

Oncoid growth and distribution controlled by sea-level fluctuations and climate (Late Oxfordian, Swiss Jura Mountains)

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Abstract Abundant lagoonal oncoids occur in the Late Oxfordian Hauptmumienbank Member of the Swiss Jura Mountains. Four oncoid types are observed in the studied sections and classified according to the oncoid surface morphology, the structure and composition of the cortex, and the texture and fauna of the encasing sediment. Micrite-dominated oncoids (types 1 and 2) have a smooth surface. Type 1 has a rather homogeneous cortex and occurs in moderate-energy environments. Type 2 presents continuous or discontinuous micritic laminae. It is associated with a low-diversity fauna and occurs in high-energy facies. *Bacinnella* and *Lithocodium* oncoids (types 3 and 4) display a lobate surface. They are dominated by microencrusters (*Bacinnella irregularis* and *Lithocodium aggregatum*) and are found in low-energy facies. The stratigraphic and spatial distribution of these oncoid types shows a correlation with the sequence-stratigraphic evolution of the studied interval, and thus with relative sea-level fluctuations. It can be shown that these sea-level fluctuations were controlled by orbital cycles with 100- and 20-kyr periodicities. At the scale of 100- and 20-kyr sequences, types 1 and 2 oncoids are preferentially found around sequence boundaries and in transgressive deposits, while types 3 and 4 oncoids are preferentially found around maximum floodings and in highstand deposits. This implies that changes of water energy and water depth were direct controlling factors.

Discrepancies in oncoid distribution point to additional controlling factors. Platform morphology defines the distribution and type of the lagoon where the oncoids flourished. A low accumulation rate is required for oncoid growth. Additionally, humidity changes in the hinterland act on the terrigenous influx, which modifies water transparency and trophic level and thus plays a role in the biotic composition and diversity in the oncoid cortex.

Keywords Oncoids · Carbonate platform · Sequence stratigraphy · Milankovitch cycles · Sea level · Palaeoenvironments · Palaeoecology

Introduction

The term “oncoid” (from the Greek *onkos*, “lump”) was coined by Arnold Heim in 1916. He thus described coated grains in Jurassic and Early Cretaceous limestones in the Swiss Helvetic nappes that differed from “oids” by non-concentric layers, not only round but also lobate shapes, dense texture with or without inclusions, and in some cases indistinct boundaries. For oncoid genesis, Heim (1916) invoked micro-organisms (bacteria) and speculated if carbonate was precipitated within the bodies of these organisms, if they only stimulated precipitation by their metabolic activity, or if they agglutinated carbonate particles. In the following decades, several authors described and classified oncoids and interpreted their depositional environments (Peryt 1981; Flügel 2004). Peryt (1983) defined them as “a group of algal (red algae excepted), cyanobacterially and bacterially coated grains, which are initiated in marine and freshwater phreatic environments”.

In our study, we focus on marine oncoids that formed during the Late Oxfordian on a part of the shallow

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carbonate-dominated platform on the northern margin of the Tethys ocean, which today is represented by the Swiss Jura Mountains. Late Jurassic marine oncoids have already been described and discussed by Dahanayake (1977, 1978, 1983) from the French Jura and by Gygi (1992) and Hug (2003) from the Swiss Jura. In this paper, the different oncoid types occurring abundantly in the Hauptmumienbank (“main mummy bed”) Member in the Swiss Jura are examined in order to better constrain the link between oncoid types and palaeoenvironmental and palaeoecological conditions. We present the results of microfacies analysis, describe the stratigraphic and lateral distribution of the different oncoid types, and discuss the main environmental factors controlling oncoid growth and distribution. We thus test if oncoids can be used as palaeoenvironmental proxy.

Stratigraphic, palaeogeographic, and palaeoclimatic framework

The stratigraphy of the Oxfordian in the Swiss Jura is well established by Gygi and Persoz (1986) and Gygi (1995, 2000a, 2000b). The oncoid-rich Hauptmumienbank Member belongs to the Vellerat Formation (Fig. 1). This lithostratigraphic unit received its name from Ziegler (1956) who was inspired by Steinmann’s (1880) comparison of microbially wrapped grains with Egyptian mummies. Laterally, it passes into the Steinebach Member characterized by ooids. Gygi (2000a, 2000b) attributes the Hauptmumienbank Member to the Late Hypselum ammonite subzone, belonging to the Bimammatum zone. In sequence-stratigraphic terms, the Hauptmumienbank and its lateral equivalents correspond to the transgressive deposits of the Ox 6+ sequence (Fig. 1). This sequence has been identified by Hug (2003) as an additional sequence and overlies the Ox 6 sequence of Hardenbol et al. (1998). Cyclostratigraphic analysis suggests that this sequence corresponds to a 400-kyr orbital eccentricity cycle (Hug 2003; Strasser et al. 2000). The Ox 6+ sequence boundary coincides with a maximum regression on the second-order scale (Hardenbol et al. 1998).

In the Late Oxfordian, the area occupied by today’s Swiss Jura was part of a complexly structured but generally shallow epicontinental domain (Fig. 2). The maximum of the second-order regression in the early part of the Bimammatum zone caused large areas to emerge. During the following transgression, abundant carbonate production compensated for the general eustatic sea-level rise and forced the Swiss Jura platform to prograde towards the southeast (Gygi and Persoz 1986). Depressions created by differential subsidence (induced by tectonic movements in the Permo–Carboniferous basement; Allenbach 2001; Hug 2003) acted as traps for siliciclastics (Pittet 1996;

Wetzel et al. 2003). The different facies occurring on this platform have first been documented by Rollier (1898) and were later described in great detail by numerous authors (e.g., P.A. Ziegler 1956, M.A. Ziegler 1962; Pümpin 1965; Bolliger and Burri 1967, 1970; Gygi 1969, 1982, 1986, 1992; Gygi and Persoz 1986). More recently, Pittet (1996) and Hug (2003) investigated, in more detail, facies and depositional sequences of the Vellerat Formation and Dupraz (1999) studied the palaeoecology of Middle–Late Oxfordian coral reefs. The encountered facies and sedimentary structures indicate depositional environments ranging from tidal flats to shallow lagoons, coral patch-reefs, and oolitic bars. Coral bioherms concentrated mainly on the platform edge and in open-marine environments, while oncoid-rich lagoons and ooid bars occupied the internal platform.

The study area was situated at a palaeolatitude of 26°–27°N (Dercourt et al. 1993). Climate was warm. Frakes et al. (1992) indicate up to 27°C for the Late Oxfordian ocean surface temperatures according to oxygen isotopes measured on planktonic foraminifera and belemnites. Based on $\delta^{18}\text{O}$ analyses of Middle Oxfordian samples from the Swiss Jura platform, Plunkett (1997) also calculated palaeotemperatures of 26–27°C. In the Middle and Late Oxfordian, abundant rainfall in the hinterland allowed vegetation growth and continental run-off of siliciclastics into the study area (Gygi 1986; Pittet 1996). Associated nutrient input periodically led to eutrophication and to crises of coral reefs (Dupraz and Strasser 1999). Towards the Late Oxfordian (Berrense subzone) and Kimmeridgian, climate became more arid, as indicated by generally less siliciclastics but occurrence of evaporite pseudomorphs (Hug 2003; Rameil 2005). The siliciclastic-dominated Röschenz Member implies humid conditions (Fig. 1). In the overlying Hauptmumienbank Member, kaolinite is abundant but detrital quartz is present only in small quantities (Gygi and Persoz 1986). This may point to a northward shift of the siliciclastic source area and/or to diminishing rainfall in the hinterland. Only the floating clay minerals and organic matter reached the study area, whereas the quartz grains were left behind (Thiry 1982). The source area is not known but supposed to have been situated in the London–Brabant, Rhenish and Bohemian massifs, and/or in the Central Massif (Fig. 2). This general climatic evolution was controlled by high-frequency climate changes related to orbital (Milankovitch) cycles. These not only controlled siliciclastic input through periodically increased rainfall but also periodic evaporite formation (Pittet 1996; Hug 2003). Furthermore, they were responsible for low-amplitude sea-level fluctuations that led to the formation of hierarchically stacked depositional sequences (Strasser et al. 1999, 2000).

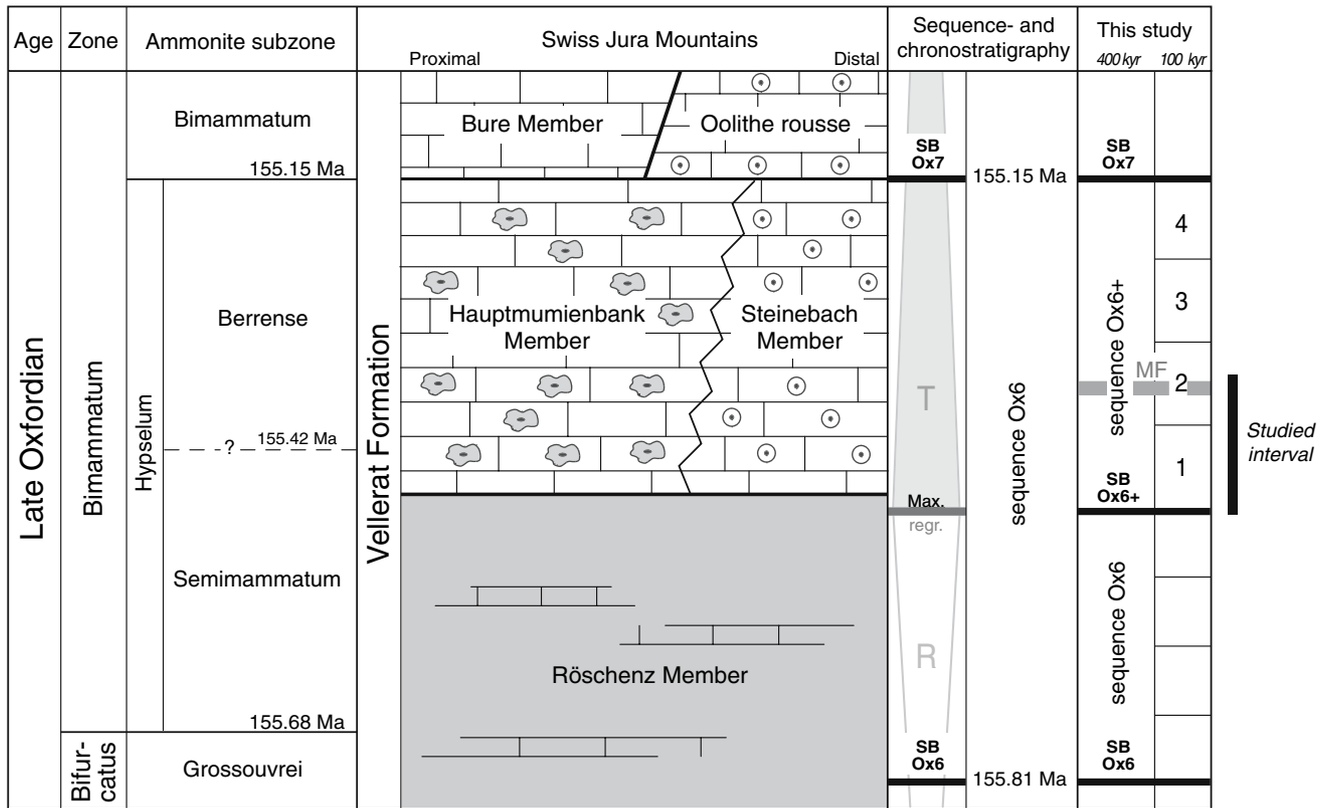


Fig. 1 Lithostratigraphic, biostratigraphic, and sequence-stratigraphic position of the Hauptmumienbank Member on the Swiss Jura platform (after Gygi and Persoz 1986; Gygi 1995, 2000a, 2000b; Hardenbol et al. 1998). The studied interval corresponds to one and a half small-scale (100-kyr) sequence between the transgressive surface of medium-scale

(400-kyr) sequence Ox 6+ at the base of the Hauptmumienbank Member and the maximum flooding of sequence Ox 6+. *SB* Sequence boundary, *MF* Maximum flooding, *R* Second-order regressive trend, *T* Second-order transgressive trend. Ages of ammonite subzone boundaries and of sequence boundaries according to Hardenbol et al. (1998)

Studied sections and methodology

The studied interval corresponds to the transgressive part of the Ox 6+ sequence (Hug 2003). Six platform sections (Voyeboeuf, Vorbourg, Hautes-Roches, Court, Savagnières, and Pertuis) were logged in detail and densely sampled (Figs. 3, 4, 5). These sections exhibit lagoonal facies: oncoïd, oïd, bioclastic, and peloid-rich deposits. Based on field observations, 300 polished sections and thin sections, plus 25 washing residues of marls, facies and microfacies have been analyzed. Analyses of the texture and relative abundance of skeletal and non-skeletal elements (Fig. 4) were performed from thin sections. These informations were then interpreted in terms of palaeoenvironments. Stacking pattern, bed surfaces, and facies were used for sequence and cyclostratigraphic interpretations (Strasser et al. 1999). This helped to correlate the sections and to trace the vertical and lateral evolution of facies types. Figure 6a shows an example of high-resolution sequence-stratigraphic correlations for two elementary sequences. Based on these correlations, platform evolution through time is reconstructed

(Fig. 6b). Based on the dominant facies, the sections are positioned from “proximal” (Voyeboeuf) to “distal” (Savagnières and Pertuis) in Fig. 7. However, because platform morphology changed through time, these attributions can only indicate the general trend.

Facies evolution and depositional sequences

The evolution of depositional environments through time is reconstructed from high-resolution facies analysis of the six studied sections. Facies evolution and stacking pattern of the beds lead to the definition of depositional sequences of different orders. Elementary sequences are the smallest units, in which one environmental cycle can be recognized (e.g., a deepening then shallowing trend, or an opening then closing of a lagoon as indicated by energy and faunal changes). In most cases, one elementary sequence corresponds to several beds. These elementary sequences stack into bundles (commonly of five elementary sequences) defining small-scale sequences, which again show charac-

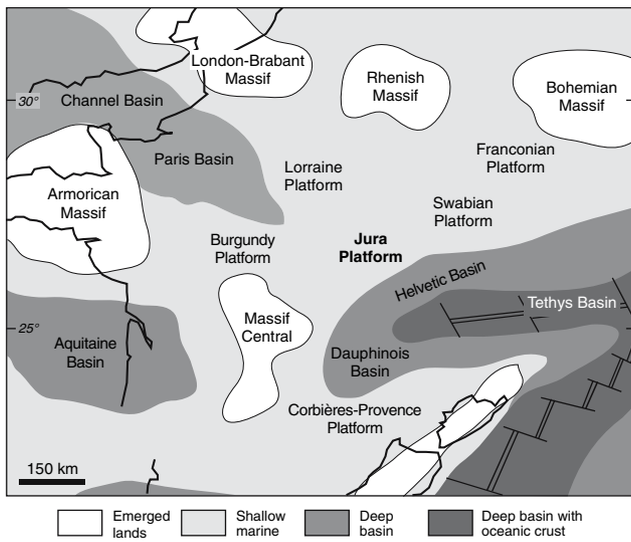


Fig. 2 Late Jurassic palaeogeography of the northern margin of the Tethys ocean, including the Jura platform, the Paris Basin, and the surrounding crystalline massifs (modified from Ziegler 1988; Thierry et al. 2000)

teristic facies evolutions. Four small-scale sequences then group into one medium-scale sequence. Depositional sequences of all scales can be interpreted in terms of sequence stratigraphy (Strasser et al. 1999). Boundaries between sequences correspond to the shallowest or most

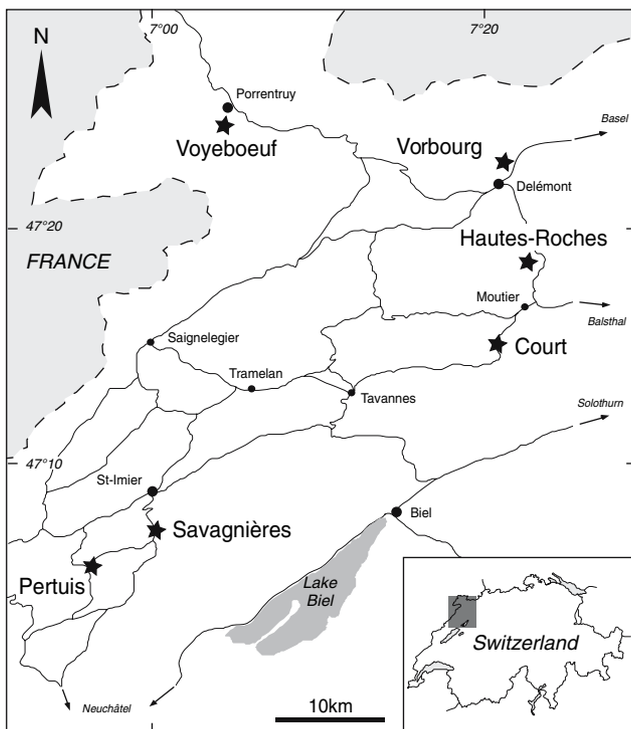


Fig. 3 Geographical position of the six studied sections in the Swiss Jura Mountains

FIELD OBSERVATIONS	COMPONENTS
<ul style="list-style-type: none"> limestone nodular limestone marl with lenses of limestone clay 	<ul style="list-style-type: none"> in thin section in the field
RELATIVE ABUNDANCE <ul style="list-style-type: none"> present common dominant + detrital quartz content - detrital quartz content 	<ul style="list-style-type: none"> • peloid type 1 oncoid type 2 oncoid type 3 oncoid type 4 oncoid oo-oncoid ○ radial ooid ● micritic ooid L <i>Lithocodium</i> B <i>Bacinella</i> × bioclast ostracode o bivalve o inoceram o oyster o gastropod ★ echinoderm o brachiopod o serpulid cluster of serpulids benthic foraminifer encrusted coral coral rubble o sponge o lithoclast o litho-oncoid OM organic matter o coal
TEXTURE <ul style="list-style-type: none"> c clay m marl M mudstone W wackestone P packstone G grainstone F floatstone R rudstone B boundstone 	
SEDIMENTARY STRUCTURES <ul style="list-style-type: none"> bioturbation bioporation cross-stratification 	
SEQUENCE- AND CYCLOSTRATIGRAPHY <ul style="list-style-type: none"> HD highstand deposits MF maximum-flooding zone TD transgressive deposits - TS - transgressive surface LD lowstand deposits - SB - sequence boundary 1.1 number of elementary sequence 	FACIES TYPES <ul style="list-style-type: none"> open - protected bioclastic peloidal oncolitic oolitic

Fig. 4 Legend to sections

protected facies, implying a drop of relative sea level. Rapid deepening or opening of facies is interpreted as a relative sea-level rise, and facies suggesting the relatively deepest water or the most open-marine conditions are attributed to the maximum flooding within the sequences. Once the depositional sequences and their hierarchical stacking are identified for each section, a correlation between the sections is attempted. Within the biostratigraphic framework of Gygi and Persoz (1986) and Gygi (1995), a fairly coherent sequence-stratigraphic correlation can be proposed. On the level of the small- and medium-scale sequences, the sequence boundary and maximum-flooding intervals in the Hauptmumienbank Member are

correlated over the whole study area (Hug 2003). On the level of the elementary sequences, correlation is sometimes difficult because facies evolution does not always display unequivocal deepening–shallowing or opening–closing trends. Furthermore, autocyclic processes partly controlled facies evolution (Strasser 1991).

Based on the absolute ages given by Gradstein et al. (1995), Hardenbol et al. (1998) dated SB Ox 6 at 155.81 Ma and SB Ox 7 at 155.15 Ma (Fig. 1). This implies a time span of 600–700 kyr between these two sequence boundaries. In the corresponding interval, Hug (2003) counted 6–8 small-scale sequences, each composed of five elementary sequences. The same number is found in the other sections that cover this interval (Pittet 1996; Hug 2003; Strasser et al. 2000). This suggests that a small-scale sequence formed within 75–117 kyr, and an elementary sequence within 15–23 kyr. These values coincide well with those of the first orbital eccentricity cycle (100 kyr) and of the precession cycle (between 18 and 22 kyr in the Oxfordian; Berger et al. 1989). It is implied that these orbital cycles induced climate changes that translated into sea-level fluctuations, which in turn controlled the formation of the observed depositional sequences.

Within this sequence and cyclostratigraphic framework, facies can now be correlated across the whole studied platform. The rapid vertical facies change between the Röschenz Member and the Hauptmumienbank Member corresponds to a major transgressive surface, which is manifest all over the Jura Mountains (Hug 2003). Facies distribution in the Oxfordian of the Swiss Jura is very heterogeneous and displays important lateral changes (Pittet 1996; Jordan 1999; Gsponer 1999; Hug 2003; Samankassou et al. 2003). Also the Hauptmumienbank Member, defined by its richness of oncoids, has a very inhomogeneous and diachronous distribution on the Jura platform (Fig. 7). Rapidly changing sediment thicknesses imply differential subsidence. This feature has also been recognized by Allenbach (2001), who related it to reactivation of ancient tectonic lineaments. Reefs formed on structural highs along the platform margin but could later find themselves in rapidly subsiding areas due to tectonic inversion. Ooid bars not only occurred on the platform margin but also in the platform interior where morphological highs created favorable conditions. Locally, wide channels brought relatively open-marine conditions into more central platform positions, generating high-energy depositional environments. Behind reefs and bars, lagoons were isolated or protected from open-marine influence and from wave or current energy (Fig. 6b; Hug 2003; Strasser and Védrine 2007). Lagoonal facies with oncoids shows a patchy distribution. No systematic relationship with the other facies is recognizable, although the most persistent ooid lagoons appear to be situated preferentially on the inner platform when compared to ooid bars and coral reefs.

Oncoids

The oncoids found in the Hauptmumienbank Member display a wide variability in size, shape, and composition. The following criteria have been used for defining types: surface morphology, size, cortex structure and composition, and sedimentological context.

Classification

Observed oncoid sizes vary between a few millimetres to a few centimetres. The oncoids have a well-defined rounded or irregular shape and smooth, wavy, or lobate contours. The final shape follows in many cases that of the nucleus, although a higher sphericity is obtained. If a nucleus is present, it is of bioclastic or lithoclastic nature and the envelope has a variable thickness. The cortex is characterized by micritic and/or organism-bearing laminations, which are continuous or discontinuous when the oncoid surface was eroded (Figs. 8, 9, 10). The types of oncoids are defined and differentiated by the surface morphology and by the presence, the nature, and the abundance of microencrusters in their cortex. Microfacies analysis of the encasing sediment furnishes the environmental context within which the oncoids were formed. The Hauptmumienbank oncoids are mainly found in wackestones, locally floatstones or rudstones, with bioclasts, ooids, and peloids. Transported micrite-dominated oncoids are also observed in ooid-rich grainstones. Texture, faunal composition, and sedimentary structures suggest a deposition in shallow-water lagoons.

Based on more than 300 thin sections and macroscopic samples from the Hauptmumienbank Member, four types of oncoids are distinguished (Fig. 11). Transitions between types also exist, suggesting a continuum.

Type 1 oncoids have diameters of a few millimetres and correspond to elliptical or spherical particles with relatively smooth contours. The cortex is micritic, homogeneous, and without microencruster inclusions (Fig. 8a, b). This pattern probably results from the trapping of fine-grained sediment on the oncoid surface by microorganisms. Laminations are difficult to distinguish. Type 1 oncoids are preferentially found in peloidal and bioclastic (foraminifera, bivalves, echinoderms) mudstones–wackestones of protected and moderate-energy lagoonal environments. This type of oncoid is relatively rare in the studied interval (Fig. 12).

Type 2 oncoids have diameters of a few millimetres (up to 1 cm) and present elliptical shapes with smooth contours. The cortex has irregular and locally truncated micritic laminae, and different growth phases can be distinguished in many cases (Figs. 8c, d, 9). This suggests relatively high-energy conditions with intermittent periods of wave agitation leading to a partial erosion of the cortex. The cortex of the micrite-dominated oncoids presents a relatively low-diversity

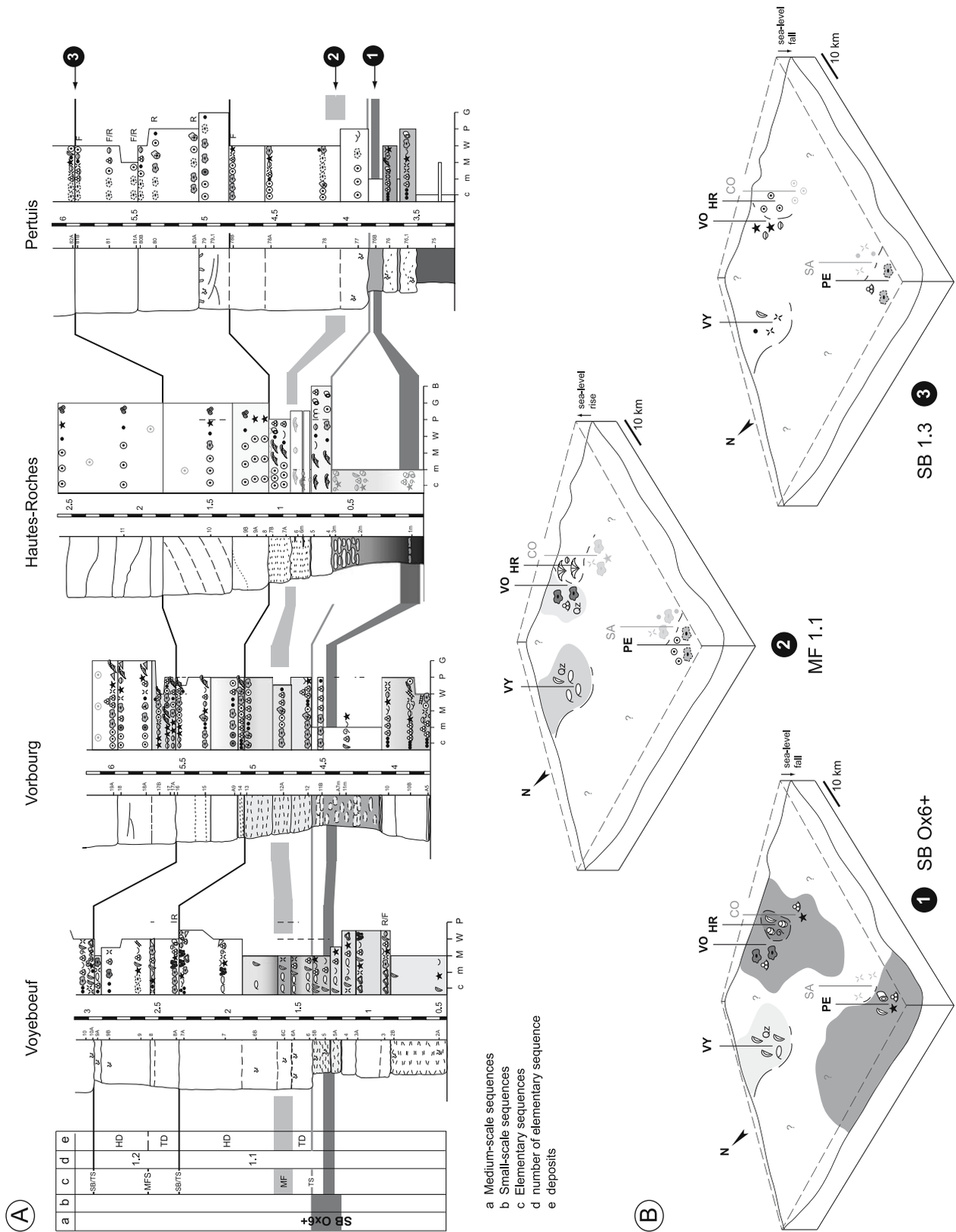


Fig. 6 **a** Example of high-resolution correlation of Voyeboeuf, Vorbourg, Hautes-Roches, and Pertuis for two elementary sequences. For symbols, refer to Fig. 4. **b** Schematic sketches showing the facies distribution on the Swiss Jura platform (1 SB Ox 6+; 2 MF 1.1; 3 SB 1.3)

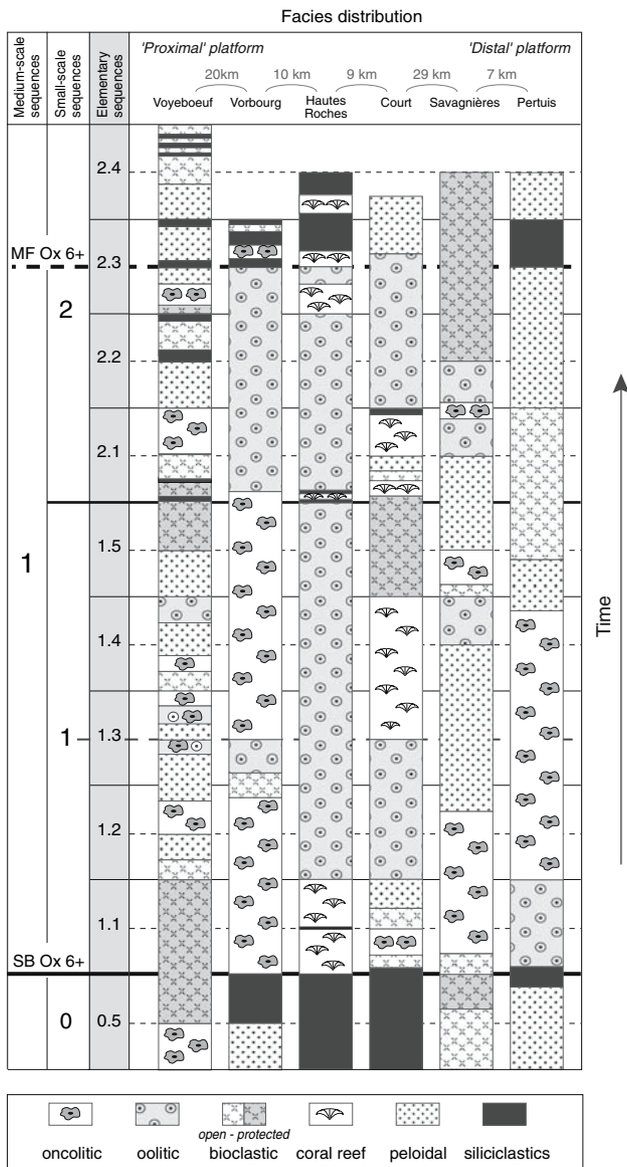


Fig. 7 Distribution of major facies types through time within the studied interval across the Swiss Jura platform. The duration of elementary sequences is considered to be equal (20 kyr)

fauna such as serpulid worms and *Bullopora*. Occasionally, ooids are incorporated in the cortex (Fig. 9c, d). These oncooids are found in packstones with normal-marine or semi-restricted fauna (brachiopods, oysters, foraminifera, echinoderms, bivalves, serpulids, ostracodes) and also in marls. Type 2 oncooids are common in all studied sections (Fig. 12).

Type 3 oncooids have diameters of a few millimetres to a few centimetres (up to 5 cm) and sub-elliptical shapes with wavy contours. The cortex is made of alternating organism-bearing and thin micritic laminations (Fig. 10). The organism-bearing laminations are formed by two microencrusters: *Bacinella irregularis* (Radoicic 1959; Fig. 10d) and

Lithocodium aggregatum (Elliott 1956; Fig. 10e). *Bacinella irregularis* is an enigmatic microencruster with an irregular micritic meshwork and interspaces filled with calcite spar. This is assumed to represent a cyanobacterial structure (e.g., Schmid 1996; Dupraz 1999; Shiraishi and Kano 2004). *Lithocodium*, originally interpreted as a codiacean alga by Elliott (1956), is characterized by inner cavities and an aggregated outer wall with numerous alveoli (complex wall structure) probably containing symbiotic and photosynthetic algae (Schmid and Leinfelder 1996). These authors thus attributed it to a loftusiacean foraminifer with an encrusting life habit. Recently, Cherchi and Schroeder (2006) interpreted *Lithocodium* as colonies of calcified cyanobacteria because of the absence of apertures connecting neighboring cavities, and because of the very irregular form and arrangement of these hollows. These *Bacinella–Lithocodium* oncooids can be compared to the porostromate oncooids of Peryt (1981) because of the presence of cyanobacteria in their cortex. The alternation of micrite laminae and organism-bearing laminae suggests calm periods during which *Bacinella–Lithocodium* could grow and more agitated periods when micrite laminae are formed by the rolling on the lagoon floor. Type 3 oncooids are preferentially found in wackestones with normal-marine fauna (brachiopods, oysters, foraminifera, echinoderms, bivalves) indicating relatively low-energy conditions. The *Bacinella–Lithocodium* association characterizes lagoonal environments with oligotrophic conditions, low accumulation rate, clear, oxygenated, shallow, and normal-marine waters (e.g., Leinfelder et al. 1993; Dupraz and Strasser 1999; Immenhauser et al. 2005). Dupraz and Strasser (1999), who examined the encrustations of Oxfordian coral reefs in the Jura, correlated the *Bacinella–Lithocodium* association with episodes of relatively high coral diversity. *Bacinella* and *Lithocodium* have also been identified in the microbial crusts associated with coral and sponge reefs of the Late Jurassic (e.g., Helm and Schülke 1998; Dupraz and Strasser 1999; Schmid and Leinfelder 1996; Olivier et al. 2004; Shiraishi and Kano 2004). This may suggest that palaeoecological conditions for *Bacinella–Lithocodium* bearing oncooids were analogous. The presence of the benthic foraminifer *Mohlerina basiliensis* (Mohler 1938; Schlagintweit and Ebli 1999; Schlagintweit et al. 2005), preferentially in wackestones with type 3 and/or 4 oncooids, suggests that oncooids and benthic foraminifera were, at least partly, controlled by similar environmental parameters.

Type 4 oncooids have diameters of a few millimetres to several centimetres (up to 10 cm) and display irregular shapes with lobate contours. Exclusively composed of *Bacinella* and/or *Lithocodium* meshwork, they have no nucleus or laminations (Fig. 10f). Type 4 oncooids are found in bioclastic wackestones, commonly associated with type 3 oncooids. Low-energy conditions with no or only rare

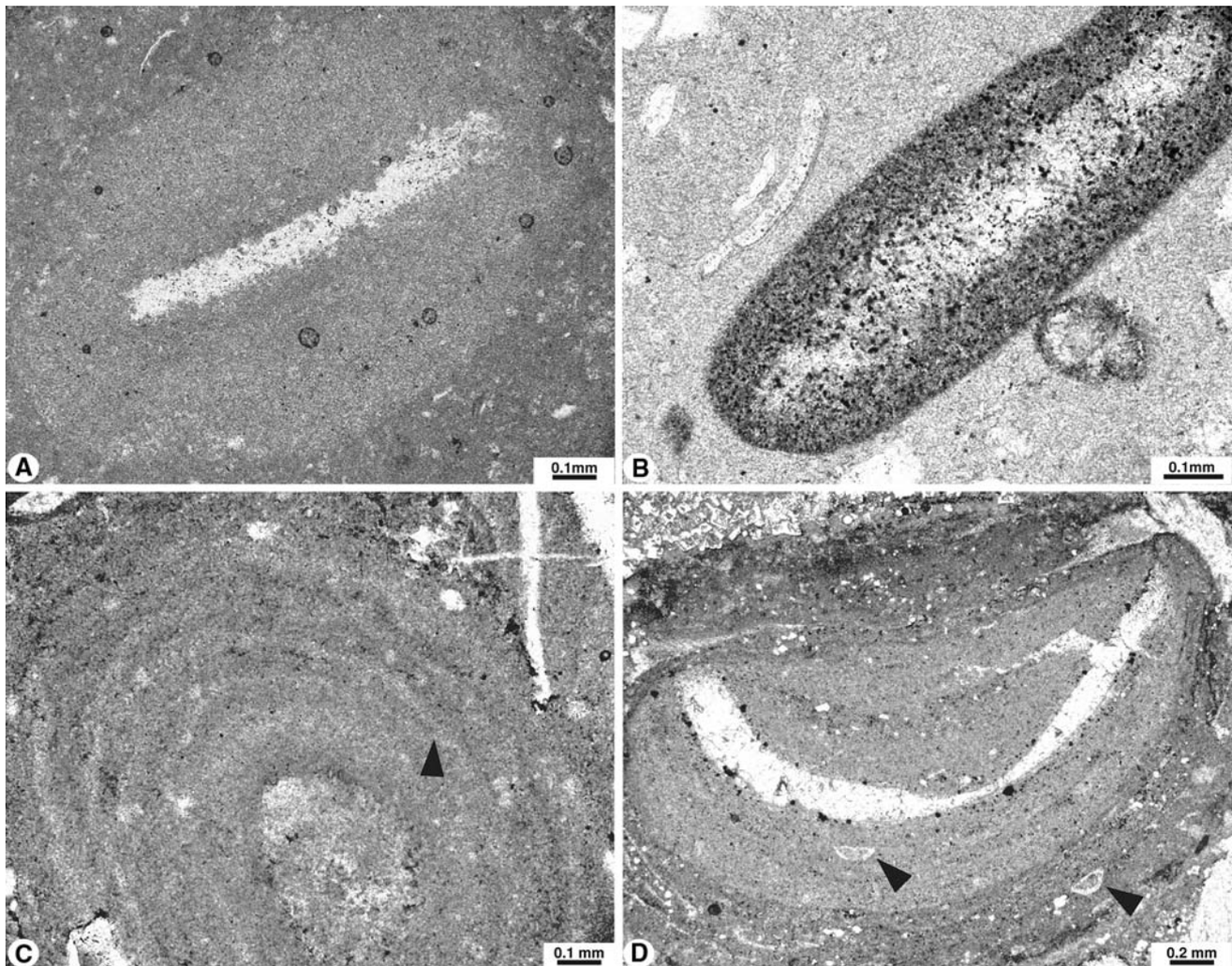


Fig. 8 **a** Type 1 oncooid with echinoderm fragment as nucleus and homogeneous, micritic cortex with barely visible laminations, found in a bioclastic wackestone. Savagnières section, thin section Sa-32C. **b** Type 1 oncooid with bioclastic nucleus and thin micritic cortex without laminations. Court section, thin section BPCo-55. *Black dots* are

pyrite. **c** Elliptical type 2 oncooid with a micritic, laminated cortex and some sparry patches (dissolved organisms?) Truncated laminae appear locally (*arrow*). Pertuis section, thin section Pe-80A. **d** Type 2 oncooid enclosing the foraminifer *Bullopورا* sp. (*arrows*). Vorbour section, thin section Vo-K

agitation are suggested. Locally, type 4 oncooids develop into carpet-like structures and build-ups of *Bacinella* (e.g., Hug 2003; Immenhauser et al. 2005).

Oncoid distribution

From the high-resolution sequence-stratigraphic framework, a time-space diagram illustrating the distribution of oncooid types is synthesized (Fig. 12). A proximal–distal trend is revealed by the greater abundance of type 3 oncooids in the “distal” platform sections (Pertuis and Savagnières); while in the “proximal” section (Voyeboeuf) types 2 and 1 oncooids predominate. Hautes-Roches and Court, which are mainly composed of ooid-rich deposits and patch-reefs, almost exclusively present type 2 oncooids. The different

oncooid types coexist, except types 1 and 4 that occur in very different environmental settings concerning mainly water energy (Fig. 11). Regardless of these general trends, it appears that local conditions (e.g., clay input) played an important role in oncooid development (Fig. 7). Migrating ooid bars locally interrupted the oncooid sedimentation.

A specific evolution of oncooid types is occasionally observed within small-scale and elementary depositional sequences. Oncooids are generally more abundant in the first small-scale sequence than in the second one (Fig. 7). Based on the high-resolution sequence-stratigraphic framework, a detailed description of the oncooid type distribution is given for the Vorbour section (Fig. 5). The lowstand and transgressive deposits of the first small-scale sequence contain abundant type 2 oncooids (Figs. 5, 7). Migration of ooid bars locally interrupts oncooid deposits in the upper part of the

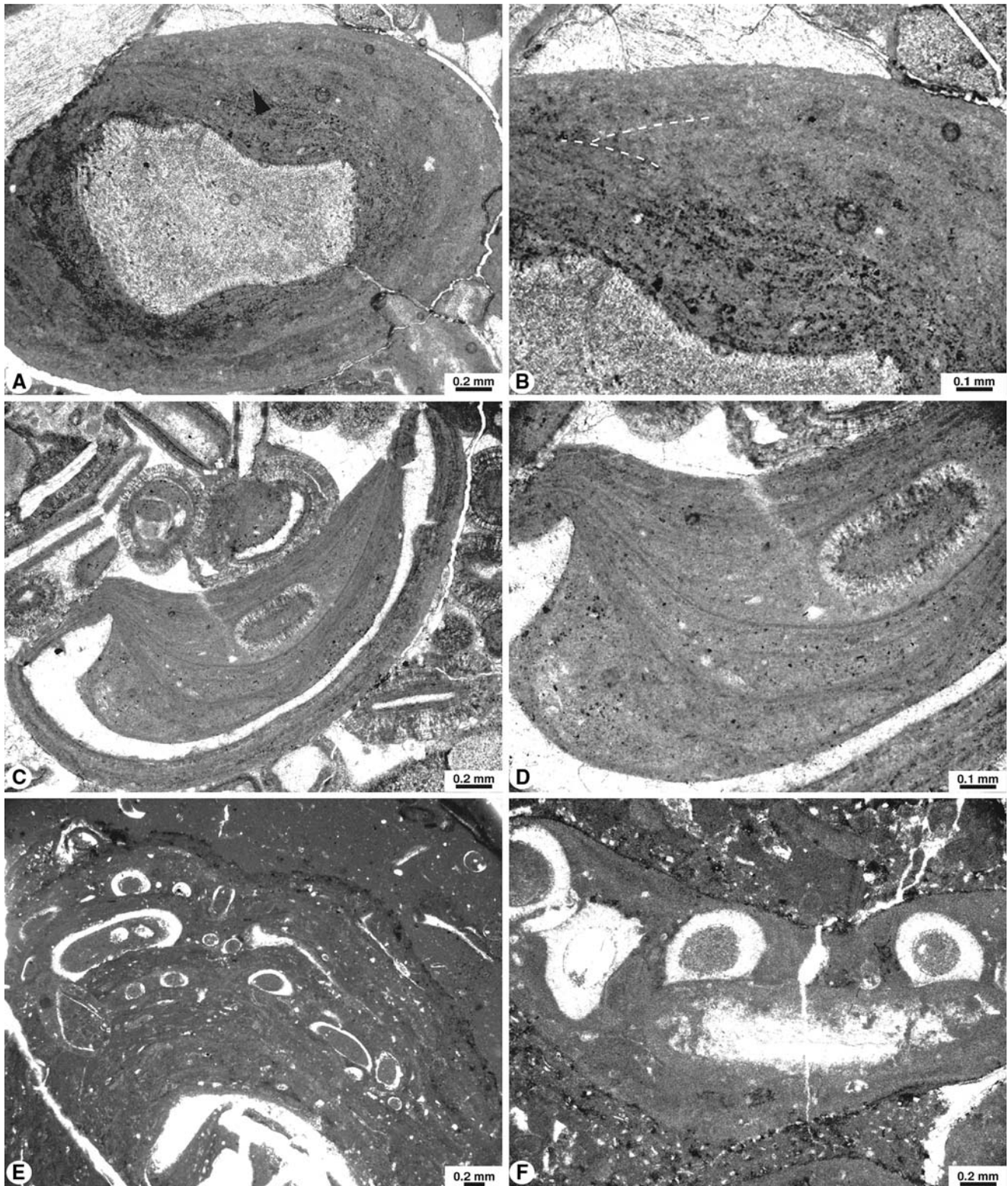


Fig. 9 Type 2 oncoids. **a** Elliptical oncoid with a large echinoderm fragment as nucleus and smooth contours. Micritic laminations of a first growth phase are overlapped by those of a second phase (*arrow*). Court section, thin section BPCo-81. **b** Detail of (**a**) showing the overlapping micritic laminae. Court section, thin section BPCo-81. **c** Elongate type 2 oncoid. In a first growth phase the laminae followed the shape of the nucleus. A radial ooid was then trapped, and in a second growth phase the cortex strived

towards a more and more rounded shape. Court section, thin section BPCo-81. **d** Detail of (**c**) showing the trapped ooid. Court section, thin section BPCo-81. **e** Type 2 oncoid with a bioclastic nucleus and irregular micritic laminae. Serpulid worms encrusted and then were incorporated into the cortex. Voyeboeuf section, thin section Vy-8A. **f** Micritic oncoid formed by several phases of growth. Large serpulid worms encrusted at different growth stages. Voyeboeuf section, thin section Vy-12Cs

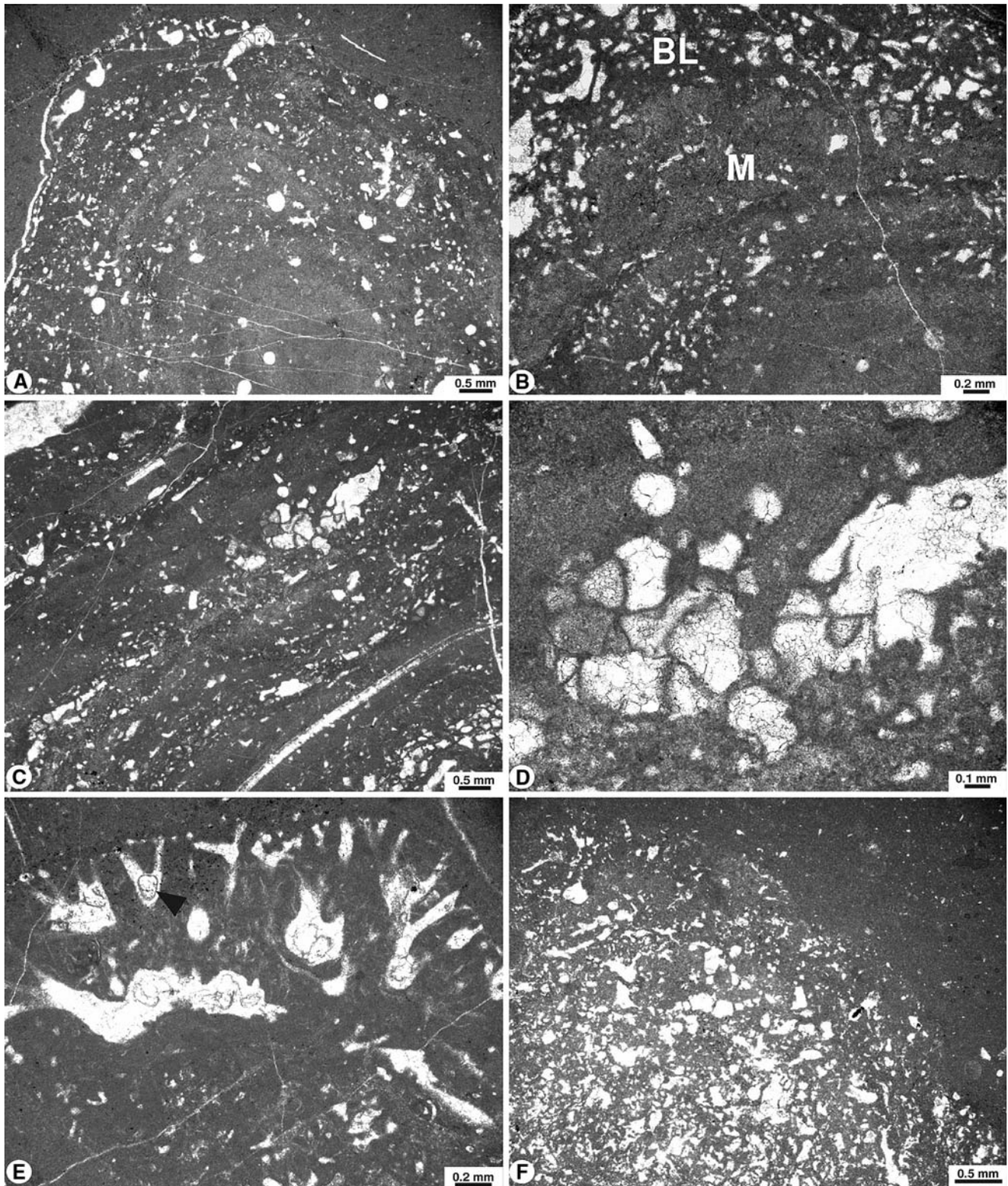


Fig. 10 Types 3 and 4 oncoids. **a** Type 3 oncooid with a cortex composed of irregular organism-bearing *Bacinella*–*Lithocodium* laminations and thinner micritic laminations. The contour is lobate. Pertuis section, thin section Pe-81A. **b** Detail of type 3 oncooid with *Bacinella*–*Lithocodium* meshwork BL and micritic laminations M. Pertuis section, thin section Pe-82A. **c** Detail of a type 3 oncooid cortex. Pertuis section, thin section Pe-82A. **d** Detail of the vesicular structure of

Bacinella irregularis. Pertuis section, thin section Pe-82A. **e** *Lithocodium aggregatum* with cavities colonized by cryptobiontic foraminifera (probably *Troglotella incrustans*; arrow) and a complex wall structure. Pertuis section, thin section Pe-84A. **f** Type 4 oncooid with lobate and diffuse contours, entirely composed of *Bacinella irregularis*. Voyeboeuf section, thin section Vy-20

		Micrite-dominated oncoids		<i>Bacinella-Lithocodium</i> oncoids	
		type 1 	type 2 	type 3 	type 4 
Morphological criteria	Prevailing size	few millimeters	few mm to 1 cm	few cm to 5 cm	few mm to 10 cm
	Shape and surface	elliptical to spherical smooth	elliptical smooth	elliptical lobate	elliptical lobate
	Nucleus	bioclastic or lithoclastic nucleus		bioclasts or without nucleus	no nucleus recognizable
Cortex configuration and composition	Lamination type	concentric and continuous but barely visible	irregular and truncated	rarely continuous	no lamination
	Microfabric	micritic laminations or homogeneous	micritic laminations and organism-bearing laminations	organism-bearing laminations and few micritic laminations	microbial meshwork
	Bioclasts	rare	common bioclasts		rare
	Microencrusters	no fauna associated	serpulid worms, <i>Bullopore</i>	abundant microencrusters: <i>Bacinella</i> , <i>Lithocodium</i>	
Surrounding sediment	Classification	mudstone - wackestone	packstone and locally wackestone	mudstone - wackestone and locally packstone	
	Associated fauna and grains	mainly benthic foraminifera, peloids, and ooids		brachiopods, commonly <i>Mohlerina basiliensis</i> , bivalves	echinoderms, commonly <i>Mohlerina basiliensis</i>
Depositional environment	Relative energy	moderate and intermittent	high and intermittent	low and intermittent	very low
	Bathymetry	subtidal			
	Sedimentation rate	low			
	Environment	protected lagoon	fully marine lagoon		

Fig. 11 Classification of the oncoids encountered in the Hauptmümbank Member. Morphological criteria, cortex configuration and composition, surrounding sediment, and depositional environmental conditions are indicated

transgressive deposits of elementary sequence 1.3. Around and after the maximum flooding of this elementary sequence, types 3 and 4 oncoids appear. In the two last elementary sequences, type 3 oncoids dominate and type 2 oncoids reappear. Small-scale SB 2 is marked by miliolid foraminifera and type 3 oncoid rudstones. Within the elementary sequences of Vorbour, no particular trend is observed. However, in the Voyeboeuf and Savagnières sections (Fig. 13), some elementary sequences display the same evolution as the small-scale sequence of Vorbour. Elementary sequence 1.5 in Savagnières presents two beds, interpreted, respectively, as transgressive and highstand deposits. The transgressive bed can be subdivided into bioclastic wackestone with normal-marine fauna and oncoid-rich wackestone-floatstone. The maximum flooding is marked by types 3 and 4 oncoid mudstone to floatstone. The highstand deposit is dominated by peloidal and bioclastic wackestone. The small-scale sequence boundary is placed at the base of a serpulid bioherm with peloids and oysters. The next elementary sequence 2.1 has a lowstand deposit rich in peloids, serpulids, oysters, textulariid foraminifera, and a few type 2 oncoids. The transgressive surface is positioned at the base of an ooid–peloid–bioclastic wackestone–packstone, corresponding to abandoned bars. In Voyeboeuf, elementary sequence 1.5 is composed of ostracode and textulariid foraminifer mudstones (Fig. 13). The highstand deposit of

this elementary sequence consists of clays with coal debris, detrital quartz, and serpulids, indicating relatively proximal and semi-restricted conditions and two thin wavy limestone beds composed of ostracode and foraminifer mudstone. The small-scale sequence boundary is placed at 6.6 m in a clay-rich level. The transgressive deposit consists of foraminifer mudstone with some type 1 oncoids. The maximum flooding is placed around thin wavy beds surrounded by marls. The highstand deposit is made of a massive bed, which is rich in types 3 and 4 oncoids and benthic foraminifera at the base. In its upper part, type 3 oncoids and foraminifera (textulariid and miliolid) dominate. The next sequence boundary SB 2.2 is marked by peloid and foraminifer (miliolid and less textulariid) wackestone rich in detrital quartz. In this lowstand deposit, some type 2 oncoids are observed.

This study on the spatial and stratigraphic distribution of oncoids in lagoonal deposits shows that types 1 and 2 oncoids have both a similar distribution and preferentially form around the sequence boundaries and during transgressive conditions of elementary and small-scale sequences, indicating that enough energy was furnished to roll the particles (Figs. 12, 14). Types 3 and 4 oncoids have an inverse occurrence because they mainly occur around small-scale and elementary maximum-flooding periods and in highstand deposits, characterizing periods of low-energy and increase of *Bacinella–Lithocodium* growth (Figs. 12, 14).

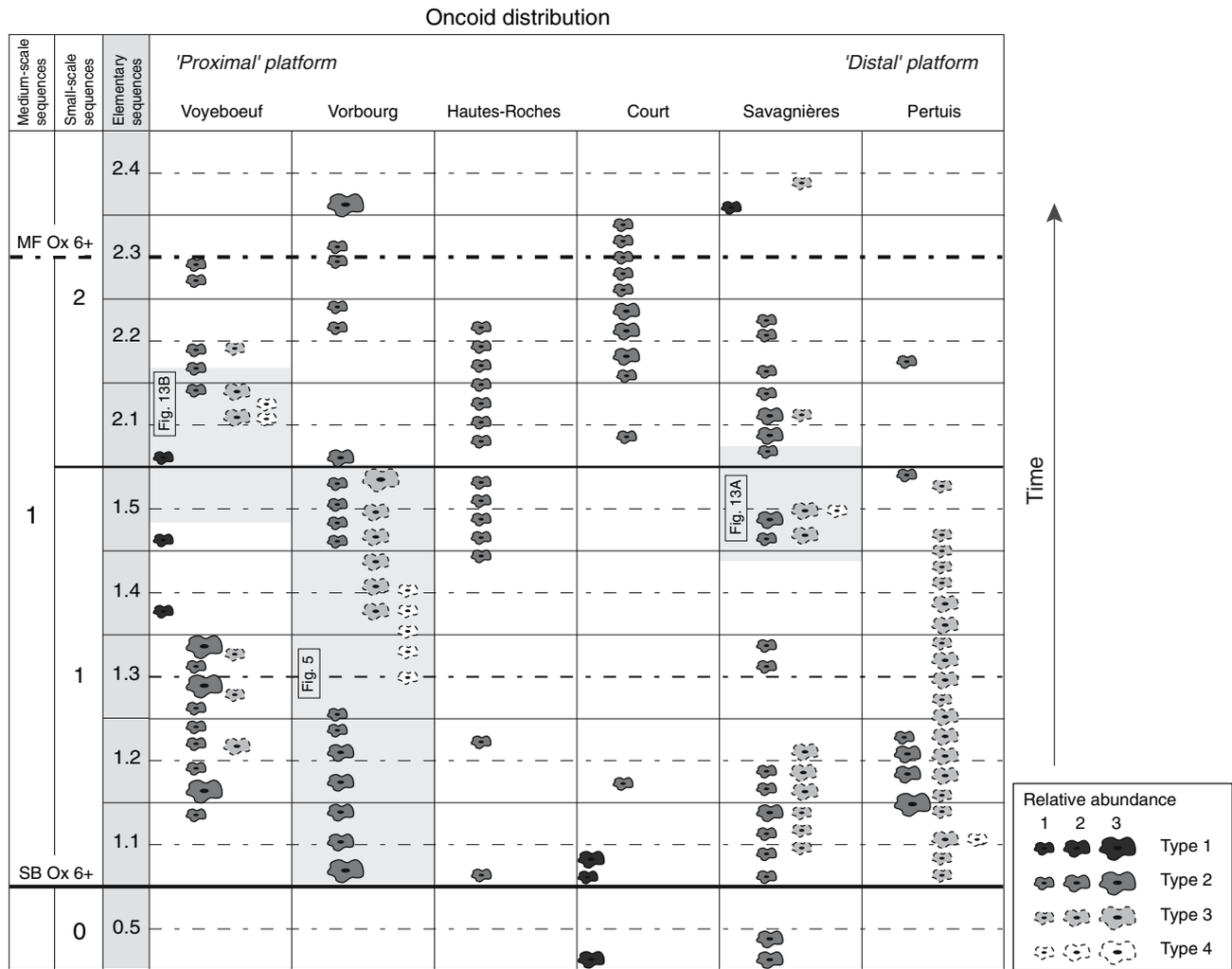


Fig. 12 Distribution of oncoïd types in the elementary sequences within the transgressive deposits of the Ox 6+ medium-scale sequence for the six studied sections. Sections are arranged from “proximal” to

“distal” platform. Relative abundance (1 present, 2 common, 3 abundant) for each oncoïd type is estimated from thin section analysis. Intervals detailed in Figs. 5, 13 are shaded

Controlling factors

Oncoïd growth and distribution are mainly controlled by sea level and climate (Figs. 14, 15). The main direct environmental parameters are water energy, water depth, trophic level, and accumulation rate. Their influence is recognized in the stratigraphic and spatial distribution of oncoïds in large-scale to elementary sequences as well as in the oncoïd surface morphology, the biotic composition of the oncoïd cortex, and the encasing sediment.

Water energy and water depth

The oncoïd-rich facies of the Hauptmumienbank Member appears at the beginning of a second-order sea-level rise, covers the entire medium-scale Ox 6+ sequence and then disappears above the third-order SB Ox 7 (Fig. 1). This stratigraphic distribution points to a link with changes of rel-

ative sea level. Sea-level fluctuations play an important role in opening and closing shallow lagoons behind barrier systems and thus strongly modify environmental factors (e.g., water energy, oxygenation, salinity, temperature, sediment transport). The specific evolution of oncoïd types within small-scale and elementary sequences confirms the link also to high-frequency sea-level fluctuations. The sub-spherical types 1 and 2 oncoïds, mainly resulting from sediment trapping by micro-organisms, preferentially occur around sequence boundaries and in transgressive deposits where higher energy conditions predominated in relatively shallow water (Fig. 14). This implies that these particles needed energy to be overturned and rolled. On the other hand, the lobate types 3 and 4 oncoïds are characteristic of low-energy environments where microbial meshwork had time to grow. Types 3 and 4 oncoïds are concentrated around maximum floodings and in highstand deposits where low-energy conditions existed in relatively deep water but still in the photic

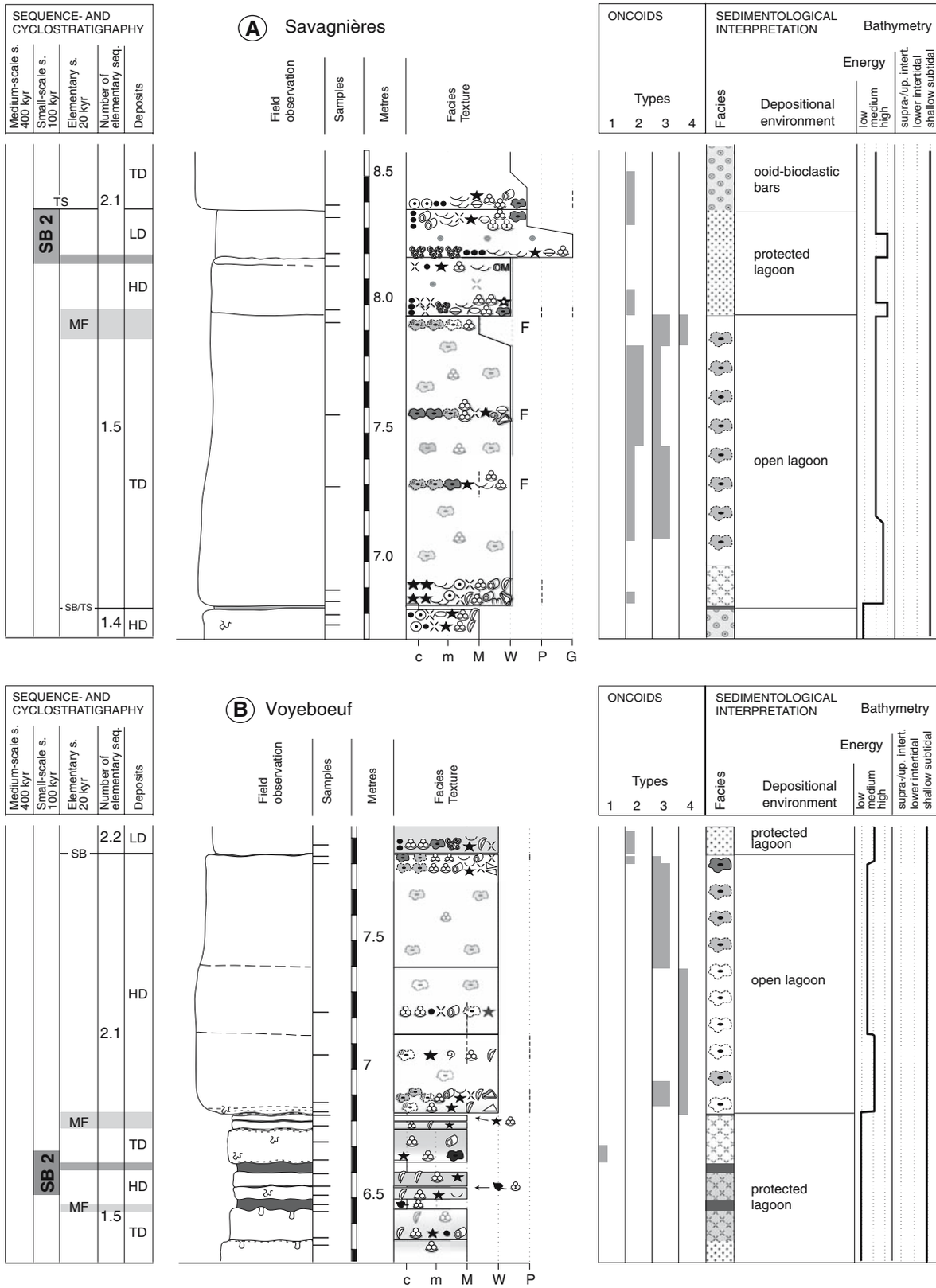


Fig. 13 Selected elementary sequences of the Savagnières and Voyeboeuf sections around small-scale sequence boundary SB 2, showing the distribution of the four oncoïd types, facies types, the palaeoenvironmental interpretation, and the sequence-stratigraphic framework. For discussion, refer to text. Legend of symbols in Fig. 4

ronmental interpretation, and the sequence-stratigraphic framework. For discussion, refer to text. Legend of symbols in Fig. 4

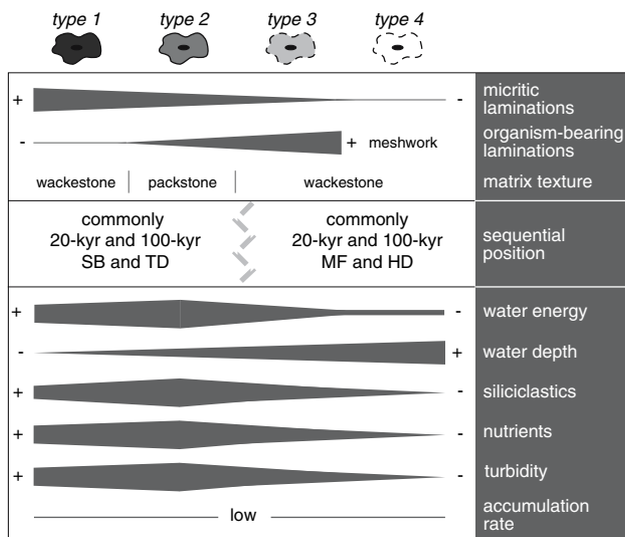


Fig. 14 Oncoïd types, sequential position, and main controlling factors. For discussion refer to text

zone (Fig. 14). Consequently, the oncoïd surface morphologies indicate the intensity of rolling on the sea floor and thus the degree of wave or current energy and, indirectly, the water depth. The biotic composition and diversity in the oncoïd cortex and the encasing sediment (e.g., Dunham texture, faunal association) provides additional information on water energy and water depth.

Platform morphology was created by differential subsidence of tectonic blocks and by differential sediment accumulation (e.g., Pittet 1996; Allenbach 2001; Hug 2003; Wetzel et al. 2003). The already existing highs and depressions were

further accentuated by the increase of carbonate production in reefs and on oïd shoals. Oncoïds were preferentially accumulated in depressions. Hallock (1988) also suggests that the point sources of terrigenous input and the platform morphology increase the spatial heterogeneity of nutrient distribution and thus trophic level. Consequently, variable platform morphology was partly responsible for the patchy distribution of oncoïd-rich facies and oncoïd types (Figs. 7, 15).

In the studied sections, some oncoïd-rich beds are capped by oïd bars, which may have been controlled by autocyclic processes (Fig. 7). On shallow carbonate platforms, such processes are inherent and involve lateral migration of sediment bodies (Ginsburg 1971; Pratt and James 1986; Strasser 1991). Consequently, autocyclic processes were superimposed on the orbitally controlled sea-level fluctuations and represented an additional factor in oncoïd growth and distribution.

Accumulation rate

Carbonate accumulation rates mainly depend of the local carbonate production but can also be modified by carbonate transport (supply or exportation). On the Swiss Jura platform, the accumulation rate of oncoïd-rich lagoons has been estimated at around 0.11 mm/year (after sediment decompaction), which is low in comparison to modern lagoonal environments (Strasser and Samankassou 2003; Strasser and Védrine 2007).

Microbial crusts commonly grow slowly. Their occurrence is considered to be diagnostic for low background

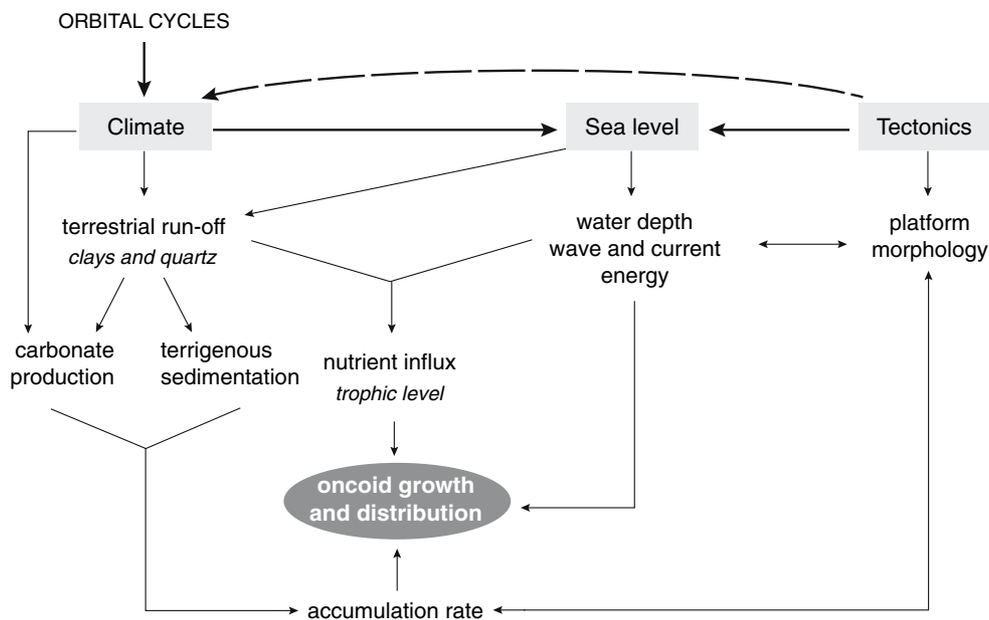


Fig. 15 Interrelated factors controlling oncoïd growth and distribution. For discussion refer to text

sedimentation rates (Leinfelder et al. 1993). Similar requirements are probable for oncoids, which are partly or totally composed of the same microencrusters such as the *Bacinella–Lithocodium* association (e.g., Peryt 1983; Dupraz 1999).

Trophic level

The deposition of the massive Hauptmumienbank limestones on top of the Röschenz marls points to a general trend toward a lower input of siliciclastics and a higher carbonate productivity. Decreasing terrigenous input may have been related to a change to less humid climate conditions in the hinterland.

In the studied sections, oncoids occur abundantly in limestones and occasionally in marls. In the limestone beds, types 1–4 oncoids are present whereas in the marly intervals only type 2 oncoids are found. The biotic composition of the oncoid cortex and the encasing sediment imply specific trophic conditions (Fig. 11). Type 2 oncoids enclose serpulids and textulariid foraminifera, which tolerate mesotrophic to eutrophic conditions (e.g., Brasier 1995; Fugagnoli 2004). The concomitant growth of type 2 oncoids with such organisms and their occurrence in clay-rich environments suggest that they were adapted to low water transparency and potentially high trophic levels. Type 1 oncoids are rare in the studied sections and found in limestone beds. They show affinity with type 2 oncoids: the scarcity of associated fauna probably indicates mesotrophic or eutrophic conditions, making life difficult for carbonate producing organisms. Types 3 and 4 oncoids are constituted of light-dependent microencrusters (i.e., *Bacinella* and *Lithocodium*) implying clear water and oligotrophic conditions (Dupraz and Strasser 1999; Fig. 14). They occur in the massive limestone beds, which formed when siliciclastic and nutrient input was reduced. Growing under water on the sediment surface, oncoids thus monitor water transparency and trophic level, which are both driven by the episodic influx of terrigenous sediment (quartz, clays, organic matter, and nutrients) onto the shallow carbonate platform.

Conclusions

Oncoid formation and distribution were studied in the Hauptmumienbank Member (Late Oxfordian, Swiss Jura) within a precise time framework obtained by sequence and cyclostratigraphy. Four types of oncoids are defined based on surface morphology, configuration and composition of the cortex, and the encasing sediment. Type 1 is sub-elliptical, micritic, and has a homogeneous cortex; Type 2 is elliptical, smooth, and displays micritic laminations with serpulid worm tubes and *Bullopora*; Type 3 is elliptical,

lobate, and shows alternating laminae of *Bacinella–Lithocodium* and micrite; Type 4 is lobate and consists mainly of a *Bacinella–Lithocodium* meshwork.

The elementary and small-scale depositional sequences, in which the oncoids occur, are related to the 20-kyr orbital precession cycle and to the 100-kyr eccentricity cycle, respectively. Thus a high-resolution time framework for the study oncoid formation and distribution is given. The stratigraphic and spatial distribution of oncoids in the six studied sections shows that each oncoid type can be attributed to a specific depositional environment. Type 1 oncoids characterize moderate-energy conditions in protected lagoons; Type 2 oncoids are related to higher energy conditions in open lagoons; Types 3 and 4 oncoids dominate in low to very low-energy settings in open lagoons. Moreover, oncoid distribution is linked to relative sea-level changes. Micrite-dominated oncoids (types 1 and 2) are preferentially found around sequence boundaries and in transgressive deposits, whereas oncoids containing *Bacinella–Lithocodium* (types 3 and 4) are found rather around maximum floodings and in highstand deposits. This implies that changes in water energy and water depth were direct controlling factors. The heterogeneous oncoid distribution through space and time implies that the variable morphology of the Swiss Jura platform was also a controlling factor. As for coral reefs and microbialites, oncoid formation required relatively low sediment accumulation rates. Humidity changes in the hinterland controlled terrigenous run-off (clays and nutrients) and thus carbonate production. Types 1 and 2 oncoids grew under mesotrophic to eutrophic conditions, while types 3 and 4 oncoids flourished in oligotrophic waters. This study demonstrates that oncoids are valuable proxies for high-resolution palaeoenvironmental and palaeoecological studies.

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