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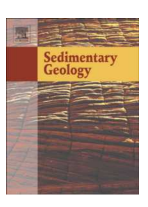
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1

Gypsum lithofacies and stratigraphic architecture of a Messinian marginal basin (Piedmont Basin, NW Italy)

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ABSTRACT

During the Messinian salinity crisis (5.97 5.. 3 Ma) the Mediterranean was transformed into the youngest salt giant in Earth history. In marginal Mediterranean subbasins, the first stage of the crisis is recorded by up to 16 it lo ogical cycles made up of shale/ gypsum couplets (Primary Lower Gypsum unit, PL3), making a downslope transition to organic-rich shales. Gypsum comprises a wide array of lithofacies, including bottom-grown gypsum, cumulate deposits (i.e., resulting from gypsum nucleation in the water column and subsequent deposition on the seaflood) and the branching selenite lithofacies. The latter appeared synchronously in many sections of the Mediterranean in the 6th PLG cycle (i.e., at about 5.87 Ma). To investigate the still enigmatic stratigraphic architecture of the lateral lithofacies transition from the shallower basin margin to the depocenter and the paleonvironmental modifications driving the development of the branching selenite lithofacies, we carried sedimentological, petrographic and geochemical analyses of the different types of gypsum lithofacies exposed along a margin-to-depocenter transect in the Piedmont Basin (NW Italy), corresponding to the northernmost Mediterranean subbasin. The results of this study suggest

that the lateral and vertical distribution of gypsum lithofacies mostly reflects the position of an oxycline with respect to the seafloor. Bottom grown gypsum (cycles PLG1-PLG5) was deposited on a temporarily oxygenated seafloor above the oxycline in the marginal part of the basin. In the case of cumulate deposits (laminar gypsum and gypsiferous mudstones, cycles PLG6-PLG15), gypsum nucleated within the water column at the oxycline and was eventually deposited on an anoxic seafloor. The branching selenite lithofacies corresponds to a syngenetic product formed below the seafloor at the expenses of cumulate gypsum.

Deposition of cumulate gypsum and formation of the branching seachite were controlled by the balance between the rate of sulfate reduction in bottom waters and nucleation of gypsum in the water column. The appearance of cumulate gypsum and the branching selenite from the 6th PLG cycle upwards reflects the intensification of water column stratification in response to increased isolation of the basin.

Keywords: Gypsum, branching selenite, rumulate deposits, water column stratification, Messinian salinity crisis.

1. Introduction

Thick successions of gy_F sum were deposited in the Mediterranean basin during the Messinian salinity crisis (MSC; 5.97-5.33 Ma), when the Mediterranean was turned into the youngest salt giant in Earth history (Hsü et al., 1973; Krijgsman et al., 1999; Roveri et al., 2014; Flecker et al., 2015). The sedimentary products of the MSC are more than 1 million km³ of evaporites (carbonate minerals, gypsum, anhydrite and halite) (Ryan, 2008, Haq et al., 2020) that are found in a variety of onland marginal subbasins as well as in offshore basinal areas of Western and Eastern Mediterranean, as suggested by seismic data (Lofi et al., 2011; Raad et al., 2020), (Fig. 1A). Messinian gypsum comprises both in situ-precipitated and

detrital deposits (Roveri et al., 2014). In situ deposits were grouped in two stratigraphic units referred to as the Lower Evaporites (containing also the thick massive halite and K/Mg salts of deep basins and Sicily), and Upper Evaporites (Rouchy and Caruso, 2006). More recently an alternative terminology was proposed, following the three stage model of the MSC events that is mostly based on outcrop studies of key sections from different Mediterranean marginal basins (CIESM, 2008; Manzi et al., 2013; Roveri et al., 2014), including the Piedmont basin (Dela Pierre et al., 2011). According to this model, that is not unanimously accepted by the scientific community (e.g., Meilijson et al., 2019), in situ primary gypsum is ascribed to (a) the Primary Lower Gypsum (PLG) unit, representing the first stage of the MSC (5.97-5.60 Ma; Manzi et al., 2013), and (b) the Upper Gypsum, recording the third stage of the MSC (5.55-5.33 Ma) (Roveri et al., 2014) (Fig. 1B). Detrital, resedimented gypsum was deposited during the second stage of the crisis (5.60-5.55 Ma) forming the Resedimented Lower Gypsum unit that apparently interfingers vi'n halite (Roveri et al., 2014). The PLG unit, the object of this paper, is composed of up 10 lithological cycles consisting of shale (and marl)/gypsum couplets (Lugli et a¹ 2 11 d). Deposition of the lithological cycles is interpreted to be controlled by orbitally-dil en climate oscillations (precession) (Krijgsman et al., 1999). Shales would be indicative of relatively humid climate during precession minima (insolation maxima), whereas gyps, m is thought to reflect more arid conditions during precession maxima (insolation minima). The PLG unit comprises a variety of gypsum lithofacies whose origin is still debated (see Ortì, 2011): (a) the massive and banded selenite, consisting of vertically-oriented palisades of bottom-nucleated selenite crystals (arrow head or swallow-tail twins, Shearman and Ortì Cabo, 1976); (b) the branching selenite, defined by Lugli et al. (2010) as consisting of "clear selenite crystals a few centimeters across with their long axis inclined or oriented horizontally grouped into decimeter-large irregular nodules and lenses separated by thin fine-grained carbonate or gypsum laminae"; and (c) the laminar gypsum,

formerly termed "balatino" (Ogniben, 1957), either representing a in situ deposit of sub-mm gypsum crystals or a fine-grained clastic deposit emplaced by low density gravity flows (Manzi et al., 2005). Across the different Mediterranean subbasins in southern Spain (i.e. Sorbas and Nijar basins), northern Italy (i.e. Piedmont and Vena del Gesso basins) and southern Italy (Sicily), these gypsum lithofacies show a similar stratigraphic stacking pattern (Lugli et al., 2010): the lower five PLG cycles are composed only of the massive and banded selenite lithofacies, whereas from the 6th cycle upwards the branching selenite is also found. In the gypsum layers of the Vena del Gesso, the branching selective is developed at the top of the massive and banded selenite (Lugli et al., 2010), where as 12 Piedmont forms entire beds and is strictly interlayered with laminar gypsum (Dela Pierre et al., 2011). The PLG unit passes laterally, in the intermediate depths and deep 'asia's, to organic-rich shales (Manzi et al., 2007), but the details of such a transition a e . 'il' poorly known. Still uncertain, moreover, are the depositional mechanisms and the maronmental conditions at the origin of the laminar gypsum and the branching selenite lithe facies, especially in light of the apparent synchronous formation of the branching selenite in different Mediterranean marginal subbasins. This paper integrates previous "trangraphic studies (Dela Pierre et al., 2011) with novel sedimentological, petrographic and geochemical data on the different gypsum lithofacies (massive selenite, branching selenite and laminar gypsum) composing the PLG unit along a margin-to-depocenter transect of a Messinian marginal subbasin located at the northernmost offshoot of the Mediterranean (Piedmont Basin, NW Italy), with the main aims to: (a) provide a detailed description of the gypsum lithofacies, in particular of the still poorly known laminar gypsum and branching selenite; (b) shed light on the paleoenvironmental conditions under which the different lithofacies formed and reconstruct the mechanisms behind the appearance of the laminar gypsum and the branching selenite in the 6th PLG cycle; and (c) discuss the lateral lithofacies transition between the basin margin and the depocenter.

2. Geological setting

During the Messinian the Piedmont Basin (northern Italy) (Fig. 1C) corresponded to a wide wedge-top basin, developed upon an orogenic wedge of Alpine, Ligurian and Adria basement units (Rossi et al., 2009; Mosca et al., 2010; Rossi, 2017). The shallow marginal zones are presently exposed in the southern (Langhe) and northern (Monferrato/Torino Hill) sectors, the latter corresponding to the uplifted sill interposed between the Piedmont Basin and the NW termination of the larger Apennine foredeep (present-day Poplain). The depocenter of the basin is buried below Pliocene and Quaternary deposit, but is well imaged by seismic data (Ghielmi et al., 2019). The Messinian succession (Fig. 1D) starts with outer shelf to slope fine-grained sediments (Sant'Agata Fossili Marls, Fortonian-lower Messinian) that record progressive restriction of the basin before the onset of the MSC (Sturani, 1973; Sturani and Sampò, 1973; Violanti et al., 2013). At 'ne basin margins this unit is overlain by in situ sulfate evaporities of the PLG unit, representing the first stage of the MSC (5.97-5.60 Ma; Dela Pierre et al., 2011). The PLG unit exhibits a lithological cyclicity interpreted to reflect precession-driven climate oscinations (Fig. 2), as elsewhere in the Mediterranean. Towards the basin depocenter, the ovolun layers make a lateral transition to shales, marls and carbonate-rich layers (Fig. 1D), representing deeper water equivalents of gypsum (Dela Pierre et al., 2012). The PLG unit and its lateral equivalents are overlain by resedimented and chaotic gypsum deposits (Valle Versa Chaotic Complex; Irace et al., 2005; Castellania Chaotic complex; Ghielmi et al., 2019), equivalent to the Resedimented Lower Gypsum unit deposited in many Mediterranean subbasins during the second stage of the MSC (5.60-5.55 Ma; Roveri et al., 2014), (Fig. 1D). In the Piedmont Basin the Messinian succession is closed by fluvio-deltaic and lacustrine deposits (Cassano Spinola Conglomerates), recording the third MSC stage (5.55-5.33 Ma; Fig. 1C). Deposition of Zanclean marine clays and marls

(Argille Azzurre Formation) marks the end of the salinity crisis at 5.33 Ma (Trenkwalder et al., 2008; Violanti et al., 2011).

2.1 The PLG unit in the Piedmont basin

Three subunits (A to C from bottom to the top) were distinguished in the Piedmont Basin according to the prevailing gypsum lithofacies (Dela Pierre et al., 2011) (Fig. 3). In all the subunits, the gypsum layers are interbedded with shales (not described in this paper), forming shale/gypsum couplets which are assumed to correspond to precessional cycles. Subunit A is composed of vertically oriented selenite crystals and includes up to five gypsum beds belonging to cycles PLG1 to 5. The gypsum beds are up to 10 m thick. Their number and thickness decreases from SW to NE, i.e., from the be in margin to the depocenter (Fig. 3). Subunit B includes a single gypsum beds (7-10 n. th.ck) belonging to cycle PLG6 and known as Sturani key-bed (SKB) (Dela Pierre et al., 2011). The SKB represents a distinct marker bed that can be physically correlated from the margin to the distal part of the basin. (Fig. 3). The SKB is composed of two lithefac. That are not observed in the underlying subunit A: laminar gypsum and branching elenite (the latter formerly described as nodular gypsum and "embrichitic" structures) (Sturmi, 1973). In the more distal sector of the basin (Govone section, the SKB passes '9 snales. Subunit C overlies the SKB and comprises up to 9 gypsum beds, about 1 m thick, composed of gypsiferous mudstones (Fig. 3). Subunit C is not present at the basin margins, since it is truncated at the top by an erosional surface (Fig. 3) corresponding to the Messinian erosional surface (Dela Pierre et. al., 2011. In the more distal part of the basin (Govone section) subunit C is the only unit recognizable and directly overlies the Sant'Agata Fossili Marls. In this section, the upper part of the Sant'Agata Fossili Marls is time equivalent of subunits A (cycles PLG1-5) and B (cycle PLG6) (Dela Pierre et al., 2016; Gennari et al., 2020) (Fig. 3).

3. Materials and methods

Field studies of the lithological and sedimentological features of gypsum lithofacies were conducted along a SW-NE transect, representing the transition from the basin margin to the depocenter of the Piedmont Basin (Dela Pierre et al. 2011). The following sections were revisited (Fig. 1C, D): Arnulfi (43°37'06''N; 7°53'41''E), Pollenzo (44°41'08''N; 7°55'33''E) and Govone (44°48'09''N; 8°07'33''E); the Monticello (44°42'39''N; 7°57'39''E) and Castagnito (44°45'44''N; 8°03'07''E) sections were studied for the first time. More detailed observations were done in the Pollenz as a tuon, where the PLG unit is exposed entirely. In the other sections the outcrops are frequently covered by vegetation or the presence of unstable cliffs corresponding to ancient quarry fronts, 10 m high and 20 m wide (Arnulfi) makes field operations dangered as.

A total of 54 oriented samples (15 from Arr Ilfi, 30 from Pollenzo, 4 from Castagnito and 5 from Govone) were taken from the various gypsum lithofacies for petrographic observations. Thirty polished slabs, 28 petrographic observations and 30 thin slices (~1 mm thick) produced by cleaving the selented crystals along the main (010) cleavage surfaces with a razor blade, were obtained from the prost representative samples and studied with transmitted, reflected, and UV-visib. Inght microscopy, using a Leika DM 2700 P microscope with excitation filters 340-380 Nm. Eight polished carbon-coated thin sections were studied for semi-quantitative elemental analyses and backscattered electron imagery (BSEI) using a JSM-IT300LV scanning electron microscope (SEM) equipped with an energy-dispersive EDS Oxford Instruments Link Systems (Department of Earth Sciences, University of Torino). Twenty-five samples of the branching selenite, laminar gypsum and gypsiferous mudstone lithofacies were studied by SEM for morphological analyses. Four samples were also analysed with a FESEM Tescan S9000G microscope at the Centre for Nanostructured

Interfaces and Surfaces of the University of Torino. X-Ray Powder Diffraction (XRPD) analyses were performed on 9 samples of laminar gypsum and gypsiferous mudstones to test for the presence of dolomite, using a Rigaku MiniFlex 600 benchtop X-ray diffractometer. A semi-quantitative analysis was performed applying the RIR (Reference Intensity Ratio) method based on the ratio of the intensities of the phases recognized into the sample with respect to the intensity of a "virtual corundum" (I/Ic). An aliquot of the same powder samples prepared for XRPD analyses was selected to assess the isotopic composition of dolomite. Carbon and Oxygen stable isotope analyses were performed at the Department of Earth Sciences, University of Milan, using an automated carbon are preparation device (Gasbench II) and a Thermo Fisher Scientific Delta V Advantage continuous flow mass spectrometer. The isotope compositions are expressed in the conventional delta notation calibrated to the Vienna Pee-Dee Belemnite (V-PDB) scale. The Yorkal Organic Carbon (TOC) contents of 18 samples representing the laminar gypsun, b anching selenite and gypsiferous mudstones lithofacies from the Pollenzo section were determined at the Institute for Geology of the University of Hamburg, using a LFCC SC-144DR Carbon Analyzer equipped with an infrared detector.

4. Results

- 4.1 Field observations
- 4.1.1 Subunit A (Cycles PLG1-5)

This subunit reaches the maximum thickness (30 m) at the basin margin (Arnulfi section, Fig. 3), The lower three beds are made up of the massive selenite lithofacies, consisting of dm-sized bottom-grown selenite crystals (Fig. 4A, B); with a uniform size throughout the bed. The 4th and 5th beds are composed of the banded selenite lithofacies, i.e cm-thick palisades of twinned crystals separated by mm-thick clay layers (Dela Pierre et al., 2011) (Fig. 4C). In the

more distal Pollenzo section, only two layers of massive selenite, up to 1.5 m thick (Figs. 3, 4A), are present, overlying a cyclic succession of shales and marls belonging to the Sant'Agata Fossili Marls (Fig. 3). Previous studies (Lozar et al. 2018) showed that these thin gypsum beds correspond to the 4th and the 5th PLG beds (banded selenite) of the Arnulfi section and that the underlying shale/marl couplets of the Sant'Agata Fossili Marls are the "deep" water equivalents of the lower three marginal PLG cycles. Basinwards (Fig. 3), subunit A is represented by one (visible) bed of massive selenite (Monticello section) and by isolated cones of gypsum growing in dolomite-rich laminated "lty clays (Castagnito section; Fig. 4D). In the more distal Govone section, subunit A passes 'aterally to dolomite-rich marls (Sabino et al., 2021).

4.1.2 Subunit B (Cycle PLG 6)

Subunit B corresponds to the SKB. The Cic'tness of this layer decreases from 10 m at the basin margin (Arnulfi section) to about 7 m in the more distal part of the basin (Castagnito section) section. At Govone the SVB Control present (Fig. 3). The SKB is composed of the laminar gypsum and branching relenite lithofacies (Fig. 5). Laminar gypsum consists of alternating gypsum-rich (1-2 cont thick) and clay-rich (100-200 m thick) laminae (Figs. 5C, 6A-D), locally organized in dm-thick packets separated by cm-thick clay-rich layers (Fig. 6C). At the outcrop scale the laminae can be followed laterally for tens of metres. No sedimentary structures related to tractive bottom currents (e.g., cross lamination) or oscillatory flows (e.g., hummocky cross stratification) have been observed. In the laminar gypsum, plant remains are abundant and insect remains (dragonfly larvae) have been documented (Sturani, 1973).

The branching selenite lithofacies consists of m-scale flattened cones (Fig. 7A-C) growing in the laminar gypsum. In section perpendicular to bedding, the larger cones appear

asymmetrical, with one side (branch) more developed than the other (Fig. 7B). Branches up to 8 m long and 1.5 m high have been observed in the studied sections (Figs. 5A, 7B, 8A). These large branches form an angle of ~10° with respect to the bedding plane (Fig. 7B). In plain view, the cones show a circular to elliptical shape (Fig. 7D). The cones are composed of clusters of dm-sized nodular aggregates (Figs. 5C, 6A), in turn formed by mm-to cm sized prismatic and twinned gypsum crystals with their long axis inclined or oriented horizontally (Fig. 7E, F). The nodular aggregates are here referred to as the "elemental units" of the branching selenite lithofacies. Within the nodular aggregates, the Crystals show a radial distribution (Fig. 7F). The excellent outcrop conditions of the SkB in Pollenzo, where it can be observed on cuts both parallel and perpendicular to the heading, allowed the reconstruction of the stratigraphic relationships between the branching selenite and the laminar gypsum (Fig. 8A). In most cases, laminar are disrupted and deformed around the nodular aggregates ("elemental units") (1 to 8B). However, thin lamina packets can also be traced within the branches or drape the spward termination of larger cones (Fig. 8C).

4.1.3 Subunit C (Cycles PLG 7 15)

The gypsum layers of submin C are up to 3 m thick and shows gradational lower and upper boundaries with the interbeaded mudstones. They are composed of gypsiferous mudstones consisting of the alternation of gypsum-poor and gypsum-rich laminae (Fig. 9A). These sediments differ from the laminar gypsum for the higher terrigenous content. The boundary between the types of laminae is transitional, through the gradual increase in the content of gypsum crystals. The gyspiferous mudstones enclose dm-sized conical structures (Fig. 9A, B), with horizontally oriented crystals (Fig. 9C). These features, up to 10 cm high and 70 dm large, are analogous to the cones observed in the SKB but never reach the size of the latter.

4.2. Gypsum microfacies

4.2.2: Vertically-oriented selenite crystals (Subunit A)

In both the massive and banded selenite lithofacies, the re-entrant angle of the twins is typified by a mm-thick lamination given by the alternation of dark turbid laminae, rich in solid inclusions, and clear lighter laminae in which the latter are scarce or absent (Fig. 10A, B). Solid inclusions are represented by fecal pellets, autofluorescent clay-rich aggregates representing marine snow floccules (Fig. 10C), and filamentous microfossils interpreted as remains of colorless sulfide-oxidizing bacteria (Dela Pierre et al. 2015; Natalicchio et al., 2021). Abundant and well-preserved remains of marine planka'c diatoms, including the nanosized *Thalassiosira* cf. *oceanica* and rarer reworked beauth's diatoms, were observed in the lower two cycles (Pellegrino et al., 2021).

4.2.2 Laminar gypsum and branching seconice (Subunit B)

In the laminar gypsum the alternation of expsum-rich- and clay-rich laminae observed in outcrop is clearly recognizable under the microscope (Fig. 11A, B). The gypsum-rich laminae are composed of a mosaic of interlocked rhombohedral to acicular crystals ranging in size from a few µm to some hund. We so f µm (Fig. 11C). The crystals are limpid, devoid of solid inclusions and are randomy distributed on the lamina surface (Fig. 11D). The clay-rich laminae show a bright autofluorescence when exposed to UV-light (Fig. 11E), which indicates high contents of organic matter. Other common components are silt-sized terrigenous grains (mica flakes, quartz and feldspar) and tiny (2-20 µm) pyrite framboids (Fig. 11F). The clay-rich laminae contain spheroidal dolomite microcrystals ranging in size from 1 to 5 µm, often typified by a hollow core (Fig. 11G, H). Quantification of the dolomite contents of this lithofacies from XPRD data was not possible (Table 1) because of the overlapping of dolomite and gypsum main diffraction peaks.

When the branching selenite lithofacies is observed on cuts perpendicular to the bedding the "elemental units" of this lithofacies appear as transparent nodular-like gypsum aggregates (Fig. 12A). These aggregates are separated by sub-mm thick layers of autofluorescent clayrich sediment including spheroidal dolomite microcrystals (Fig. 12B). The nodular aggregates are composed of mm- to cm-sized densely interlocked prismatic and twinned crystals that give rise to a sutured mosaic (Fig. 12A, C). On cuts parallel to the bedding, the crystals show a unidirectional orientation (Fig. 12C). In the aggregates grouped to form larger asymmetrical cones, the long axis of the crystals is preferentially oriented parallel to the elongation of the branches (Fig. 12D). The gypsum crystals show a concentric 2 ning marked by a fluorescent core surrounded by a not-fluorescent rim (Fig. 12E, F) The core is rich in solid inclusions, including fluorescent loose peloidal sediment, individual pellets (Fig. 12F), gypsum crystals identical to those observed in the laminar gyps an. 1: nofacies (Fig. 12G), terrigenous grains and dolomite microcrystals, i.e., the san, components of the clay-rich laminae in the laminar gypsum. Strings of fluorescent fine-grained sediment are observed between the nodular aggregates (Fig. 12A, B). This sediment appears to have been deformed by the displacive growth of the nodular aggregates.

4.2.3 Gypsiferous mudsines (Subunit C)

The gypsiferous mudstones of subunit C consist of alternation of gypsum-poor and gypsum-rich laminae. The gypsum-rich laminae are 1.5 to 2 mm thick and are composed of equant to prismatic limpid crystals, already described in laminar gypsum, mixed with clay and silt-sized terrigenous grains (mica, quartz, feldspars) (Fig. 13A). The gypsum-poor laminae, 150-200 m thick, show a bright autofluorescence when exposed to UV-light and are typified by high contents of spheroidal dolomite microcrystals (up to 30 wt%, Table 1). The gypsiferous mudstones show evidences of modification of the original depositional fabric; two end

members can be distinguished: (a) the original lamination of the sediment is still recognizable, but laminae are deformed by the overgrowth of gypsum crystals (Figs 13A). These crystals show prismatic habit, are isolated in the sediment and range in size from 1.5 to 3 mm (Fig. 13A). Larger crystals are found along the lamina boundaries; and (b) the original fabric has been obliterated and the bed is composed of mm-scale interlocked crystals forming cm-scale nodular aggregates (Fig. 13B) enclosed in an autofluorescent and dolomite-rich fine-grained matrix. As described for the branching selenite lithofacies, aggregates can be grouped into larger, dm-sized cones (Fig. 9).

4.3. TOC contents and bulk rock C and O stable isotopo composition

The TOC contents of gypsum beds of subunit B (branching selenite and laminar gypsum) and C (gypsiferous mudstones) from the Pollenzo ecolor ranges from 0.1 to 2 wt% (Table 1). Laminar gypsum and gypsiferous mudstones show the highest TOC contents (ranging from 1 to 2 wt%), with small differences between the two subunits. The ¹³C values of dolomite (¹³C_{dol}) ranges from –4.2 to +0.1% FDB, whereas the ¹⁸O_{dol} values range from –5.0 to +4.2 % PDB (Table 1).

5. Discussion

- 5.1 Modes of gypsum deposition
- 5.1.1 Vertically-oriented selenite crystals (subunit A)

The vertically-oriented selenite crystals were deposited on a bottom overlain by calcium and sulfate saturated brines (Fig. 14A). The large and constant size of the selenite crystals that typifies the massive selenite lithofacies suggests a relatively low degree of supersaturation of the brines, allowing the crystals to grow larger (Ortì, 2011). In contrast the smaller crystals in the banded selenite lithofacies (found at the basin margin in the 4th and 5th PLG cycle) reflect

an increase in supersaturation of bottom brines, enhancing the nucleation of crystals and therefore more competition for space among the nuclei (Lugli et al., 2010). The lamination in the re-entrant angle of the twins observed in both massive and banded selenite lithofacies (Fig. 10) possibly reflects short-term (annual? seasonal?) climate oscillations causing cyclic changes of gypsum saturation state in the water mass intercepting the sea floor. During phases of saturation, gypsum growth incorporated sediment and biogenic material accumulated on the top of the crystals during the previous unsaturated phase (turbid laminae) and generated the inclusion-free overgrowths (clear laminae). Gyosum deposition occurred when the sea bottom was oxygenated (Garcia Veigas et al. 2018); oxygenation hampered the activity of sulfate-reducing bacteria able to promote gynsum undersaturation through sulfate consumption (e.g., Babel, 2007; De Lange and Krijg ma. 2010). No clear depth indications are provided by these facies, due to the absence of modern analogues. The assignment of the filamentous fossils to colorless sulfide-o. idizing bacteria like *Beggiatoa* and *Thioploca* (Natalicchio et al., 2021) do not provide any depth constraint, since these prokaryotes can live in a wide range of water depth from bathyal to peritidal settings (e.g. Bailey et al., 2009). In the selenite crystals of cycles PLG1 and 2, the massive occurrence of planktic nanosized diatoms (*Thalassiosira* oceanica) nowadays thriving in coastal to open ocean waters agrees with the notion unit pottom-grown gypsum does not necessarily reflect shallow depositional conditions (Pellegrino et al., 2021).

5.1.2 Laminar gypsum and large branching selenite cones (subunit B)

Laminar gypsum, has been described from several Messinian (Hardie and Eugster 1971; Schreiber et al., 1977; Marsaglia and Tribble, 1999; Lugli et al., 2015) and Badenian (Bąbel, 1999) basins as well as from Miocene lacustrine deposits from Spain (Ortì et al., 2014). The laminar gypsum lithofacies was interpreted either as a primary in situ precipitate, or as a

detrital deposit emplaced by low density turbidity currents (Schreiber and El Tabakh, 2000). The latter interpretation is based on the occurrence of soft sediment deformation, bottom current structures (i.e., cross lamination), normal grading, smoothed and rounded crystal boundaries or broken crystals (e.g., Hardie and Eugster, 1971; Babel, 1999; Schreiber and El Tabakh, 2000). In the studied case, features indicative of a detrital origin are totally absent. On the contrary, the following evidence indicates that laminar gypsum represents an in situ cumulate deposit, resulting from gypsum nucleation at the top or within the water column and subsequent deposition on the seafloor (Hardie and Lowenstein 2004; Lugli et al., 2015): (a) the fine lamination and lateral continuity of the gypsum lamin e and their cyclic alternation with fine-grained settling material (clay, silt, organic matter) (Fig 5), suggesting that the seafloor was periodically draped by a "pelagic" rain of covstals (Fig. 14B); and (b) the absence of solid inclusions in the crystals, that ag acs with nucleation in the water column. The varved aspect of this lithofacies suggests the influence of short-term climate oscillations from more arid (gypsum-rich laminae) romore humid (clay-rich laminae) conditions. Every lamina can be the product of a distinct seasonal sedimentary episode, as documented in modern salinas (e.g., Ortì et al., 1984) or inferred from Messinian gypsum cumulate deposits (Manzi et al., 2012).

The nucleation of tiny gosum crystals at the top of the water column or within the latter requires high supersaturation conditions favouring the formation of new nuclei, inhibiting them from growing larger (i.e., homogeneous nucleation; Alimi et al., 2003; Alimi and Gadri, 2004; Reiss et al., 2017). High supersaturation may be favoured by: (a) strong seasonal evaporation, which also increases the salinity of surface waters (e.g., Babel and Schreiber, 2014) and promotes the nucleation of gypsum at the air-water interface (e.g., Schreiber and El Tabakh, 2000); or (b) mixing of water masses of different chemical composition and density in stratified basins, which can promote gypsum precipitation in the water column

even from a brine that was undersaturated with respect to gypsum before mixing ("salting out" effect; Raup, 1982; Babel and Bogucki, 2007). Mixing occurs at the pycnocline (Raup, 1982) which commonly coincides with an oxycline separating an oxygenated upper water layer from an anoxic, denser lower layer, as observed in both modern (Wakehamn et al., 2007) and ancient stratified basin (e.g., Sabino et al., 2021, and reference therein). An additional mechanism known to increase sulfate concentration and gypsum precipitation is the oxidation of sulfide accumulated in organic-rich bottom sediments through bacterial sulfate reduction and diffused upward in the water column until it maches oxygenated waters above the oxycline (e.g., Tiffany et al., 2007). This process was described in the Salton Sea, a Californian hypersaline lake where gypsum crystals form periodically in the water column (Ma et al., 2020).

Nucleation at the air-water interface and the preservation of settling gypsum at the bottom would imply that the entire water colum. was saturated; such conditions are consistent with a very shallow basin (i.e., few meters), but it is rather unlikely for the studied deposits since:

(a) the absence of physical and paleot congical evidence of very shallow water conditions, such as desiccation features, statetures related to tractive bottom currents or bird footprints. These features were instead described in very shallow water laminar gypsum deposits from other Messinian Medite, ranean subbasins, such as the "balatino" gypsum from Sicily (Hardie and Eugster 1971) and the Marmara gypsum from Cyprus (Manzi et al., 2016); and (b) the high lateral continuity of the SKB, pointing to uniform depositional conditions across the basin, in line with a relatively deep basin, although a reliable depth estimate is not possible. Therefore, it is reasonable to assume that gypsum nucleated within the water column, at the oxycline of a stratified basin, through brine mixing (Raup, 1982), reoxidation of sulfide, or a combination of the two.

Deposition of laminar gypsum from settling crystals mixed with fine-grained sediment was associated with the formation of the branching selenite. This lithofacies was considered as a bottom grown structure (Lugli et al., 2010) comparable with the subaqueous selenite supercones described in the Sorbas basin (Spain) by Dronkert (1985). The horizontal crystal arrangement was explained with a depressed pycnocline (close to the sea floor), hampering the vertical growth of the crystals (Lugli et al., 2010). The same interpretation was proposed for the branching selenite of the Piedmont Basin studied here (Dela Pierre et al., 2011). However, revisitation of this lithofacies through the field and potrographic analyses presented here reveals that: (a) the crystals of the branching selenite include abundant solid inclusions, which implies their initial poikilotopical growth within the sediments; (b) the crystals commonly show a fluorescent core and a limpid non fluorescent rim devoid of solid inclusion (Fig. 12F), which suggests subsequent overgre with in the sediments, to form nodular aggregates of interlocked crystals representing the "elemental units" of the branching lithofacies; and (c) at the contact with the nodular aggregates of the branching selenite the laminae of laminar gypsum are bended or even disrupted, suggesting that the crystals continued to grow displacively annibalizing part of the laminar gypsum crystals and pushing away the enclosing laminated addiment that is now squeezed between the nodular aggregates. These observations suggest that the "elemental units" composing the branching selenite lithofacies actually grew below the sediment water interface, within a sediment consisting of a mixture of "pelagic" gypsum crystals, clay and organic matter, i.e, the laminar gypsum (). The fact that locally laminar gypsum and clay-rich layers drape the larger branches (Fig. 8C), suggests that the topmost part of the cones locally reached the seafloor, forming low-relief positive structures draped by particularly intense episodes of accumulation of pelagic rain of gypsum. The asymmetric growth of these structures was explained with the presence of flowing brines causing the preferential growth of the crystals upstream (Babel and Bogucki,

2007; Lugli et al., 2010). The absence of tractive current structures in laminar gypsum suggests, however, that brines were not able to move the loose sediment deposited at the sea bottom.

Abundant dolomite microcrystals were observed in the fluorescent laminae of laminar gypsum and around the nodular aggregates in the branching selenite lithofacies (Fig. 11G, H). The spheroidal habit and hollow core of the dolomite microcrystals and the slightly ¹³C depleted isotope composition of the sediment (¹³C values –2.4‰ in average) agree with a microbial origin of dolomite (Fontes et al. 1987; Warthmann et al., 2000; Van Lith et al., 2003; Lindtke et al., 2011). The presence of organic matte in he laminar gypsum may suggest that dolomite precipitation was induced by organoclastic bacterial sulfate reduction (Petrash et al., 2017). This process increases alkalinity, and favours the precipitation of ¹³C depleted carbonate minerals, including dolomite 125., Vasconcelos et al., 1995; Sanz-Montero et al., 2009; Bontognali et al., 2017). In addition, such process releases hydrogen sulfide that, reacting with iron, can sink so pyrite (that is actually abundant in the laminar gypsum, see Fig. 11F). Since sulfate reducing bacteria are prevalently anaerobic heterotrophs (e.g., Londry and Des Marais, 2003), it is very likely that the branching selenite grew in anoxic pore waters and that accosition of laminar gypsum took place when the sea bottom was anoxic. The formation and preservation of gypsum in such conditions is in contrast with the notion that bacterial sulfate reduction decreases sulfate concentration, ultimately causing gypsum dissolution (e.g., Babel, 2007; De Lange and Krijgsman, 2010). However, gypsum supersaturation can be maintained even in a sulfate reducing environment, if the sulfate sink via microbial reduction is smaller than the supply of sulfate from other sources (e.g., Torfstein et al., 2005). In the case studied here, calcium and sulfate ions could have been continuously shuttled to bottom waters by the settling of solid gypsum microcrystals nucleated in the water column (e.g., Torfstein et al., 2005). These crystals were initially

susceptible to dissolution in the unsaturated lower water column and seafloor. However, when the rate of gypsum nucleation was high, sulfate consumption by microbial sulfate reduction below the oxycline was outweighted by sulfate replenishment through dissolution of solid gypsum. In these conditions, saturation at the sea bottom was immediately attained and maintained, allowing accumulation of tiny crystals forming the gypsum lamina (Fig. 14B, step 1). In contrast, when nucleation of gypsum in the water column was limited or absent and the supply of organic matter enhanced, sulfate removal was not balanced by sulfate replenishment by the pelagic gypsum rain. These conditions, that coincide with the formation of the clay (and dolomite)-rich lamina, favoured the momentary undersaturation of the lower water column and at the seafloor, with consequent dissolution of the rare gypsum crystals settling down from the oxycline and of the crysuls deposited during the previous phase (Fig. 14B, step 2). When the pelagic gyr sum ain resumed, saturation at the sea bottom was again attained and calcium and sulfa in previously provided by dissolved crystals to interstitial brines could precipitate, proporting the syngenetic growth and further enlargement of the crystals observed in the nodular aggregates of the branching selenite lithofacies (Fig. 14B, step 3). Repetition of this mechanism through time drove the formation of clusters of aggregates, that could locally reach the seafloor and be sealed by pelagic gypsum during intervals of enhanced nucleation in the water column. On the contrary, in the distal deeper portions of the basin, reduced shuttling of gypsum microcrystals and enhanced bacterial sulfate reduction hampered gypsum preservation (laminar gypsum) and nucleation in the shallow subsurface (branching selenite; Fig. 14B).

In conclusion, we propose that the branching selenite lithofacies is a syngenetic product formed within the sediment, at the expenses of gypsum crystals nucleated in the water column and deposited at the seafloor. These crystals acted as shuttles of calcium and sulfate ions that fostered gypsum deposition in reducing pore waters.

5.1.3 Gypsiferous mudstones and small branching selenite cones (subunit C)

Gypsiferous mudstones share many characteristics with laminar gypsum (lamination, lateral continuity of the laminae, absence of evidence of bottom currents) and are interpreted as cumulate deposits. However, in the gypsiferous laminae the pelagic gypsum crystals are mixed with clay and organic matter. Since the TOC contents are the same as in the SKB (~ 1.5% in average) (Table1), mixing is most likely due to reduced nucleation of gypsum in the water column and enhanced input of clay in the basin, rather them to increased flux of organic matter (Fig. 14C). As described for subunit B, undersatura ion of the lower water column and at the seafloor following unbalanced bacterial sulfate reduction, drove dissolution of pelagic gypsum. The subsequent saturation of interstitial brines promoted the syngenetic growth of gypsum crystals in bottom sediments, deforming he lamination and eventually forming nodular aggregates of interlocked crystal. ("clemental units") that were grouped together to form the observed cones. Interestingly, evpsum beds of subunit C are thinner than the SKB and large cones are not observed. Such a change reflects most likely the reduced availability of gypsum settling down through the water column, whose dissolution provided calcium and sulfate ions for the growth of crystals in the sediment.

5.2 Mechanisms controlling the distribution of gypsum lithofacies

The progressively younger ages of the first gypsum bed from the basin margin to the depocenter defines an overall progradational stacking pattern reflecting shallowing of the basin from the base to the top of the PLG unit and the consequent shift of gypsum deposition basinwards. This shallowing-upward trend is shared with other PLG marginal subbasins (Vena del Gesso; Lugli et al., 2010; Reghizzi et al., 2018; Sorbas; Roveri et al., 2020).

At the onset of the MSC, bottom grown selenitic gypsum (subunit A) was deposited in the shallowest part of the Piedmont basin (Arnulfi section), whereas marls with filamentous microfossils (Dela Pierre et al., 2012) or rich in dolomite (Sabino et al., 2021) formed at intermediate depth and in the deepest parts of the basin, respectively (Pollenzo and Govone sections). Such lateral facies change was interpreted to reflect the stratification of the water column and the position of the oxycline with respect to the seafloor (Sabino et al., 2021). Bottom grown selenitic gypsum (massive selenite lithofacies) formed only above the oxycline, in mixed and periodically oxygenated calcium and surface saturated waters, not affected by bacterial sulfate reduction at times of gypsum precipitation (Fig. 14A). In the specific case where the oxycline intercepted the sea botton, suboxic conditions prevailed favouring, the growth of microbial mats dominated by sulfide-oxidizing bacteria (Natalicchio et al., 2017). In the deeper sector of the basin, the sea bottom was below the oxycline and bottom grown gypsum did not form, sinc ir cense bacterial sulfate reduction (in this case not balanced by "pelagic" gypsum deposition) caused gypsum undersaturation and the formation of authigenic dolomite (Babel and Ro, waki, 2007; De Lange and Krijgsman, 2010; Sabino et al., 2021). The onset of gypsun. deposition in the distal parts of the basin (Pollenzo, Monticello and Castagnito sections) coincides with the vertical facies change from massive (cycle PLG3) to banded resente (cycle PLG4) at the basin margins. Such a change reflects most likely shallowing of the basin, enhancing supersaturation in the shallowest part and the shifting of the oxycline basinwards. Accordingly, gypsum could start to grow on the previously suboxic or anoxic seafloor. The "distal" selenitic gypsum layers are however thinner and discontinuous (at Castagnito) compared to their marginal counterparts (Fig. 3). Such thickness and lateral continuity reduction possibly results from exhaustion of dissolved calcium and sulfate ions in the distal bottom brine, following massive gypsum precipitation at the basin margins.

The appearance of laminar gypsum and the branching selenite lithofacies (Fig. 14B) reflects a significative paleonvironmental and paleohydrological change at about 5.87 Ma (6th PLG cycle). Interestingly, the branching selenite lithofacies apparently formed synchronously in the 6th PLG cycle in sections located several hundreds of km apart (SE Spain, Sicily, Northern Apennines, Piedmont Basin) (Lugli et al., 2010). This event was possibly related to the reduction of the inflows of oceanic waters and/or to increasing riverine runoff in marginal Mediterranean sub-basins (Reghizzi et al., 2018). Deposition of the 6th PLG cycle correlates with an eccentricity minimum (Manzi et al., 2013), associated with a cooling phase and global sea-level fall (Miller et al., 2011; Van der Laan et a'., 2012). These processes, along with ongoing tectonic restriction of the hydrological exchanges with the Atlantic Ocean (e.g., Flecker et al., 2015; Capella et al., 2018) resulted in furn or restriction of the Mediterranean Sea (e.g., Reghizzi et al., 2018) that consequently became more sensitive to the influx of freshwater from river discharge (or even 'ro'n the Paratethys; Grothe et al., 2020). An effect of freshwater inflows was the intensific, tion of water column stratification (e.g., Simon and Meijer, 2017). In the Piedmont basin Calanced stratification was associated with the rise of the oxycline in the water colunn, that led to homogeneous environmental conditions and to deposition of laminar gypsum, and the branching selenite lithofacies, under the control of short term climate fluctuation. Deposition of subunit C (Fig. 14C) reflects most likely continuous restriction and shallowing of the basin, which agrees with the progradational stacking pattern of gypsum layers, that were deposited even in the more distal Govone section. Shallowing and restriction resulted in closer sediment entry points and enhanced sensitivity to riverine runoff, with consequent increase of the input of clay, reduction of gypsum nucleation in the water column and persistent stratification. Stratification is possibly reflected by the partial offset of the ¹⁸O_{dol} values between sediments deposited in more marginal (Pollenzo section, ¹⁸O_{dol} as low as -4.9% PDB) and deeper sectors (Govone

section, ¹⁸O_{dol} up to +4.2% PDB) of the basin; accordingly, ¹⁸O-enriched, denser and likely colder bottom waters occupied the deeper sectors of the basin, whereas ¹⁸O-depleted, more diluted and warmer bottom waters were present in more marginal sectors.

No gypsum, but only time equivalent shales, are reported from the buried basin depocenter on the base of seismic data (Ghielmi et al., 2019). This pattern is possibly explained by reduced supply of gypsum from the water column in more open sea and higher rates of bacteria sulfate reduction, deriving from permanent seafloor anoxia and larger availability of organic matter.

6. Conclusions

The analysis of gypsum deposits exposed in the Pied non. Basin provides new insights on the lateral facies transition between the margin and the deposition of a Messinian marginal subbasin. Bottom-grown gypsum was detected above the oxycline from overlying saturated brines not affected by bacterial sulfate reduction. Cumulate deposits (laminar gypsum and gypsiferous mudstones) originated from gypsum nucleation at the oxycline and subsequent deposition on an anoxic sea bottom. Deposition of these lithofacies under reducing bottom conditions was controlled by the balance between the rate of sulfate reduction (removing sulfate) and the rate of sulfate replenishment through dissolution of settling gypsum crystals. The branching selenite lithofacies is interpreted as a syngenetic product formed in the sediment at the expenses of dissolved cumulate gypsum. The appearance of cumulate deposits and the branching selenite lithofacies in the 6th PLG cycle (i.e., at about 5.87 Ma) records further restriction of the basin and the intensification of water column stratification. This study suggests that fluctuations of the oxycline in the water mass was a key parameter controlling the stratigraphic distribution of gypsum lithofacies in an ancient stratified basin during the MSC.

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FIGURE AND TABLE CAPTIONS

Table 1

Total Organic Carbon (TOC) contents, dolomite contents, $^{13}C_{dol}$ and $^{18}O_{dol}$ values of representative samples.

Fig. 1

(A) Distribution of Messinian evaporites in the Mediterranean basin (modified from Manzi et al., 2013). (B) Chronostratigraphic framework of the Messinian salinity crisis (modified from Manzi et al., 2013). (C) Structuard sketch map of the Piedmont Basin (modified from Bigi et al., 1990) with location of the Gratigraphic sections shown in Fig. 3 (numbers): (1) Arnulfi; (2) Pollenzo; (3) Monticardo; (4) Castagnito; (5) Govone. TH: Torino Hill; MO: Monferrato; gray dotted lines indicate the isobaths of the base of Pliocene deposits. The black dotted line is the trace of the cross section shown in panel D. (D) Schematic cross section showing the distribution of the Messinian sediments (from Dela Pierre et al., 2011), with locations of the stratigraphic sections mentioned in the text (numbers). The trace of the cross section is indicated in panel C. M/P: Miocene-Pliocene boundary. PLG: Primary Lower Gypsum unit; RLG: Resedimented Lower Gypsum unit; UG: Upper Gypsum; MES: Messinian erosional surface.

Fig. 2

(A) Aerial view of the Pollenzo section, showing the lithological cycles of the PLG unit.

More competent layers correspond to gypsum, softer layers to shales. The white rectangle indicates the position of panel B; (B) Detail of A.

Fig. 3

Correlation scheme of the sections cited in the text (see Fig. 1C for their location). Datum plane: base of the Sturani key-bed. SAF: Sant'Agata Fossi'i Marls; PLG: Primary Lower Gypsum unit; MES: Messinian erosional surface. PLG1-FIG 15: Primary Lower Gypsum cycle.

Fig.4

Subunit A. (A) The gypsum layer of cycle PLG5, composed of massive selenite, and the overlying cycle PLG 6 with the Stural i key-bed (Pollenzo section). (B) Dm-sized vertically-oriented twinned selenite crystal from cycle PLG5 (Pollenzo section). (C) Banded selenite lithofacies (cycle PLG4, Arnalif. section); the yellow dotted lines indicate clay layers between crystal palisades. (D) m reale gypsum cone composed of vertically-oriented selenite crystals (cycle PLG5, Castagnito section). The yellow dotted line indicates the base of the cones, the white dotted lines indicate bedding of hosting shales.

Fig.5

Sturani key-bed in the Arnulfi (A) and Monticello (B, C) sections. Note in (A) the coalescent nodular aggregates (arrows) growing in laminar gypsum (white lines) and a large asymmetric cone (yellow dotted line); in (C), laminar gypsum (LG) and the branching selenite (BS)

lithofacies are clearly visible. Note that the branching selenite is formed by clusters of nodular aggregates.

Fig. 6

Laminar gypsum (LG) and branching selenite (BS). (A) A m-scale branch, internally composed of smaller nodular aggregates (arrows) above laminar gypsum (Pollenzo section). (B) Close up of the laminar gypsum; note the cm-thick packets separated by clay-rich layers (Pollenzo section). (C) Detail of B, showing a five cm-thick laminated packet, bordered by clay-rich layers (arrows). The gypsum-rich laminae are tv₁ ifier by a whitish to pinkish colour. (D) The laminar gypsum in the Monticello section.

Fig. 7

(A) Plane view of the Sturani key-bea in the Pollenzo section: note the large asymmetrical cone (yellow dotted lines) and the eto ig ition of the more developed arm (branch) towards the left side of the picture (south). B) Fransversal view of the same branch of (A) (yellow dotted line). Note that the laminae in the laminar gypsum (white dotted lines) can be traced within the branch where they are conjugate to the bedding. The sketch on the left side indicates the 3D shape of the cone and the orientation of the thin sections shown in Fig. 12A-C. (C) Metersized conical structures at the base of the Sturani key-bed in the Pollenzo section, observed from below. (D) Aerial view of asymmetrical cones. (E) Detail of (D), showing clusters of nodular aggregates parallel to the elongation of a branch (arrows). (F) Detail of (E), showing mm-sized gypsum crystals with their long axis oriented parallel to the elongation of a branch.

Fig. 8

(A) Stratigraphic log and outcrop view of the Sturani key-bed in the Pollenzo section, showing the stratigraphic relationships between laminar gypsum (LG) and the branching selenite (BS) lithofacies. A large branch cross-cutting laminar gypsum (yellow dotted lines) is visible. The thick black lines in the log indicate clay-rich layers which subdivide the Sturani key-bed in twelve gypsum-rich intervals. (B) Bended and deformed laminae (arrows) around the nodular aggregates. (C) Detail of (A), showing the draping of the laminae on the upward termination of a branch (white dotted line). Note the nodular aggregates composing the branch.

Fig. 9

Gypsiferous mudstones. (A) Outcrop view of cycle PLG? (Pollenzo section). Note the lamination in the gypsiferous mudstones and the small cones they enclose (arrows). (B) A single cone enclosed in gypsiferous mudstones (Cycle PLG15, Govone section). The X, Y and Z axes refer to the orientation of the sections reported in (C). (C) Transversal and planar section of the same cone of (B); arrow indicate the growth direction of the crystals. Note the radial distribution of the crystals.

Fig. 10

Photomicrographs in transmitted (A, B) and UV-light (C) of vertically-oriented twinned selenite crystals (subunit A). (A) Alternation of turbid and limpid laminae in the re-entrant angle of the twin. (B) Turbid lamina (TL) and light lamina (LL) from the same crystal of (A). The red arrows indicate the filamentous microfossils. (C) Fluorescent clay-rich aggregates corresponding to marine snow floccules. Note the tiny pyrite grains in the center.

Fig. 11

Laminar gypsum (subunit B). (A, B) Photomicrographs in transmitted light. Note that thicker gypsum-rich laminae are separated by thin clay-rich laminae. (C) Close-up of a gypsum lamina (polarized light): note the sutured boundaries in two large crystals (arrow). (D, E) Photomicrographs in transmitted (D) and UV (E) light of a cut slightly oblique to lamination. Note in (D) the random orientation of the gypsum crystals and in (E) the high autofluorescence of a clay-rich lamina. Pollenzo section. (F) Backscatter SEM image of the clay-rich matrix between the gypsum crystals (gy). Note the silt-size terrigenous grains (t) and the abundant, bright grains corresponding to pyrite (arrowe). Follenzo section. (G) SEM image of a clay-rich lamina showing clustered dolomite microcrystals with rounded shape and hollow core and some terrigenous grains (t). Monticent section. (H) SEM image of dolomite microcrystals with hollow core.

Fig. 12:

Branching selenite (subunit B). (A) The next itted light photomicrograph of a cut perpendicular to the elongation of the branch represented in Fig. 7B; nodular crystal aggregates separated by thin strings of sediment are recognizable. (B) UV-light photomicrograph showing a detail of (A). Note the zoned crystal typified by a fluorescent core and a non-fluorescent rim and the high fluorescence of the surrounding clay-rich sediment. (C) Plane view of the branch represented in Fig. 7B (transmitted light photomicrograph); note that the crystals (arrows) are disposed parallel to the elongation of the branch; the inset shows a twinned crystal. (D) Transmitted light photomicrograph of a cut parallel to the elongation of the branch (Fig. 7B). The crystals (arrows) are disposed parallel to the elongation of the branch as in (C). Nodular crystal aggregates are recognizable. (E, G) Photomicrographs in transmitted (E) and UV (F) light of interlocked crystals composing the nodular aggregates; note the solid inclusions (peloids) in the core of the crystals (arrows) and in (F) the different fluorescence intensity of

the core (c) and the rim (r). (G) Elongated prismatic primary gypsum crystal (arrow) enclosed in a larger crystal (polarized light). The yellow faces of the rectangle in panels A, C and D indicate the orientation of the thin sections (see also Fig. 7B).

Fig. 13

Gypsiferous mudstones (subunit C). (A) Alternation of gypsum-rich and gypsum-poor laminae. Note the larger size of the crystals compared to laminar gypsum. (B) Nodular aggregates composed of interlocked crystals-

Fig. 14

Schematic representation of the depositional conditions of the three subunits composing the PLG unit. (A) Subunit A (vertically-oriented cryptor15); (B) Subunit B (laminar gypsum and branching selenite; (C) Subunit C (gypsitor as mudstones). The panel below (B) shows the interpreted mode of deposition of laminor gypsum (steps 1 and 2) and the branching selenite lithofacies (step 3). The light blue arrow on the right indicates the influx of Mediterranean (Med) water in the basin. See wet for details.

| Sample | Section | PLG cycle | Lithofacies | TOC (%) | Dolomite (wt%) | $\delta^{13}C_{dol}$ | $\delta^{18}O_{dol}$ |
|----------|----------|------------|--------------------------|------------|----------------|----------------------|----------------------|
| Ar1 | Arnulfi | PLG6 (SKB) | Laminar gypsum | n.a. | - | -2,4 | -4,9 |
| SKB-A2 | Pollenzo | PLG6 (SKB) | Laminar gypsum | 0.97 | 5.3 | -2,4 | -2,9 |
| PDZ5.7 | Pollenzo | PLG8 | Gypsiferous mudstones | 2,03 | 29.9 | -2,3 | +3,9 |
| PDZ6.9 | Pollenzo | PLG9 | Gypsiferous mudstones | 1,03 | 11.2 | -4,2 | +1,9 |
| PDZ7.8 | Pollenzo | PLG10 | Gypsiferous mudstones | 1,50 | 11.8 | -2,9 | -4,2 |
| PDZ8.4 | Pollenzo | PLG11 | Gypsiferous mudstones | 1,18 | 9.9 | +0,1 | -4,9 |
| SG-M7 | Govone | PLG8 | Gypsiferous mudstones | n.a. | 28.7 | -3,0 | +4,2 |
| SG-M10 | Govone | PLG9 | Gypsiferous mudstones | n.a. | 21.7 | -2,3 | +3,8 |
| PDZ.3.10 | Pollenzo | PLG6 (SKB) | Branching selenite | 0,14 | ¬ a. | | |
| PDZ.3.12 | Pollenzo | PLG6 (SKB) | Branching selenite | 0,22 | n.a | | |
| PDZ.3.15 | Pollenzo | PLG6 (SKB) | Branching selenite | 0,13 | 11.d. | | |
| PDZ.3.16 | Pollenzo | PLG6 (SKB) | Branching selenite | 30,0 | n.a. | | |
| PDZ.4.12 | Pollenzo | PLG7 | Branching selenite | 0.17 | n.a. | | |
| PDZ.4.13 | Pollenzo | PLG7 | Branching selenite | ^41 | n.a. | | |
| PDZ.5.8 | Pollenzo | PLG8 | Branching selenite | 0,1) | n.a. | | |
| PDZ.6.10 | Pollenzo | PLG9 | Branching selenite | J. 18 | n.a. | | |
| PDZ.6.8 | Pollenzo | PLG9 | Branching selenita | 0,18 | n.a. | | |
| PDZ.7.8 | Pollenzo | PLG10 | Branching selen 'e | 0,08 | n.a. | | |
| PDZ.7.9 | Pollenzo | PLG10 | Branching sel nite | 0,08 | n.a. | | |
| PDZ.8.5 | Pollenzo | PLG11 | Branchir & S. len. | 0,23 | n.a. | | |
| PDZ.9.5 | Pollenzo | PLG12 | Branch. 2g selenite | 0,39 | n.a. | | |

n.a.: not analyzed

Declaration of interests

| ☑ The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper. | |
|--|--|
| □The authors declare the following financial interests/personal relationships which may be | |

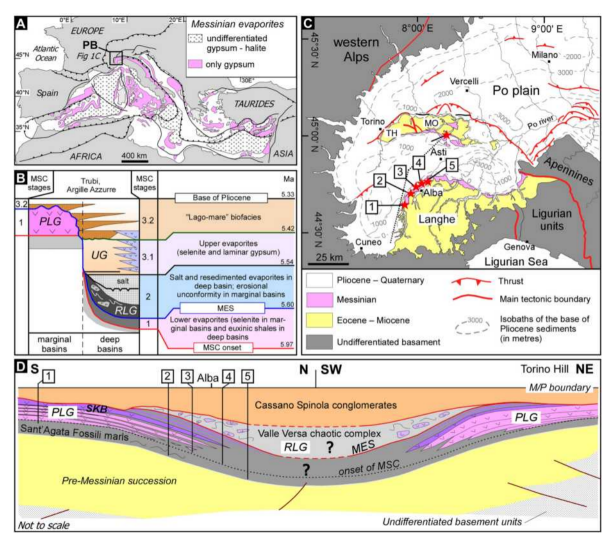


Figure 1



Figure 2

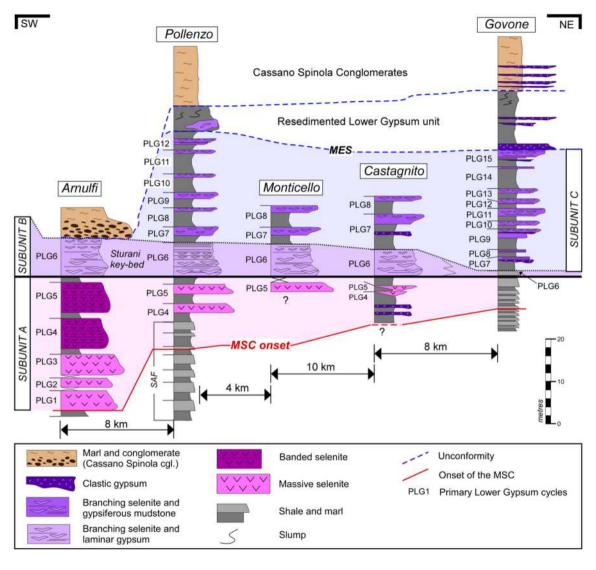


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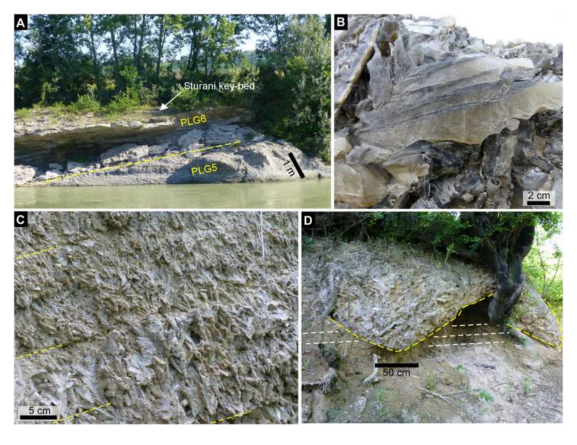


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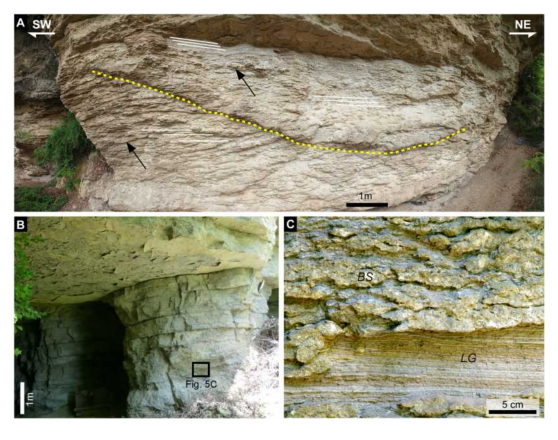


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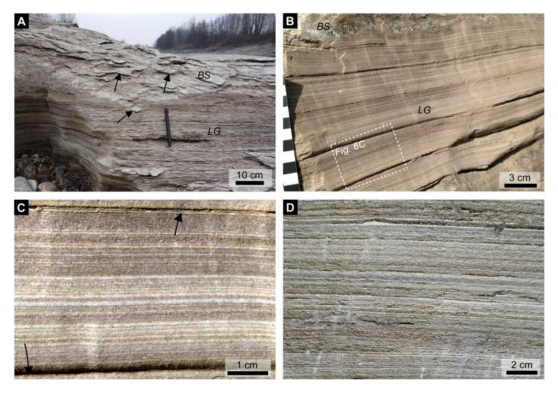


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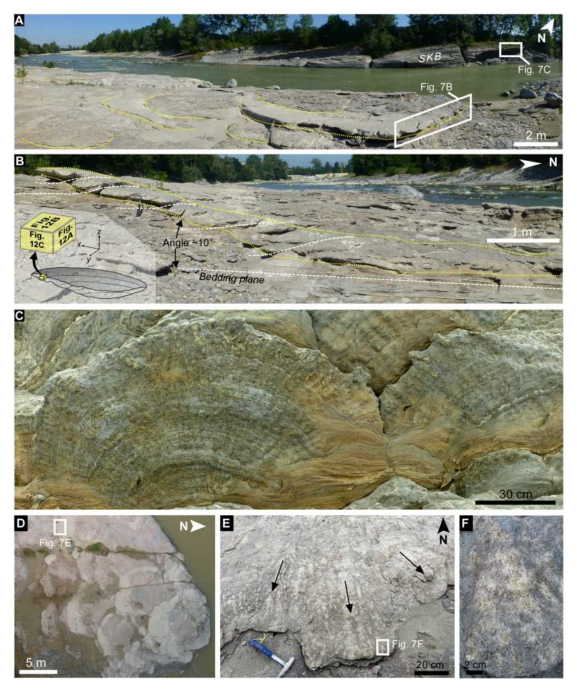


Figure 7

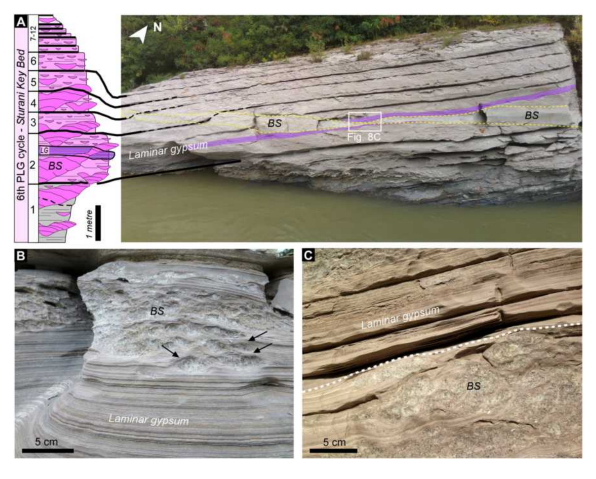


Figure 8

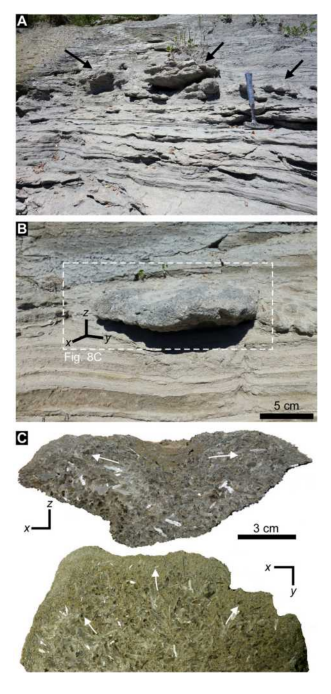


Figure 9

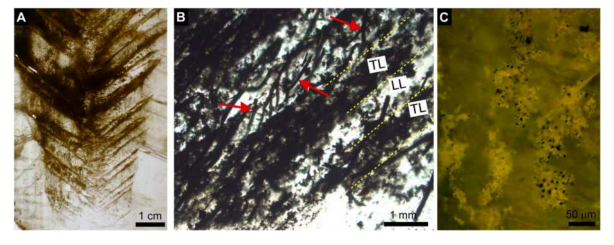


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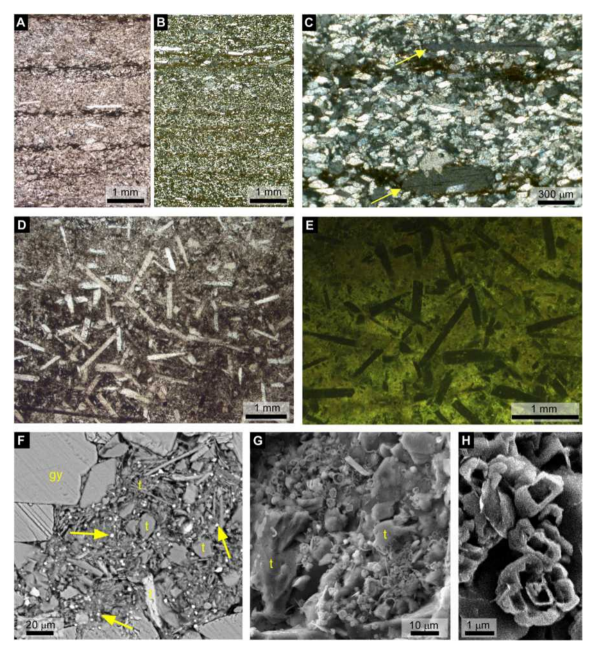


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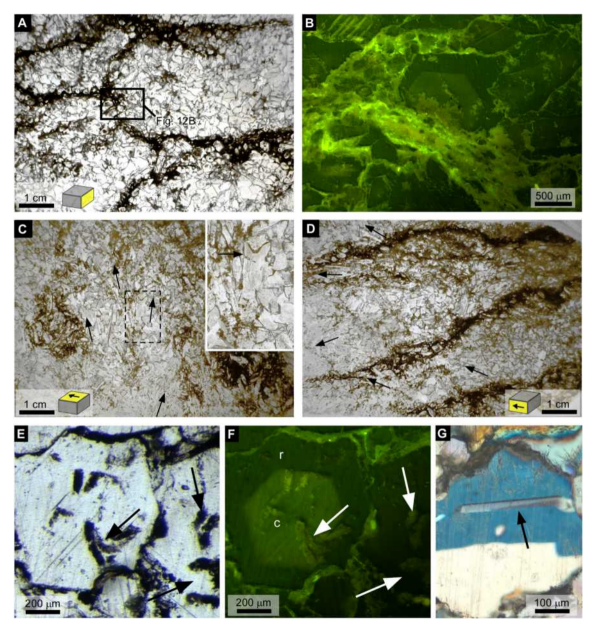


Figure 12

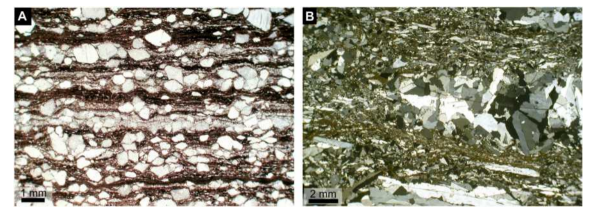


Figure 13

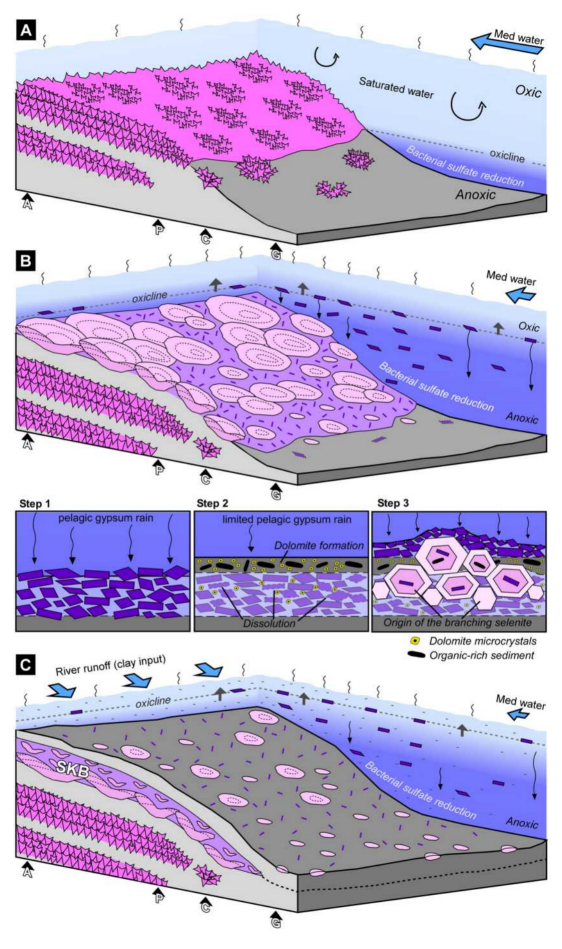


Figure 14