 1996; Hovland and Judd, 1988; Plaza-Faverola et al., 2011). In particular, the morphology and vertical variation of amplitude anomalies are useful to reconstruct qualitatively the history of fluid leakage intensity (Ho et al., 2012).

One key process leading to lateral development of seep carbonates is “self-sealing” as conceptualized by Hovland (2002); concretions growing at seep sites block vertical migration along the initial venting domain and promote its lateral shift, which leads to precipitation of laterally extensive slabs of methane-derived carbonate. Self-sealing has been interpreted from oceanographic data in the Black Sea (Naudts et al., 2008) and in outcrop by Agirrezabala et al. (2013). Gay et al. (2020) interpret a similar phenomenon at the scale of individual carbonate lenses in the Oxfordian pseudobioherms of Beauvoisin. On the pore-scale, vertical gas migration is blocked by precipitation of seep carbonates, leading to a downward growth of concretionary crusts in the order of 1–10 cm (Bayon et al., 2009; Greinert et al., 2002; Peckmann et al., 2002). In contrast to the downward growth pattern, Liebetrau et al. (2014) have documented continuous upward growth of seep carbonates over >3 m. Lateral gas deflection due to clogging of the vertical pore network has also been reported from continental slope settings with stable gas hydrates as their growth can block fluid migration pathways causing upslope shift of the gas leakage zone (Casenave et al., 2017). Hydrocarbon leakage indicators appear in a number of basins as stacks of small, 20–200 m-diameter subcircular or elongate seismic amplitude patches described as “pipes” (Bünz et al., 2003; Berndt, 2005; Petersen et al., 2010; Løseth et al., 2011) or “chimneys” (Heggland, 1998; Ligtenberg, 2003; Løseth et al., 2009; Hustoft et al., 2010; Ho et al., 2016). Such stacks of seismic amplitude patches can exceed 1 km in thickness (Løseth et al., 2011). They highlight situations where self-sealing is restricted to patches being 20–200 m in diameter whereas vertical growth predominates on a large scale (Plaza-Faverola et al., 2011). Seep carbonate
bodies at all scales thus reflect a balancing between lateral and vertical growth (Hovland et Judd, 1988). Self-sealing has been invoked as the dominant process causing lateral growth, but what governs vertical growth of a seep carbonate body?

It is the purpose of the present study (1) to describe a well-exposed 15-m-thick, 8-m-diameter columnar carbonate body (initially described as a “pseudobioherm”) and its sedimentary/diagenetic architecture, (2) to decipher the cm-to-m-scale processes fostering the vertical growth/aggradation of the pseudobioherm, and (3) to emplace this small-scale architecture in the frame of the permeability field in and around the pseudobioherm.

2. Geological Setting

The studied mid-Callovian pseudobioherm is located near the village of Aurel, about 50 km to the ESE of Valence city in SE France. During the Callovian, the area was part of the SE France Basin, a ca. 150 km-wide embayment along the northern margin of the Alpine segment of the opening Tethys Ocean (Fig.1A–B).

2.1. Geodynamic context

The SE France Basin resulted from the Triassic rifting of the nascent Alpine Tethys (Lemoine et al., 2000; Masini et al., 2013). Triassic transgressive deposits above the peneplaned magmatic and metamorphic Variscan basement and Permo-Carboniferous basins entrenched therein consist of shallow-water siliciclastics and evaporites. The evaporites acted as decollement level during the structuring of the basin. Deepening of the depositional environment during the Lower Jurassic coupled with large lateral thickness variations implies synsedimentary tilted block tectonics linked to the paroxysmal phase of rifting (Lemoine et al., 1986). More uniform Bathonian to the Tithonian deposits exceeding several kilometers in thickness are attributed to the onset of ocean spreading and associated thermal subsidence of the continental margin (Fig.1B–C). During Triassic and Jurassic times, the SE France basin was elongated along a NE-SW axis inherited from Variscan structures, whereas during the Cretaceous the basin narrowed and acquired an E-W elongation due to the centripetal progradation of carbonate platforms. E-W folding of the basin, initiated during the Aptian-Albian within a N-S compressional regime induced by the Pyrenean-Provencal orogeny, led to emersion during the Santonian. From the Oligocene onward, the Alpine orogeny overprinted previous structures by E-W compression.

2.2. Structural and stratigraphic setting

A 50 km long, E-W oriented seismic section, crossing through the studied outcrop at Aurel, allowed the description of the local structural setting from the edge to the center of the basin (Fig. 1A; Roure et al., 1992; Wannesson and Bessereau, 1999). About 10 km east of Aurel, these authors interpreted an offset in the basement reflection as evidence for two major westward-dipping normal faults rooted in the Paleozoic substratum. Early Jurassic tilted blocks are visible in the western part of the section, whereas they have been reactivated as reverse structures during compressive phases in the eastern part. The outcrop of Aurel is located in the middle of a large-scale domal structure formerly known as the ‘Aurel anticlinorium’ (Artru, 1972)

An exploration borehole drilled to a depth of 2800 m less than 1 km away from the outcrop provided information on the stratigraphic column below the outcropping Callovian strata, down to Toarcian-Aalenian deposits (COPEFA, 1967) (Fig. 1C–D). The uniform Toarcian-Aalenian 'Marly Lias' has been partially penetrated
for 1.5 km; the overlying 600 m thick ‘Calcareous Dogger’ is a limestone-marl alternation dated Bajocian to Bathonian pro parte. The Bathonian to Oxfordian Terres Noires Formation is a uniform package of shales about 1.9 km thick in the Aurel area; it grades upward into limestone-marl alternations locally known as ‘Argovian’ facies (Artru, 1972).

2.3. Potential oil and gas sources

Potential hydrocarbon source rocks of the SE France Basin have been studied in a report by the Institut Français du Pétrole (IFP, 1982a, b, c) and later by Wannesson and Bessereau (1999) and Mascle and Vially (1999). In stratigraphic order, the potential source rocks are:

- Carboniferous coals in post-Hercynian basins, known from outcrops and coal mines around the SE-France Basin but speculative in the vicinity of Aurel.
- The “Lias marneux” Formation (Marly Lias) is >1500 m thick in the Aurel borehole and contains on-average 0.6% TOC (Wannesson and Bessereau, 1999).
- The Terres Noires Formation yielded TOC values of 0.5–1%. Common occurrence of oil in septarian concretions indicate that some hydrocarbons were generated in the Terres Noires Formation (Montenant et al., 1968; Oddou 2004)

2.4. Seep carbonates in the Jurassic SE France basin

Lenticular carbonate bodies scattered in the Terres Noires Formation have been intensively studied over the past 60 years, starting with Orgeval and Zimmerman (1957). Artru and Gauthier (1966) and Artru (1972) proposed a first interpretation as sponge bioherms based on the abundance of disarticulated spicules. The lack of in-situ frame building organisms and the absence of evidence for relief above the seafloor, however, led Flandrin (1974) to question this hypothesis. To account for this uncertainty, Gaillard et al. (1985) and Rolin (1987) coined the term ‘pseudobioherm’, defined as “carbonate anomalies of early diagenetic origin, characterized by the predominance of endobenthic organisms”. Rolin (1987), Rolin et al. (1990) and Gaillard et al. (1992) noticed similarities between the numerous assumed chemosynthetic bivalves often associated with the pseudobioherms and the fauna discovered a few years earlier at cold hydrocarbon seeps (Suess et al., 1985). Since then, the hypothesis of hydrocarbon seepage has been developed for the outcrop of Beauvoisin, located ca. 50 km to the south of Aurel where 19 pseudobioherms are vertically distributed over 350 m inside an 800-m diameter perimeter (Gay et al., 2019, 2020). Gay et al. (2020) interpreted the Beauvoisin cluster as a long-lived giant pockmark and assigned the multiple generations of pseudobioherms to self-organized lateral migration of the fluid conduit over time.

3. Material and Methods

The paleogeographic, structural and stratigraphic context of the Aurel pseudobioherm has been determined at regional scale based on the 1/50.000 geological map (Flandrin, 1974) and the Aurel borehole (COPEFA, 1967). At basin scale, the context has been evaluated using seismic data (Roure et al., 1992; Wannesson and Bessereau, 1999), isopach and facies maps (Debrand-Passard, 1984), and a few exploration boreholes compiled by Wannesson and Bessereau (1999).
In total, 20 rock slabs were cut and polished, 6 to 25 cm long (average 13 cm), and 6 of them were oriented. They were studied in natural light and 365 nm ultraviolet light (UV) emitted by a Vilber VL-6 (Eberhardzell, Germany). The 3-D geometry of tubular structures was approached by comparing closely spaced sections such as those on opposite sides of 3 mm-wide sawcuts. Thirty thin sections were investigated microscopically using plane and cross-polarized light, cathodoluminescence (CL) and epifluorescence UV (epi-UV). The CL was generated by a CITL system (Hatfield, United Kingdom), model CCL 8200 ink4 (12kV, 450 mA). The epi-UV was emitted by a Leica (Wetzlar, Germany) EL 6000-light mounted on a Leica DMRXP microscope (bandpass 340–380nm).

For X-ray diffraction (XRD), 27 samples were crushed manually in an agate mortar. The powders were analyzed with a Rigaku (Tokyo, Japan) Ultima IV diffractometer system equipped with a Cu X-ray tube, operated at 40 kV and 40 mA, and with a D-Tex linear detector. Scans were run from 5° to 70°2θ, with a step interval of 0.01°2θ and a goniometric speed of 20·120s⁻¹. The identification of all minerals was performed using the Rigaku PDXL2 software package and the ICDD Powder Diffraction File 2014 database (International Centre for Diffraction Data).

For stable oxygen and carbon isotope analyses, 40 samples of carbonate cements were selected. They were taken from polished blocks and rock chips using a handheld microdrill under the alternation of natural and UV light. Due to the size of the drilling bit, only thick enough cements could be sampled individually, whereas grainstone was taken as bulk, including cement and grains. Samples were analyzed using a Kiel III automated carbonate preparation device coupled to a Finnigan MAT (Bremen, Germany) 252 isotope-ratio mass spectrometer. Carbonate material was reacted with 100% phosphoric acid for 10 min at 70°C. The produced CO₂ was then passed through the isotope-ratio mass spectrometer for masses 44, 45, and 46 measurements alternately with the measurement of a calibrated reference CO₂ gas. Instrumental precision was monitored by analysis of NBS 18, NBS 19, and LSVEC reference material. Precisions are –0.05‰ for carbon and –0.14‰ for oxygen. All isotope data are given relative to the Vienna Peedee Belemnite standard (V-PDB).

4. Results and interpretation

Four pseudobioherms have been identified less than 1 km away from the Aurel borehole (Artru, 1972, Fig.1D). Three of them near the Vaunière ruin are covered by vegetation and the relationships to the surrounding marls are not visible. In contrast, the pseudobioherm at “La Touche” is well exposed in lateral continuity with its background sediments; it is the focus of this paper. The host sediment consists of an alternation of mid-Callovian marls and thin platy limestone beds dipping 15° towards ESE (log in Fig. 2). Two massive limestone bodies are present ca. 30 m apart along a small thalweg roughly oriented E-W. The upper one, on the northern bank of the thalweg, is a 7–10 m-wide columnar body that interfingers with background marl-limestone alternations. Only 15 m is exposed over vertically above ground. The top is truncated by erosion and its lower part is cut by a low-angle fault (strike 040, dip 38W) filled by a cm-thick calcite vein. The vein shows crude striations with an azimuth 095–125, without direct evidence for normal vs. reverse movement. The lower exposure is a westward-dipping subcircular section ca. 8 m in diameter on the southern bank of the same thalweg. It exposes the same facies as the upper one over 6 m. The relationships between the limestone body and its host sediments are poorly exposed and its basal contact with the marls is not visible. The low angle and the orientation of the fault, perpendicular to the trend of the exposures, suggest that these are two parts of the same columnar body offset by a reverse fault.
4.1. Architecture of the pseudobioherm

Mapping of the southern flank (Fig. 3A–B) and eastern flank (Fig. 3C–D) shows that the columnar pseudobioherm is made up of stacked lenses. Individual lenses are a few tens of cm thick and most of them pinch out within the massive body. They are separated by cm-thick marl layers with or without nodules or they merge laterally or vertically. The center of an individual lens generally consists of massive limestone, which grades outward into nodular limestone that in turn splits laterally into several wedges grading into nodular marl. Nodules feather out into background marl over 1–2 meters. Lenses in lateral continuation with limestone beds in the background sediments have the greatest lateral extension and are the most indurated. Overall, the pseudobioherm can be subdivided vertically into 3 units comprising from base to top:

- **Unit 1.** At the base, it is partially truncated by a fault. It mostly exposes nodular limestone grading laterally into nodular marls. Locally some limestone lenses tend to be stacked vertically. The contact to Unit 2 above is defined by an abrupt lateral offset by 1–2 meters westward.
- **Unit 2.** The lowermost bed continues laterally into a prominent limestone bed in the background deposits classified as ‘marker bed A’. The middle portion of Unit 2 is made up of lenses consisting of nodular to massive limestone facies, locally organized in vertical stacks. The top of Unit 2 becomes increasingly massive. The transition from Unit 2 to Unit 3 is, once again, affected by an apparent offset by 1–2 meters northward, similar to the transition from Unit 1 to Unit 2.
- **Unit 3.** The lowermost bed of Unit 3, classified as ‘marker bed B’, continues laterally into a limestone bed of the host sediment and is particularly massive. It is almost entirely made up of massive limestone along the oblique southern flank of the pseudobioherm.

4.2. Facies of the pseudobioherm and background sediments

4.2.1. Facies 1 (F1): Background, marl-limestone alternation

In the studied area, the typical middle Callovian facies consists of marls with intercalated limestone beds, a few cm to 20 cm thick, regionally known as “Marnes à plaquettes rousses” (marl with platy, rusty weathered beds) (Artru, 1972; Flandrin, 1974) (Fig. 4A). Individual limestone strata commonly show plane bed lamination and current ripples, as well as occasional burrows at the base of a bed (Fig. 4B). The only common macrofossils are shells of *Bositra sp.*, a bivalve typically associated to dysoxic settings (Meesook et al., 2009).

In thin sections, both the background marl and the limestone beds exhibit a dominant wackestone fabric having a similar grain content composed of microfossils and silt-grade quartz representing a calcilutite. They differ with respect to the matrix, being shaly in marls (Fig. 4C) and microsparitic in limestones (probably recrystallized micrite, Fig. 4D). Laminae in limestone beds appear as mm-thick alternations of wackestone and packstone (Fig. 4E). The most abundant microfossils are spheres with a diameter of ca. 100 μm, likely calcispheres and/or recrystallized radiolarians, and sponge spicules (Fig. 4F). Echinoderm ossicles are rare. Most microfossils have been recrystallized into sparite. Sinuous filaments, up to several hundreds of μm in length and less than 10 μm wide correspond to the structures identified as fragments of exfoliated shells of *Bositra sp.* (Negra et al., 2011) (Fig. 4C).
4.2.2. Facies 2 (F2): Nodular marl

This facies comprises the transition between background sediments and the pseudobioherm limestone body (Fig. 5A). It consists of micritic limestone nodules embedded in a marly matrix exhibiting a matrix/marl-supported floatstone texture. The most common nodules are isolated prolate ellipsoids with the long axis parallel to stratification. The nodules are 1–2 cm in diameter and up to 5 cm in length. Some show rod-like, straight, or curved protuberances (Fig. 5B). Nodule density increases towards the limestone body. At higher concentration, adjacent nodules commonly aggregate into peanut-shaped clusters of two or three, in a pattern similar to that described in Beauvoisin by Gay et al. (2019, 2020). In thin sections, nodules show a wackestone texture containing the same grain types as the encasing marl and limestone beds described above, such as microfossils and detrital silt. As peloids represent a key component of F4 (see below), we paid particular attention to their possible presence. Neither direct observation, nor fluorescence or cathodoluminescence provided any evidence of their (former) presence in F2. Nodules are weakly fluorescent under UV light (Fig. 5C–D). F2 shows a gradual transition to facies F3 comprising nodular limestone as the concentration of nodules increases towards the axis of the pseudobioherm and individual nodules may become amalgamated. A threshold of ~50% nodules marks the limit between F2 that is sensitive to weathering as being a nodular marl and F3 that remains rather massive and less prone to weathering. The different weatherability of the two facies was used for pseudobioherm facies mapping of hardly accessible zones (Fig. 3).

4.2.3. Facies 3 (F3): Nodular limestone

Across the transition of F2 to F3, nodules become more frequent and the matrix between the nodules is strongly indurated and shows pervasive rusty speckles (Fig. 6A). The nodules are weakly fluorescent in a non-fluorescent matrix as in F2 (Fig. 6B). In contrast to nodular marl, the nodular limestone represents a nodule-supported rudstone texture. In thin sections, the rusty speckles consist of euhedral crystals of saddle dolomite about 100 μm in size, locally replacing and overgrowing carbonate bioclasts (Fig. 6C). The rusty color of the crystals is due to oxidation of iron along cleavage planes and gives the matrix its speckled macroscopic appearance.

4.2.4. Facies 4 (F4): Massive limestone

4.2.4.1. Macroscopic observations and macrofauna

Facies F4 consists of gray to brownish limestone that typically interfingers with the nodular limestone. In contrast to the background sediments, no primary sedimentary structures or laminations have been observed, although laminated limestone beds of the surrounding host sediment occur in continuity with prominent lenses of the pseudobioherm. Severe weathering hinders observation of facies details in the field, except on a few isolated surfaces. One of these, ca. 30 cm x 70 cm, exposes the feathering out of F4 into marker bed A that extends farthest away from the axis of the pseudobioherm (Fig. 7A). The surface shows three articulated bivalves (Fig. 7B–D). In addition, a single, internal mold of a 6 cm-long articulated specimen found on the margin of the pseudobioherm was identified as a lucinid, possibly Beauvoisina carinata (Kiel et al., 2010) although it seems to be more asymmetrical than the type specimen and the diagnostic carinate lunule could not be recognized on the mold (Fig. 7E–F). A few small gastropods were also observed in the outcrop and in cut samples (Fig. 7G), indicating that benthic mollusks were relatively common and diverse in F4. Facies 4 commonly shows ~5-mm-diameter cement-filled tubular structures with the characteristic branching pattern of Thalassinoides/Spongeliomorpha (Fig. 7H).
4.2.4.2. Fabric and bioturbation

The fabric is best characterized by using combined plane and UV light on cut sections. Macroscopically, most samples show a complex mosaic of patches, which under UV light appear to consist of low-fluorescence micrite and calcarenite exhibiting various levels of fluorescence (Fig. 8). At hand lens magnification, the calcarenitic limestone consists of peloid grainstone having fluorescent cement. The proportions of micrite and calcarenite vary, the former ranging from 20–50%, the latter from 80–50%. Near the transition to facies F3, where micrite is enriched (~50%) it forms continuous domains, whereas calcarenitic material constitutes a network of 2–3 cm-wide patches surrounded by a darker, commonly fluorescent halo (Figs. 8C–D, 9A–C). This texture evidences pervasive bioturbation, with two populations of burrows. Calcarenitic domains 2–3 cm in width are referred to as “large burrows” and 0.5–1.5 cm-sized ones as “medium burrows” in the following. In addition, numerous mm-diameter burrows (“small burrows”) filled by fluorescent cements are scattered throughout the section. In calcarenite-rich samples, micrite occurs as low/moderate-fluorescence patches, 1 mm to 2 cm in diameter, angular to subrounded, commonly surrounded by a dark halo. The micrite domains are interpreted as remnants of the initial sediment that were not reworked by burrowing. They are described as so-called ‘pheno-intraclasts’ floating in the dominant peloid grainstone (Fig. 8E–G), in contrast to true intraclasts defined as reworked particles. The overall fabric of calcarenite-rich areas reflects pervasive bioturbation, including but likely not restricted to the large burrows.

a) Large burrows

The margins of large burrows, as defined by their dark halo, are rather irregular. The geometrical arrangement of the large burrows suggests that they constitute a complex branched network (see Cunningham et al., 2012). They are filled with sediment, typically peloid grainstone/wackestone identical to that of the background indicative of passive fill. Large burrows have variable intensity of fluorescence, with peloid grainstone being the most fluorescent. At the transition from F3 to F4, granular/fluorescent material becomes progressively enriched. The more granular/fluorescent patches dominantly crosscut less fluorescent ones. (Fig. 8D)

b) Medium burrows

The morphology of the medium burrows is best observed by narrow-spaced parallel sections, on opposite sides of a sawcut (Fig. 9A–B). Some burrows show abrupt changes in direction and/or diameter between closely spaced parallel sections (Fig. 9A–B, D). Burrows commonly show a dark halo around the cement fill, which in some cases can be followed into the sediment, defining a subcircular shape (Fig. 9E). Its lower part is filled with sediment and the upper part with cement. Within the burrows, the contact between sediment within burrows and cement above is planar, with an apparent dip of 0°–30° (Fig. 9E), indicating partially passive fill. Some burrows show laminated sediment fill recording multiphase passive infill. Macroscopically, the lumen of most medium burrows shows a concentric bipartite cement fill with a ca. 1 mm-thick tan, translucent, continuous outer rim around the lumen and a white to yellow/gold final axial fill (Fig. 10A–B). Both the dark halo and the outer rim are brightly fluorescent, in contrast to the low-fluorescent final fill (Figs. 9C, 10A–B). Burrow margins commonly show irregularities that might have resulted from burrow excavation, corrosion by fluids circulating in the burrows, in particular when they contain (hydrogen)sulfide, or local collapse of the burrow top. The distance between
medium burrows ranges from 2–10 cm with an estimated average of 5 cm. Burrows show subvertical, oblique and subhorizontal segments, without preferential orientation. The abrupt changes of orientation and diameter over short distances and the coexistence of vertical, oblique, and horizontal segments are diagnostic of burrows of the Thalassinoides – Spongeliomorpha suite (e.g., Cunningham et al., 2012; Knaust, 2017). These are produced by decapod crustaceans that may penetrate as deep as two meters or more below seafloor (e.g., Sarnthein, 1972; Pemberton and Buckley, 1976). Thalassinoides corresponds to smooth-walled burrows, being lined with mucus in soft sediment and unlined in sediment stiff enough to remain open and Spongeliomorpha to burrows scratched by body appendages indicating firm sediment (Wetzel and Uchman, 1998). Medium burrows occur preferentially in calcarenite and appear to get around pheno-intraclasts (Fig. 8A–B, E–G). However, in calcarenite patches, they do not follow specific patterns that could suggest spatial relationship between large and medium burrows (Fig. 8).

c) Small burrows

Small, cement-filled burrows are better visible under UV light. Some of these small burrows are directly connected to medium burrows while most appear isolated in vertical section (Fig. 9A, C, F), but out-of-plane contact with medium burrows appears likely. Their geometry, size, and fill match the characteristics of Trypanites representing borings in hard substrate. Small burrows may cut across pheno-intraclasts and/or peloid limestone, or locally follow the contact between the two.

4.3. Relative timing and tiering

Based on cross-cutting relationships, it is possible to distinguish successive phases of burrowing evidenced by intensely mixed, homogeneous sediment, overprinted by abundant shallow-produced large Thalassinoides, in turn cross-cut by deep, but scarcer and smaller Thalassinoides or Spongeliomorpha; occasional Trypanites borings entrenched into the cemented margins of the latter traces represent the last phase of bioturbation. In addition, the non-fluorescent wackestone occurring in patches has a grain content similar to the background sediment of Facies 1 or to the nodular facies F2 and F3. In contrast to the background sediment, it never shows lamination, nor stratification on the scale of limestone-marl alternations of the host sediment. Nonetheless, the processes that produced layering and lamination must have been active at the pseudobioherm site like in the surroundings. The uniform texture of the wackestone, therefore, reflects efficient mixing of the uppermost soupy/soft sediment by near-surface burrowing organisms producing so-called biodeformational structures that overprint pre-existing structures but do not constitute distinct traces (e.g., Schäfer, 1956; Wetzel, 1991). Consequently, it is possible to distinguish four phases of endobenthic activity recorded by

- tier 1 comprising biodeformational structures homogenizing soft sediment into wackestone (Phase 1),
- tier 2 constituted by burrows emplaced in soft to firm sediment mainly by large decapod crustaceans and filled by peloids (Phase 2),
- tier 3 characterized by burrows produced by smaller decapod crustaceans in sediment so stiff that the tunnels remained open (Phase 3), and occasionally,
- tier 4 solely evidenced by Trypanites borings originating at the walls of tier-3 burrows (Phase 4).

4.4. Microfacies and diagenesis

Tier 1: Micrite pheno-intraclasts
Marl/calcilutite homogenized by biodeformational bioturbation in tier 1 before it was cemented is only found as isolated remnants between burrows of deeper tiers. In thin section, these pheno-intraclasts consist of shaly-silty-bioclastic wackestone, with the same grain types as in background sediment (Fig. 4C), such as calcispheres, sponge spicules, and bivalve fragments (Fig. 10C).

**Tier 2: Large burrows and their infill**

Phase-2 burrows are filled with peloid grainstone. Peloids have a narrow size-range of fine-to-medium sand, typically 200–300 μm (Fig. 10D). Their morphology is ovoid, and their internal fabric is identical to that of pheno-intraclasts of F4 and nodules of F2, F3, consisting of silt/microfossil wackestone. Epifluorescence microscopy confirms the macroscopic observations that fluorescence mainly comes from microsparite cementing the grainstone (Fig. 10E).

**Tier 3: Medium burrows and their infill**

Cements composing the dark halo at the burrow boundary are commonly interlayered with sediment and thus, coeval with passive burrow infill (Fig. 9E). In thin section, the dark halo shows the same succession of cements irrespective of their position, between sediment laminae or at the sediment/cement boundary (Figs. 10A–C). It starts with a sediment-coating, 0.1–0.3 mm-thick layer of brown microsparite (BMSpar), commonly covered by 0.2–0.5 mm-thick clear microsparite (CMSpar, Figs. 11A, C, D, Fig. 12A). CMSpar, in turn, is locally covered by sets of radiaxial gray carbonate crystals (RAx; Fig. 11D). Sediment within burrows can show geopetal features above this succession, in some cases covered by a similar BMSpar-CMSpar couplet. The sediment-BMSpar-CMSpar succession has been observed up to 3 times (Fig. 12A). Individual microspar crystals are continuous across the boundary between brown and clear microspar (Figs. 12B–C). At high magnification, the brown color of BMSpar is due to bush-forming sets of brown filaments (Fig. 12D) whose diameters are close to the limit of optical resolution of ca. 1 μm. Individual “bushes” are about 100 μm in diameter and have a smooth, convex margin. Neither the abundant inclusions that characterize BMSpar, nor the calcite cementing the filaments are fluorescent, in contrast to CMSpar that shows both general mineral fluorescence and brightly fluorescent fluid inclusions (Fig. 12E–H). Fluorescent inclusions appear most abundant along the contact between BMSpar and CMSpar (Fig. 12F). Early diagenetic cements are, thus, arranged into repetitive sequences made of sediment/BMSpar/CMSpar, occasionally capped by radiaxial sets of calcite crystals; these will be referred to as “sediment-cement sequences”. Above the last sediment-cement sequence, the dominant cement is sparry calcite (Spar-1; Figs. 11C–D), with the occasional presence of botryoidal chalcedony (Figs. 11E–F).

Above the sediment-cement sequences and occasional Spar-1, there is a continuous 100–300 μm-thick rim of white cement, which in thin section consists of a high-relief light brown mineral (Figs. 10C, 11A) that shows the typical texture of flamboyant chalcedony in cross-polarized light (Sander and Black, 1988; Figs. 11B, F). This mineral, noted Chal-2, generally occurs in association with saddle dolomite (Dol-1) and is commonly covered by euhedral quartz crystals (Figs. 11D–F). Flamboyant chalcedony commonly cross-cuts single sparite crystals (Fig. 11C). It locally replaces older cryptocrystalline botryoidal chalcedony (Chal-1; Figs. 11 E–F). Below the continuous rim, discontinuous Chal-2/Dol-1 patches commonly replace the upper part of sediment-cement sequences thereby defining a complex, digitated surface of replacement. Silica replacement never occurs above a surface characterized by the occurrence of euhedral quartz crystals but can locally affect any of the underlying
cements, except those cementing grainstone. We refer to this surface as the “main silicification surface”, which can be traced as the envelope of euhedral quartz crystals (see Figs. 11 D, F). Cements above the main silicification surface are dominantly sparry calcite (Spar-2) and/or saddle dolomite (Dol-2), in variable proportions (Figs. 10C, 11A). Subordinate minerals include fracture-filling microcrystalline chalcedony and occasional sulfates (barite-celestite mixtures).

4.5. Stable isotopes

The stable carbon and oxygen isotope data of the studied carbonates exhibit a trend between the marl sample with δ¹³C of 1‰ and δ¹⁸O of -2.2‰ and a cluster centered at δ¹³C = -10‰ and δ¹⁸O = -1.5‰ (Fig. 13A; Appendix 1). Samples of the ¹³C-depleted cluster comprise microsparitic cements of the sediment-cement sequences (BMSpar and CMSpar) and Spar-1; all these cements pre-date the main silicification event (Fig. 13B–C). It also includes some peloids and some nodules. The majority of nodules and peloids and all micrite pheno-intraclast samples follow a mixing trend between marine sediment (F1 marl) and the ¹³C-depleted cluster. The δ¹⁸O values vary in a narrow range from -1.8‰ to -2.2‰ for early cements precipitated at seafloor temperature and from -1.1‰ to -2.1‰ for peloids and micrite pheno-intraclasts. Five of the 6 samples of saddle dolomite have similar values (-0.9‰ to -2.2‰) whereas the last one is the most depleted showing -7.5‰.

5. Discussion

The main proxies to identify seep carbonates in the rock record are the presence of specific methanotrophic fauna (e.g., Kiel, 2010) and/or the carbon isotope signature of the carbonate minerals, generally considered diagnostic if δ¹³C depletion exceeds -25‰. In Aurel, no sample meets the latter criterion. Similarly, the scarcity of CH₄-related macrofauna with one single specimen of a lucinid does not provide a compelling proof for the nature of the whole carbonate body. The most convincing evidence comes from combined analysis of bioturbation, early diagenesis in and around burrows and isotope geochemistry.

5.1. Alternatives to the bioturbation interpretation

The homogeneous character of micrite pheno-intraclasts and the tubular structures are interpreted as the result of bioturbation, affecting sediment of soupy consistency in the upper few cm of the seafloor and firm sediments below, respectively. Could these features be abiogenic and simply result from gas escape itself?

Homogenization by ascending fluid, known as fluidization, occurs in non-cohesive material when fluid ascent velocity exceeds the settling velocity of the particles (Allen, 1992). In the case of cohesive material interacting with gas, however, observations systematically indicate bubbling at discrete points rather than pervasive ebullition (Dupré et al., 2020). Both analogue and numerical modeling have shown that individual bubbles move as discrete lenticular hydrofractures within the cohesive material (Boudreau, 2012; Katsman et al., 2013). On the other hand, homogenization by bioturbation by meiofauna in the first few cm below seafloor is a well-documented phenomenon (Wetzel, 1983; Wetzel and Uchmann, 1998, and references therein). Moreover, the presence of discrete open conduits in the sub-seafloor ensures that any upcoming fluid (liquid or gas) will migrate along these conduits up to discrete venting points rather than homogenize the whole shallow sub-seafloor (Blouet et al., 2021c).
Could the tubular structures interpreted as burrows be related to other structures, for instance to abiogenic gas conduits? Generally speaking, due to its buoyancy gas tends to migrate vertically in homogeneous sediment. As observed for sand injectites and demonstrated by analogue modeling, deflection to horizontal can occur where changes in lithology/cohesion and/or permeability make it more energy-effective for ascending gas to follow a permeable, non-cohesive bed and lift the cohesive overburden than to fracture it. This phenomenon typically results in sill development, as observed for sand or magma injectites and modeled by Mourguès et al. (2012), and could explain limited subhorizontal segments (Boudreau, 2012). Abiogenic gas conduits have been interpreted by De Boever et al. (2009) in permeable sandy sediments, where free gas migrates vertically as long as the sediment is homogeneous and it is deflected laterally below less permeable stringers. An abiogenic origin has also been proposed for tubes in the Teepee Buttes Fm., whose dimensions and character are closer to what is observed in Aurel. The interpretation as abiogenic, 3-phase (gas-oil-water) flow is based on both the overall subvertical character, which is not the case in Aurel, and on the presence of silty laminae around the lumen of the tubes, interpreted to reflect deposition of clastic microparticles “adher[ing] to the surface of methane gas bubbles and entrained in the wake of the bubbles” (Krause et al., 2009). Neither criterion matches our observations, whereas the similarity with *Thalassinoides/Spongeliomorpha* is clear for most of the tubes. We therefore consider a biogenic origin as the most likely hypothesis in the case of Aurel.

5.2. Nature of the pseudobioherm

The co-occurrence of peloids and pervasive bioturbation strongly suggests a genetic link. In addition, the uniform size and regular ovoid morphology of the peloids, their internal grain content identical to that of pheno-intraclasts and adjacent nodules imply in-situ recycling of local sediment by deposit feeders. We thus interpret the peloids as fecal pellets, produced either by the organisms that dug Phase-2 burrows close to the seafloor or passively transferred into the burrows by bottom currents. The latter might also have formed the ripples in the background sediment.

The pseudobioherm appears as an oasis of thriving benthic life, essentially endobenthic, within an area marked by much lower abundance of benthos and scarce bioturbation. One key factor to support benthos is the availability of food/energy. Below the photic zone, benthos depends on the availability of organic matter produced in the photic zone and settling to the seafloor or brought in by currents, or, alternatively, by chemosynthesis. Very localized spots of sustained benthic life like the Aurel pseudobioherm (< 10 m in diameter) are found in particular at seep sites (e.g., Jensen et al., 1992). The Aurel pseudobioherm contains neither frame-building nor sediment-binding organisms. The vertical extent of the pseudobioherm of at least 15 m largely exceeds the maximum penetration depth of burrowing organisms.

In F4, the final fill of burrows generally consists of saddle dolomite, which is known to precipitate from saline fluids at temperatures in the range 80–160°C (Spötl and Pitman, 1998). The network of Phase-3 burrows, thus, remained open until this temperature value was reached supposedly at a burial depth of a few kilometers. During early stages of burial when the pseudobioherm still grew, the fine-grained background sediment had a strongly anisotropic permeability. Vertical permeability was low due to the dominance of laterally continuous marl strata whereas lateral permeability was high due to the presence of the intercalated calcilutite beds providing pathways for lateral fluid migration. In contrast, the absence of laminated limestone in the pseudobioherm and the pervasive biodeformational sediment mixing and hence, homogenization during the very first stages of burial.
imply an isotropic low permeability of the pseudobioherm deposits in the first 5–10 cm below seafloor. In contrast, the open burrow network of F4 provided localized natural conduits for fluid flow during compaction, making the whole pseudobioherm an effective vertical drain at large scale. Therefore, we interpret the vertical development of the Aurel pseudobioherm to have resulted from dominantly vertical fluid seepage facilitated by burrows and making it a fluid seep carbonate body.

5.3. Nature of the fluid

Stable carbon isotope analysis is the most commonly used method to elucidate the origin of fluids from which seep carbonates are derived (Campbell, 2006); a “classical seep carbonate” signature is in the range of -60 to -30‰ for δ¹³C and -2 to +8‰ for δ¹⁸O (Campbell 2006: fig. 9A). There appears to be a consensus that any authigenic carbonate exhibiting δ¹³C values lower than -30‰ can be considered diagnostic of methane oxidation, whereas less negative values may result from mixing of bicarbonate derived from other sources, in particular seawater (Campbell, 2006). In the Aurel pseudobioherm, the most δ¹³C-depleted samples exhibiting about -10‰ are those predating the emplacement of the main silicification surface and being large enough to be sampled without mixing. Samples comprising both sedimentary carbonate and diagenetic cement have δ¹³C values along the trend between marine sediments and the most depleted cements (Fig. 13A).

In this study, the most ¹³C-depleted cements are significantly less depleted than the commonly accepted diagnostic δ¹³C threshold value of -30‰. This threshold results from the successive changes in ¹³C depletion during the maturation of organic matter to hydrocarbons and eventual precipitation of carbonate when anaerobic oxidation of these hydrocarbons took place. The δ¹³C of sedimentary organic matter ranges from -10 to -35‰, with an average of -25‰ in marine environment. During burial, microbial methanogenesis leads to further depletion. Microbially generated methane shows a δ¹³C depletion from -120‰ to -50‰ (Botz et al. 1996, Whiticar, 1999), whereas thermogenic maturation yields methane with δ¹³C values from -50 to -35‰ (Fuex, 1977).

Comparing carbon isotopy of recent seep carbonates with that of gas bubbling from the corresponding seep sites evidenced enrichment in δ¹³C by ca. 20‰ on average upon carbonate precipitation, interpreted to result from mixing with dissolved inorganic carbon (Peckmann and Thiel, 2004). Combining the successive phases of depletion/enrichment, seep carbonates should have typical values of -100‰ to -35‰ when derived from microbial methane and -35‰ to -15‰ when sourced from thermogenic gas. A recent review of methane carbon isotopy based on more than 20,000 samples indicates that the δ¹³C of thermogenic methane ranges from -65‰ to -25‰ and depends very much on the maturity of CH₄ (Milkov and Etiope, 2018). In detail, oil-associated methane matches the values proposed by Fuex (1977), whereas early mature values are in the range of -65‰ to -50‰ and late mature ones cluster around -30‰. Assuming that the average mixing effect with seawater is in the range of 20‰ as observed by Peckmann and Thiel (2004), we interpret the δ¹³C of -10‰ in the early cements of the Aurel pseudobioherm to indicate that anaerobic oxidation of late thermogenic methane fueled seep carbonate precipitation. The values measured at Aurel show a dominant trend in the negative δ¹³C zone, in sharp contrast with the wider range of values reported from the neighboring Beauvoisin pseudobioherms, from +17‰ to -24‰ (Gay et al., 2020). At the same time, the influence of AOM has been evidenced by the presence of lipid biomarkers in Beauvoisin (Peckmann et al., 1999), and the most ¹³C-depleted samples fall much closer to the generally admitted diagnostic threshold than in Aurel. We interpret the difference to result mostly to a difference in the
source of fluids, microbial methane from the Bathonian-Oxfordian in Beauvoisin (Gay et al., 2020) and late thermogenic methane in Aurel, hypothetically from a deeply buried Carboniferous source rock.

Oxygen isotope values exhibit a poorly organized pattern, especially as regards saddle dolomite. The latter is known to have precipitated at 80°C at least (Spötl and Pitman, 1998), typically resulting in $\delta^{18}O$ values below -6‰. Five of the 6 values obtained here for saddle dolomite are inconsistent with precipitation from hot fluids. The apparently erratic character of oxygen isotopes in saddle dolomite could be due to the fact that oxygen is much more affected than carbon by isotope exchange during diagenesis of carbonates (Peckmann et al., 2003; Zwicker et al., 2015). We acknowledge the problem but consider that the issue goes beyond the scope of this study.

Nonetheless the Aurel pseudobioherm is interpreted as a column of seep carbonate precipitated at the top of a stationary, 5–10 m-diameter fluid chimney supplying late mature thermogenic methane.

5.4. Facies interpretation

5.4.1. Facies F1, F2, F3

Background sediment records hemipelagic settling of fine-grained siliciclastic and carbonate particles forming marl and episodic density currents that transported material from the slope or the shelf downward constituting calcareous, often calcilutite beds. Hydrocarbon-charged fluids seeping up through the limestone column does not appear to have affected sedimentation or early diagenesis more than a few meters away from the venting site. Only the 'normal', basin-wide burial compaction affected the diagenesis of F1. For F2 and F3, the isotope signature of nodules show an input of AOM-generated bicarbonate to levels above the precipitation threshold of calcite, the more as the nodules become more abundant and coalesce closer to the axis of the pseudobioherm (Fig. 14). Calcite precipitation occurred in the SMTZ that has been shallower above the pseudobioherm than in the surrounding 'host' area due to the increased methane supply (Fig. 14, cf. Paull and Ussler, 2008). In F3, the saddle dolomite crystals in the matrix between coalescing nodules indicate percolation of late diagenetic fluids through the limestone-dominated column that still acted as a local fluid conduit even if burial depth was sufficient to reach a temperature of at least 80°C.

5.4.2. Facies F4

Carbon isotope data of the Phase-3 burrow infill indicates that the main silification surface separates AOM-influenced early diagenetic cement from late diagenetic saddle dolomite as prominent mineral. Anaerobic oxidation of methane can only occur if sulfate is available, that is present at a seep site only within the first few meters below seafloor at the deepest. The high temperatures required for saddle dolomite precipitation, however, imply that late diagenesis took place at depths largely exceeding 1 km, more likely 2 km assuming geothermal gradients of 30°–40°C km$^{-1}$.

In F4, the early, AOM-dominated diagenesis prevailed whereas the late diagenetic phase only affected Phase-3 burrows. The F4 fabric records the interplay between sedimentation, bioturbation and early diagenesis. Sedimentation at the seep site must have been identical to that recorded by background facies F1, since hemipelagic settling and current deposition acted on a larger scale than the 10-m diameter of the pseudobioherm. Homogenization by bioturbation was the first process that modified the surface sediment, followed by very early cementation that started stiffening the deposits. Bioturbation phases 2 and 3 occurred after enough seep carbonate
had been precipitated to maintain burrows open and preservable (e.g., Wetzel and Uchman, 1998). The open burrows in turn guided upward methane migration and promoted AOM and further gradual cementation around the burrows (Wetzel, 2013). Leaving aside the effects of silicification, the cement infill of Phase-3 burrows and their host sediment is quite simple: AOM-related carbonate precipitation in the SMTZ is expressed by diagenetic micrite within the pores of homogenized 'depositional' micrite, by microspar crystals in the pores of pellet grainstone and in the first 0.1–0.5 mm off the burrow walls (BMSpar-CMSpar) and by spar-size crystals (Spar-1) in the remaining free space of the lumen. The radiaxial sets of gray carbonate crystals that occasionally terminate the sequence of cements (RAx) may represent recrystallized equivalents to the aragonite fans described by Blouet et al. (2017) in the Panoche Hills (CA, USA) or by Peckmann et al. (1999) from Beauvoisin. Calcite crystals continuing across the BMSpar-CMSpar contacts indicate that the brown color of the former simply reflects the inclusion of the filamental bushes into the growing microspar. These bushes are reminiscent of the dumbbell-shaped crystal aggregates reported by Peckmann et al. (1999) from the Miocene seep carbonates of Marmorito (Italy), and are most likely filamentous, bush-shaped microbial colonies having grown over free surfaces of micrite pheno-intraclasts or pellets at burrow walls. The presence or absence of geopetal sediment in the burrows simply reflects the capacity of a burrow segment to trap pellets avalanching from above. Pellets could fall down due to biological activity above or bottom currents inducing circulation within burrows. The variability of $^{13}$C depletion of micrite (nodules and pheno-intraclasts) reflects variations in the amount of marine carbonate mixed with the AOM-derived cement. In multi-phase samples such as pellet grainstone containing AOM-derived cement, the variability may result from varying amounts of pellets derived from micrite with a variable isotope signature and cement (Figs. 8D, 10C).

At seep sites, the SMTZ crops out at the seafloor (Paull and Ussler, 2008). In the Aurel pseudobioherm, the cm-scale geometry of the SMTZ was most likely complex, with pervasive downward digitations following the open burrow network that provided connection to the sulfate-bearing open water. In addition to this complex geometry, tube ventilation by seawater is known to be induced by bottom currents such as tidal currents (“passive ventilation”: Wetzel et al., 2014; Gingras and Zonneveld, 2015; Rodríguez-Tovar et al., 2019) and by the burrowing organism itself (“active ventilation”). Active ventilation has been shown to be used by decapod crustaceans to promote microbial growth on burrow walls (Savrda, 2007). The microbial colonies of BMSpar may result from one or the other type of ventilation. The key information provided by late diagenesis about the vertical growth of the pseudobioherm, is that the tube network constituted by burrow lumina remained open long after the pseudobioherm was buried, at least as long as the pseudobioherm was growing.

Silica diagenesis occurred in two phases: in the first one, chalcedony botryoids precipitated locally above a first succession of AOM-related cements was in turn covered by AOM-related sparite Spar-1; the second phase post-dates all AOM-related carbonate precipitation and is coeval with the precipitation of saddle dolomite. Silica is ubiquitous in the host sediment as sponge spicules and so it was readily available within and all around the pseudobioherm. Precipitation of AOM-related carbonates followed by silicification and in turn precipitation of non-AOM related carbonates has been reported from Paleozoic–Cenozoic fossil seep carbonates hosted in siliceous microfossil-bearing sediments by Kuechler et al. (2012) and Smrzka et al. (2015). These authors interpret the silicification event to reflect silica dissolution by alkalinity/pH increase during AOM, favoring dissolution of the silica, followed by reprecipitation when AOM ceased (Kuechler et al., 2012) or the zone of main methane flux shifted laterally (Smrzka et al., 2015). The first phase of silica precipitation in Aurel is coeval with AOM, and thus
does not fully match the observations of Kuechler et al. (2012) or Smrzka et al. (2015). However, early silica could have precipitated during temporary, local interruption of AOM by redirection of methane(-bearing fluids) to another of the numerous conduits in the pseudobioherm that was followed by another episode of AOM when the flow was re-established. The second, more pervasive phase of silicification was coeval with precipitation of saddle dolomite and largely post-dated the demise of AOM. We hypothesize that it corresponds to a specific step of silica diagenesis affecting the sponge-rich interval of the pseudobioherm, perhaps upon entering the Opal C-T-Quartz transformation window around 55–85°C (Keller and Isaacs, 1985), which overlaps with the lower range of precipitation temperature for saddle dolomite.

5.5. Parameters controlling the stacking pattern

The Aurel seep-carbonate body comprises vertically stacked carbonate lenses interfingering with the host marls. The stacking pattern and the uniform character of F4 over the whole thickness of the pseudobioherm indicate overall upward growth keeping pace with deposition. Lateral compensation between successive lenses, due to self-sealing and lateral shift, remains well within the diameter of the carbonate body, except across marker beds A and B where the axis of the pseudobioherm shifted by several meters and stabilized in its new position (Fig. 3). Downward growth, if any, would be restricted to individual lenses a few tens of cm thick. Which parameters controlled the vertical growth of the pseudobioherm between the marker beds? The role of Thalassinoides in focusing migration of methane(-charged fluids) has been identified by Wetzel (2013) and later by Wiese et al. (2015), Zwicker et al. (2015) and Blouet et al. (2017). Evidently, in spite of likely overprinting by shallower-tier burrowers, individual Phase-3 burrows remained open until burial reached at least 1 km and provided obvious fluid migration pathways as deep as the burrowing organisms penetrated, likely in the range of 1–3 m as known for present-day decapod crustaceans (see above). Within tier 1, mainly small-sized organisms (meiofauna < 300 μm) that represent the most efficient 'homogenizers' (e.g. Wetzel, 1983) probably avoided the shafts of Thalassinoides not to fall into them. Therefore, at least some shafts remained open after burrow abandonment. Burrowers of tier 2, however, could have intersected the upper part of abandoned tier-3 burrows and shed sediment and pellets down the intersected shafts. The estimated average spacing of ~ 5 cm between tier-3 burrows implies a tunnel density of about 400 per m², in the mid-to-high density range defined by D’Andrea and DeWitt (2009). As the burrows have a complex branching geometry made of segments with highly variable azimuth and inclination, there is room for many intersections between active and abandoned Phase-3 burrows. Thalassinoides-/Spongeliomorpha-type burrows generally show variable orientation on a cm-scale (e.g., Griffis and Suchanek, 1991; Dworschak and de Rodrigues, 1997; Stamhuis et al., 1997; Ziebis et al., 1996), as observed in the Aurel samples (Fig. 15A). The gently inclined segments provide dead ends or traps for sediment avalanching in or entering the burrows; as soon as sediment derived from above reached a segment dipping less than the angle of repose of granular material, passive fill is hindered and the tunnel network underneath remains open (Fig. 15B). The geometric arrangement of the tier-3 burrows with subhorizontal segments thus represents an important factor ensuring that these burrows provided an open tube network over the whole thickness of the pseudobioherm as long as there were organisms active in providing the final opening to the seafloor.

Vertical growth is the consequence of such fluid circulation 'highways'. Fluids expelled during compaction and buoyant hydrocarbons, once they reached the base of the pseudobioherm, were funneled within the open-burrow network up to domains close to the seafloor where seep carbonate precipitation occurred in the
SMTZ. Free gas trapping by any local permeability restriction inside a tube would initiate buildup of a continuous gas column below. Its buoyancy could overcome the mechanical resistance of the plug as soon as the height of the column reached about the thickness of the plug (Cathles et al., 2010). The resulting sudden fluid release may have episodically flushed the tube system, contributing to keeping it open. At any time during active seep carbonate growth, the tier-3 burrows provided connection with the seawater on the one hand and brought up methane on the other, ensuring both seep carbonate precipitation in the shallow sub-seafloor and multiple bypass pathways preventing self-sealing. Increasing cementation around the open tubes prevented their collapse. Decapod crustaceans are known to dig deep into the sediment and periodically ventilate their burrow systems with seawater that contains sulfate and to induce cross-stratal fluid flow (e.g., Forster and Graf, 1992; Ziebis et al., 1996). Once initiated, vertical growth became a self-sustaining process.

What caused the offset of the seep carbonate column upon crossing marker beds A and B? Event deposition of relatively a few tens of cm thick marker beds should not have significantly affected burrowers within tiers 2 and 3 as they have the capability to maintain the connection to the seafloor (e.g., Bromley, 1996) and evidently did so, and vertical communication up to the seafloor as previously described must have been reestablished rapidly. Because massive calcilutite beds in the pseudobioherm became homogenized by bioturbation in the very shallow, 0–10 cm sediment depth (Wetzel, 1981), it is highly likely that bioturbational mixing extinguished the continuity of thinner limestone beds by displacing grains into the adjacent marly mud, whereas sufficiently thick beds would just show blurred contact at the top limestone/marl contact but preserve a permeable lower part. Any restriction to fluid flow above the permeable bed likely deflects at least part of incoming methane-bearing fluid along the permeable bed, where migration would follow the local gradient at the top of the permeable bed (regional slope combined with local irregularities at the base of the bioturbated, impermeable top-of-bed interval). Upslope migration of hydrocarbon-charged fluids, in turn, shifts the zone of maximum hydrocarbon concentration laterally upslope and also the habitat of burrowers feeding on chemosynthetic microbial communities along their burrow walls and on the seafloor.

In summary, the main factor governing seep carbonate stacking in Aurel appears to be the contrast in m³-scale permeability between the pseudobioherm and its host sediment (Fig. 15C). Sedimentation of the background facies defines a strongly anisotropic permeability tensor on m³-scale, with low vertical permeability due to the fine-grained marls whereas high lateral permeability occurs before lithification due to the presence of calcilutite limestone beds. At the same scale in the seep carbonate column, the Thalassinoides/Spongeliomorpha tube network results in high vertical permeability (e.g., Cunningham et al., 2012; Gingras et al., 2012) whereas lateral permeability is generally lowered by bioturbational blurring of thin granular limestone beds into the marls. Only granular beds significantly thicker than the mixing depth of near-surface endobenthic organisms remain to a large proportion in their original state (e.g., Wheatcroft, 1990; Wetzel, 2009). They can retain sufficient permeability to act as drains, leading to lateral deflection of incoming fluids and lateral shift of the stacking pattern.

6. Conclusion

The development of the Aurel pseudobioherm, a 15-m-thick, 8-m-diameter columnar seep-carbonate body encased in a >500 m-thick marl succession, was modulated by factors that promoted its vertical growth and limited lateral shift induced by self-sealing. The following two points help explain one key factor that appears to be intense bioturbation by decapod crustaceans:
a) The specific morphology of the burrows produced in a deep tier includes low-inclination to horizontal segments that could hinder passive infill by sediment falling from the seafloor or shallower bioturbation tiers, preserving a network of open tubes below.

b) The areal density of burrows of ca. 400 burrows m\(^2\), was sufficient to ensure cross-cutting between successive generations of burrows, providing a connected network extending deep below the base of the SMTZ and thereby fostering unrestricted vertical fluid percolation through the whole carbonate body due to their high vertical permeability.

The host sediment is dominated by fine-grained marls resulting in very low vertical permeability. In contrast, intervening cm-thick laterally continuous calcilutite beds make up to a few percent of the formation and create a relatively high lateral permeability. At the location of the pseudobioherm, these thin beds were mixed with mud through biodeformational bioturbation in the upper 5–10 cm below seafloor, blocking lateral fluid circulation inside the pseudobioherm and into the host sediment.

Conversely, the few >10-cm-thick calcilutite beds were not fully homogenized, leading to (possibly upslope) lateral pseudobioherm migration by a few meters along the thicker beds.

The critical factors thus appear to be the adequate type and abundance of burrows; they foster compaction fluid flow into the pseudobioherm with positive feedback on the development of endobenthic life. Decapod crustacean-promoted focusing of fluid flow into a seepage area could be the antidote to self-sealing and promote sustained vertical growth of seep carbonate bodies. This type of bioturbation, known to be common at hydrocarbon seep sites, may thus be a small-scale but critical factor fixing a seepage site at the same point for long spans of time, as has commonly been observed on seismic sections worldwide.

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References


Figures

Figure 1: Geological setting of the studied outcrop. A) Cross-section from the margin to the center of the Dauphinois Basin traversing the Aurel borehole (after Wannesson and Bessereau, 1999, modified). B) Facies map of the Dauphinois Basin for the Callovian (Debrand-Passard, 1984); for the basin facies represented by the Terres Noires Formation (middle Bathonian up to lower Oxfordian) in the central part of the basin isopachs are shown (after Beaudrimont and Dubois, 1977). Outcrops of the Terres Noires Formation (after Artru, 1972) and the location of the major pseudobioherms marked by red stars (after Rolin, 1987). C) Lithologic log of the Aurel area (Flandrin, 1974). The Aurel borehole penetrated the carbonate-dominated Dogger and the marly upper Lias. The pseudobioherms in the Aurel area are located in middle Callovian sediments; the pseudobioherms in the Dauphinois Basin occur in upper Bathonian to middle Oxfordian deposits. D) Geological map of the Aurel area (after Flandrin, 1974, modified). Several pseudobioherms are poorly exposed near the Vaunière ruin; the studied pseudobioherm is located near La Touche farm.
Figure 2: Lithologic log of the middle Callovian deposits in the pseudobioherm-bearing interval, logged about 20 m to the west of the upper part of the pseudobioherm, not affected by faulting; note marker beds A and B, easily recognizable in both host sediment and pseudobioherm.
Figure 3: A) and B), Image and interpretation of the southern flank of the Aurel pseudobioherm; dashed lines represent assumed axis of the pseudobioherm. C) and D), Image and interpretation of the eastern flank of the pseudobioherm. LST = limestone.
Figure 4: Background facies F1. A) Typical alternation between marl and limestone showing platy bedding. B) Surface of a slab with radiating burrow (*Gyrophyllites multiradiatus*); inset shows the bivalve *Bositra*, the only common fossil in F1. C) Marl with numerous filamentous structures interpreted as disaggregated *Bositra* shells. D) Limestone showing the same microfossil content as the marl but in microsparitic matrix. E) Limestone consisting of alternating packstone and wackestone laminae. F) Marl with numerous recrystallized microfossils in a micritic matrix; circular objects are interpreted as calcspheres or radiolarians; inset showing a tubular structure interpreted as a sponge spicule.
Figure 5: Facies F2, nodular marl. A) Outcrop view. B) Individual nodules and nodule clusters. C) and D) Cross-section of a nodule under natural (C) and UV light (D); darker (C) non-fluorescent (D) vertical parting in the middle is suggestive of previous boundary between two, now clustered nodules.
Figure 6: Facies F3, nodular limestone. A) and B) Vertical section under natural (A) and UV light (B); arrows point upward. C) Photomicrograph showing contact between a nodule and matrix (marked yellow dashed line). Grain packing in matrix is denser than in nodule, suggesting cementation of the nodule prior to compaction.
Figure 7: Facies F4, massive limestone, and associated fauna. A) Weathered surface showing three bivalves shown in (B) to (D) in detail, apparently belonging to different species. D) Bivalve with geopetal filling. E) and F) Two valves of the only lucinid found (*Beauvoisina carinata*, Kiel et al., 2010†). G) Section of a small gastropod. H) Bedding plane view of *Thalassinoides*; smaller tape measure graduations in mm.
Figure 8: Fabric of intensely bioturbated sediment. A) Natural light and B) UV fluorescence light, Facies F4. White arrows in B) indicate micrite pheno-intraclasts. Note the variability of pheno-intraclasts in roundness and size. C) Natural light and D) UV fluorescence light, transition from F3 to F4. In both samples, medium burrows (MB) appear independent from variations in fabric around them; black rectangles drawn for thin section selection. E) Natural light, F) UV fluorescence light and G) Line drawing, facies F4, unoriented vertical section. White arrows in G) show points where burrow margin tangentially passes pheno-intraclasts.
Figure 9: Burrow architecture in F4. A) and B), Opposing sides of a sawcut, ca. 3 mm apart; Note 3 types of burrows: Large burrows (one marked as Ba) surrounded by white dashed line; medium burrows, (partly) filled with white-yellowish cement (labelled B1–B6); small burrows (t1–t3) encircled in red. C) UV-light view of the section shown in (A). Yellow lines follow main dark halos, as marked in (A). D) Line drawing of (C) showing with marked burrows and halos; yellow dotted line marking outlines of B1–B6 on opposite section B), highlighting variations in size/orientation across the 3-mm gap. E) Close-up of B6 marked in (A) showing burrow surrounded by dark halo (bold arrows) and half-filled with geopetal sediment; thin white arrows show halos in the geopetal filling. F) Cross-cutting relationships between burrows (UV fluorescence). White circles point at small burrows connected to a medium burrow. White rectangle highlights longitudinal cross-sections of small burrows. Black rectangles drawn for thin section selection.
Figure 10: see caption next page
Figure 10: A) and B) Close-up of Figs. 8 A, B showing the details of burrow infill. Orange arrows in (A) mark dark halo surrounding the burrow on the left side and expanding on the right side into peloid grainstone (PG); yellow arrows mark thin, continuous, milky white cement rim following the dark halo on the left side of the image but the sediment-cement contact on the right side. B) UV fluorescence view of the same section; the blurry appearance results from diffusion of light emitted from fluorescent parts into non-fluorescent ones. Int = micrite pheno-intraclasts, EC = early cements, LC = late cements; white rectangle refers to detail shown in C. Pheno-intraclasts, like nodules in F2 and F3 with low level of fluorescence, in contrast to brightly fluorescent peloid grainstone (less fluorescent spots therein are individual peloids). The brightly fluorescent rim coincides with the cement ring limited on either side by an outer dark rim and an inner white rim. C) Microphotograph of the same sample; dark halo (orange arrows) corresponds to superposition of brown and clear microsparite layers; brown microspar (BMSpar) in direct contact with burrow wall. The white rim (yellow arrows) consists of beige chalcedony (for details, see Fig. 11). White arrows point to shell fragments, PW = peloid wackestone, Dol 2 = late saddle dolomite. D) and E) Photomicrographs of peloid grainstone in F4. D) Internal texture of the peloids in plane polarized light. E) as (D) in UV epifluorescent light.
Figure 11: Cements within Phase-3 burrows and host sediment. A) and B) General view of burrow margin in plane light (A) and cross-polarized light (B). Note continuous rim of beige chalcedony (Chal-2) associated with saddle dolomite (Dol-1). C) Detail of relationship between Chal-2 and carbonate cements in cross-polarized light; white rectangle: Chal-2 cutting across a single sparite crystal. D) Two stacked sediment-cement sequences in plane polarized light showing alignment of euhedral quartz Q roughly parallel to burrow margin wall, as well as radiaxial gray calcite crystals (RAx) locally capping Sed-BMSpar-CM Spar succession; blue zigzag line marks upper limit of euhedral quartz. E) and F), main phases of silicification in plane light (E) and cross-polarized light (F); Chal-1 = botryoidal chalcedony precipitated freely at the tube wall, Q = euhedral quartz also overgrown on a free surface, Chal-2 and the associated saddle dolomite Dol-1 replaced both pre-existing carbonates and Chal-1. Note the extinction pattern of Chal-2 characterizing “flamboyant chalcedony”. The blue zigzag line marks the main silicification surface. For details see text.
Figure 12: Microphotographs showing details of sediment-cement sequences. A) Repeated sequence of sediment (sed), brown microsparite (BMSpar), and clear microsparite (CMSpar), locally overprinted by chalcedony (Chal-2). B) Detail of BMSpar-CMSpar contact (marked by black dashed line); BMSpar exhibiting filamentous “bushes”. C) as (B) in cross-polarized light, showing individual microspar crystals straddling across BMSpar/CMSpar boundaries (marked by black arrows); white rectangle marks area shown in detail in (D). D) Close-up of filamentous bush; note elongate brown inclusions within clear crystal. E) and F) BMSpar/CMSpar contact in plane light (E) and UV epifluorescent light (F); fluorescent inclusions near the margin of BMSpar following oblique contact between BMSpar and CMSpar into the thin section. G) and H) Typical CMSpar in plane light (G) and UV epifluorescent light (H), fluorescence originating from both fluid inclusions (bright white) and microspar (dull yellow).
Figure 13: Stable C and O isotopes and microfacies. A) Sedimentary and diagenetic carbonate phases encountered in the Aurel pseudobioherm in $\delta^{13}C$-$\delta^{18}O$ plot; position of samples of facies F1 to F3 is shown in Fig. 14. MCSW = Middle Callovian seawater composition; LMCSW = subset for the lower part of the Middle Callovian (Pellenard et al., 2014). B) Fabric and microfacies of F4 recording successive phases of bioturbation. C) Schematic geometric relationships between sediment and various cement generations and their stable isotope values within facies F4; black boxes mark minimum size of samples drilled for isotope analysis, some likely mixing different phases like peloid grainstone and BMSpar/CMSpar. GST = grainstone, MSS = main silicification surface, for details, see text.
Figure 14: Change in depth of the sulfate methane transition zone (SMTZ) and methane-derived authigenic carbonates (brown) from host sediment towards the pseudobioherm; vertical dashed line marking the axis of the pseudobioherm, symbols referring to the isotope data shown in Fig. 13A. Not to scale.
Figure 15: Fluid flow in and around the pseudobioherm (PBH). A) Typical architecture of a recent decapod crustacean burrow (*Callianassa truncata*, after Ziebis et al., 1996, modified); gray surfaces indicating hypothetical limits between bioturbation tiers 1 to 3 in Aurel (tier 4 below shown interval). The blue frame marks the shallowest chamber acting as trap for sediment shed from above and minimizing further downward sediment transfer. B) Close-up of trapping chamber; once lateral spill-out from the chamber reaches angle of repose (ca. 30°, Allen, 1992), additional sediment from above piles up in the shaft and deeper parts of the burrow remain open. C) Sketch of fluid circulation in and around the Aurel pseudobioherm (PBH) during its formation. Lower part representing hypothetical deep structure focusing late thermogenic methane into the pseudobioherm; red dashed arrows at depth indicate hypothetical gas migration pathways from a source rock into the structure and further up to the base of the pseudobioherm. Upper part, circulation pattern of gas and fluids within the pseudobioherm; orange zigzag lines represent crustacean burrows providing a connected tube network down to the bottom of the pseudobioherm. White ellipses represent permeability tensors in the pseudobioherm and host sediment, with the principal axes of minimum and maximum permeability shown by black arrows; red arrow within the pseudobioherm marks vertical flow pattern of the gas (-charged fluids), likely with episodic escape of gas at the seafloor (red bubbles in water column).
Highlights

- A 15 m-thick and 10 m-wide columnar carbonate pseudobioherm shows typical facies architecture.
- The pseudobioherm is characterized by a dense network of subvertical and horizontal crustacean burrows.
- About 200–500 burrows m² remained long time open after burial of the column.
- Early cements within burrows are depleted in $^{13}$C, indicating AOM shallow in sediment.
- The open burrow network promoted vertical growth of the methane seep carbonate pseudobioherm.
## Appendix 1

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