INTRODUCTION

Primary bottom-grown gypsum is an excellent archive of ancient bio-signatures (e.g., Schopf et al., 2012) and represents a promising target for understanding early life on Earth and for exploring the possibility of life on Mars (Allwood et al., 2013; Benison & Karmanocky, 2014). Fast growth of gypsum crystals allows for rapid entombment of organic material, including microalgae and prokaryotes, often the only fossils found in ancient marine evaporites (Dela Pierre et al., 2015; Schopf et al., 2012). A prominent archive is the primary...
gypsum having accumulated in the Mediterranean Basin during the Messinian salinity crisis (MSC; 5.97–5.33 Ma; Roveri et al., 2014), when the Mediterranean was turned into the youngest salt giant of Earth history (e.g., Hsu et al., 1973). The Messinian gypsum deposits are grouped into three stratigraphic units (CIEMS, 2008): the Primary Lower Gypsum unit, representing the first phase of the MSC (5.97–5.60 Ma); the Resedimented Lower Gypsum unit, consisting of large blocks of the Primary Lower Gypsum unit emplaced by gravity flows during the second phase of the MSC (5.60–5.55 Ma; Roveri et al., 2014); and the Upper Gypsum unit, deposited during the final phase of the crisis (5.55–5.33 Ma). The environmental conditions controlling the deposition of Messinian gypsum are still debated, especially due to the lack of modern analogs. The onshore Messinian primary gypsum deposits are largely composed of bottom-grown selenitic (i.e., larger than 2 mm and up to some meters in size) crystals. Among the best examples of bottom-grown laminated, selenite gypsum is the so-called “stromatolitic” gypsum deposits (sensu Rouchy & Monty, 2000 and Allwood et al., 2013) of Cyprus and Crete. Their formation has been explained by a combination of abiotic and microbially driven processes (cf. Allwood et al., 2013).

Intricate networks of filamentous microfossils (Dela Pierre et al., 2015; Panieri et al., 2010; Rouchy, 1982; Rouchy & Monty, 1981, 2000; Schopf et al., 2012; Vai & Ricci Lucchi, 1977), also described as “shoestring-” and “spaghetti-like” structures (Vai & Ricci Lucchi, 1977), are the most striking feature of many gypsum crystals. The phylogenetic affiliation of the microfossils has not been unequivocally determined. It was proposed that the filaments represent brine shrimp fecal pellets (Schreiber & Decima, 1976), algae (Vai & Ricci Lucchi, 1977), or cyanobacteria (Rouchy & Monty, 2000). An assignment to cyanobacteria was further substantiated on the basis of the extraction of 16S rRNA from fossiliferous gypsum of northern Italy (Panieri et al., 2010). Later on, Schopf et al. (2012) suggested that the filaments may rather represent the remains of colorless sulfide-oxidizing bacteria like *Thioploca* or *Beggiaota*. The dense fossil accumulations have not resulted from settling of planktic filamentous organisms, but represent former benthic assemblages of microbial filaments (Dela Pierre et al., 2015). The recognition of abundant aggregates of microcrystalline pyrite and associated polysulfide—the latter representing a diagnostic criterion for this clade of bacteria—within the filaments agrees with the assignment of the filamentous fossils to the colorless sulfide-oxidizing bacteria (Dela Pierre et al., 2015). Based on this circumstance, an unequivocal identification of the phylogenetic affiliation of these microfossils would allow the reconstruction of the depositional environment of bottom-grown Messinian gypsum. Should the assignment to phototrophic cyanobacteria be correct, gypsum would have been deposited within the photic zone, that is, at a maximum depth of 200 m (Lugli et al., 2010). In contrast, colorless sulfide-oxidizing bacteria do not provide any depth constraints—these prokaryotes can live at any depth from benthic to peritidal settings (Bailey et al., 2009)—but can shed light on the redox conditions at the seafloor and the presence of an active biogeochemical sulfur cycle in Messinian brines (cf. Schulz & Jørgensen, 2001).

The analysis of molecular fossils (lipid biomarkers) is a well-established approach to reconstruct the composition of microbial communities preserved in evaporites (e.g., Grice et al., 1998; Jahnke et al., 2004; Turich & Freeman, 2011). The rapid entrapment of lipids within evaporitic minerals tends to favor preservation for up to hundreds of millions of years (e.g., Schinteie & Brocks, 2017; Summons et al., 1999). Gypsum and microbial organosedimentary carbonate deposits are common sedimentary products of modern hypersaline settings (Aref & Taj, 2012; Dupraz et al., 2009; Pettrash et al., 2012; Vogel et al., 2010). These deposits exhibit a typical molecular fossil association dominated by short-chain n-alkanes, fatty acids, and hopanoids, commonly including abundant cyanobacterial lipids (e.g., Bühring et al., 2009; Jahnke et al., 2014). In contrast to modern gypsum deposits, the molecular fossil inventory of the Messinian gypsum has only been rarely investigated and only some groups of lipids (i.e., n-alkanes, alkenones, and isoprenoids) have been used as proxies for paleoclimate (Vasiliou et al., 2017) or for paleoenvironmental reconstruction (Christeleit et al., 2015).

Here, we provide a comprehensive overview on the inventory of molecular fossils preserved in filament-bearing, primary bottom-grown selenitic gypsum formed during the first stage of the MSC in four marginal basins across the Mediterranean. The excellent preservation of molecular fossils, along with the determination of compound-specific carbon stable isotope compositions, enables us (1) to recognize the main groups of micro-organisms inhabiting Messinian aquatic ecosystems at times of gypsum formation and (2) to contribute to the identification of the enigmatic filamentous microfossils.

### 2 | GEOLOGICAL SETTINGS OF THE SAMPLED SECTIONS

Messinian primary gypsum from five sections was collected and analyzed for this study. The samples are from four Mediterranean peripheral basins, (1) the Nijar Basin (southeastern Spain) at the western margin of the Mediterranean, (2) the Vena del Gesso Basin (Italy; two sections) located in the northern sector of the Mediterranean, and the (3) Heraklion (Crete) and (4) Psematismenos (Cyprus) basins in the eastern Mediterranean (Figure 1). The studied samples are from various stratigraphic levels, but they share the presence of densely packed filamentous fossils.

#### 2.1 | Nijar basin

The Nijar Neogene sedimentary succession fills a SW-NE elongated basin located in the internal zone of the Betic Cordillera (southeastern Spain; Fortuin & Krijgsman, 2003). The Messinian succession comprises (a) pre-MSC marine strata of the Abad Marls (a member of the Torre Formation; Van de Poel, 1991), (b) sulfate evaporites of the Yesares Formation (e.g., Fortuin & Krijgsman, 2003), which is considered as the local equivalent of the Mediterranean Primary
Lower Gypsum unit (sensu Roveri et al., 2008), and (c) post-evaporitic continental and lagoonal deposits of the Feos Formation (Fortuin & Krijgsman, 2003; Omodeo Salé et al., 2012). The studied gypsum was sampled from the Yesares Formation exposed in the Gafares section (37°01′28″N, 1°59′17″W); here the Yesares Formation is approximately 70 m thick and consists of a cyclic alternation of at least eight gypsum and laminated marl couplets (Lu, 2006). The Gafares section starts with large selenitic gypsum crystals (the coarse and palmate twinned selenite facies of Lu, 2006) and shows an upward trend toward smaller gypsum crystals (grass-like selenite facies of Lu, 2006). The filament-rich selenitic gypsum was sampled at the base of the Yesares Formation. A dominantly marine origin has been suggested for the Yesares selenites based on $^{87}\text{Sr}/^{86}\text{Sr}$ ratios, trace element distribution, and the marine microfossil assemblages of the intercalated marls (Lu & Meyers, 2003; Lu et al., 2001, 2002).

2.2 | Vena del Gesso basin

The Vena del Gesso Basin is located in the northern Apennines (Italy), exposing a complete succession of the Primary Lower Gypsum unit along a ridge (Figure 2a). This succession consists of 16 cycles composed of gypsum and shale couplets (Lugli et al., 2010; Regghizzi et al., 2018; Vai & Ricci Lucchi, 1977), cut at the top by the Messinian erosional surface (Monticino quarry). The ideal cycle starts with bituminous shales, followed by stromatolitic limestones (Rouchy & Monty, 1981, 2000; Vai & Ricci Lucchi, 1977), which directly underlie the filament-bearing massive selenite facies (Figure 2b; Lugli et al., 2010). The size of the selenite crystals decreases upward, with the appearance of banded selenite and the branching selenite (the latter starting from the 6th cycle) facies completing an ideal cycle. Two samples of filament-bearing selenite gypsum were taken from the basal part of the third Primary Lower Gypsum cycle in the Monte Tondo quarry (44°15′04″N, 11°40′13″E) and from the second cycle in the Monticino quarry (44°13′29″N, 11°45′43″E).

2.3 | Heraklion basin

In the Neogene Heraklion Basin (central Crete), samples of filament-bearing gypsum were taken from the Tsangaraki section, approximately 1 km northeast of Profitis Ilias mountain (Allwood et al., 2013; Rouchy, 1982). Gypsum containing filaments was taken from dome-shaped bodies with stromatolitic morphology, the so-called “selenitic domical or columnar stromatolites” (Allwood et al., 2013). These deposits belong to the Lower Evaporites (Delrieu et al., 1993), which are the equivalent to the Primary Lower Gypsum. According to Rouchy (1982), they formed in a very shallow hypersaline lagoon. In the Tsangaraki area, the 2–3 m thick stromatolitic gypsum interval is located at the base of a gypsum unit (up to 80 m thick) containing different types of gypsum (selenite crystals, nodular and laminated gypsum, and gypsarenites). This unit overlies a chaotic interval of laminated fossiliferous carbonates and a rhythmic succession of laminated marls, diatomites, and fine-grained carbonates.

2.4 | Psematismenos basin

The Psematismenos Basin, located in the southeastern part of Cyprus, is a Neogene sedimentary basin surrounding the Troodos ophiolitic massif (Manzi et al., 2016). In this basin, Messinian evaporites comprise a thick gypsum unit resting on an alternation of pre-MSC sapropels, marls, and carbonate beds (Pakhna Formation). This gypsum unit has been subdivided in the western Polemi Basin into a Lower Gypsum, an Intermediate Breccia, and an Upper Gypsum subunit (Orszag-Sperber et al., 2009). The correlation of these subunits with the MSC chronostatigraphic model is still debated (Krijgsman et al., 2002; Manzi et al., 2016; Orszag-Sperber et al., 2009). According to Orszag-Sperber et al. (2009), in the Kalavasos area of the Psematismenos Basin only the Lower Gypsum is preserved and it correlates with the Primary Lower Gypsum unit of the western Mediterranean, representing the first phase of the MSC. Manzi et al. (2016), however, re-interpreted the Lower Gypsum subunit as an equivalent of the Resedimented Lower Gypsum unit, deposited during the second phase of the crisis (5.60–5.55 Ma; Roveri et al., 2014). The gypsum studied herein was collected in the Tokhni section (34°46′26″N, 33°18′54″E) and consists of selenitic filament-bearing gypsum with domal lamination (Figure 2c,d). According to Manzi et al. (2016), it represents a block of the Primary Lower Gypsum assigned to the Resedimented Lower gypsum unit.

3 | METHODS

Thin sections and thin platy gypsum crystals were studied with transmitted light on a Leica DM4500 P LED optical microscope equipped with a digital camera (Leica DFC 450 C). A FEI Inspect S equipped with an energy dispersive X-ray detection unit (EDAX Apollo XV) was used for scanning electron microscopy (SEM) and semi-quantitative element recognition. For SEM analyses, thin
gypsum slices were treated with deionized water to partly dissolve the selenitic gypsum. Mineralogical composition was determined with X-ray diffraction using a Panalytical X’Pert PRO diffractometer (CuKα radiation, 40 kV, 40 mA, step size 0.0167, 5 s per step; Table 1). The samples were loaded into the sample holders as oriented powder.

One representative gypsum sample from each location (approximately 200 g each) was selected for lipid biomarker analysis. The samples were carefully cleaned with acetone and ground with pestle and mortar to a fine powder. The extraction and separation procedure was carried out following Hoffmann-Sell et al. (2011). The powders were saponified in 6% KOH/MeOH (3 h, 80°C) and three times extracted with a microwave extraction system (MARS X, CEM Corporation) at 80°C and 300 W with dichloromethane (DCM):MeOH (3:1). The resulting total extracts were cleaned by separation into an n-hexane soluble fraction (maltenes) and a DCM-soluble fraction (asphaltenes). Maltenate fractions were separated into four fractions of increasing polarity by column chromatography (hydrocarbons, ketones and esters, alcohols, carboxylic acids). In a second step, the remaining extracted gypsum powder was dissolved in clean water, enriched in NaCl, which had been annealed before, and subsequently extracted and fractionated as described above. The extracts of (1) gypsum powder (average 71 wt% of all extracted lipids) and (2) gypsum powder after dissolution (average 29% of all extracted lipids) yielded the same lipid distribution; for this reason, the total amount of identified compounds (Table 2) represents the sum of quantifications after both extractions.

For GC-MS analysis, alcohols were analyzed as their trimethylsilyl ether (TMS-) derivatives and carboxylic acids as their methyl ester (ME-) derivatives. Compounds from the hydrocarbon, alcohol, and fatty acid fractions were examined by gas chromatography flame ionization detection (GC-FID) and gas chromatography–mass spectrometry (GC–MS) using a Thermo Electron Trace DSQII GC–MS instrument at the MARUM, University of Bremen. Molecules were identified based on retention times and in comparison with published mass spectra. Contents are given in ng/g rock (dry weight). Compound-specific carbon isotope analysis (irm-GC/MS) was performed with a Thermo Electron GC-combustion-III-interface linked to a Thermo Electron Delta-plus XP MS mass spectrometer at the MARUM, University of Bremen. The δ13C values of alcohols and carboxylic acids were corrected for the addition of trimethylsilyl and methyl derivatives, respectively. Each measurement was calibrated using several pulses of carbon dioxide gas with known composition at the beginning and the end of the run. The precision of measurements was checked with a mixture of n-alkanes (C_{15}–C_{25}) with known isotopic composition. The analytical standard deviation was smaller than 0.4‰.

Glycerol dibiphytanyl glycerol tetraethers (GDGTs) were prepared according to the procedure described in Hoffmann-Sell et al. (2011). An aliquot of each non-derivated alcohol fraction was dissolved in n-hexane:propan-2-ol (99:1, v/v). Measurements were performed with high-performance liquid chromatography/ atmospheric pressure ionization mass spectrometry (HPLC/APCI-MS) using an Agilent 1200 series HPLC system coupled to an Agilent 6120A single quadrupole mass spectrometer at the MARUM, University of Bremen. Identification of GDGTs was obtained by screening m/z 400–1400. For GDGT peak integration, single ion monitoring (SIM) of the [M + H]+ ions (m/z 1302, 1300, 1298, 1296, 1292) was used (dwell time 76 ms). Archaeol (m/z 654) was measured on both GC-FID and HPLC-APCI-MS and was used as reference compound. On the GC-FID, it was quantified with n-C_{15} alcohol as internal standard. The quantification of the GDGTs was carried out by correlating GC-FID and HPLC-APCI-MS data for each sample with archaeol. The response factors of archaeol and GDGT-0 on the HPLC-MS-APCI were 1.5 and 1.0, respectively.

4 | RESULTS

4.1 | Petrography and mineralogy

The studied gypsum samples show two types of morphologies: (1) decimeter-sized twinned selenite crystals (arrow-head or swallow-tail twins; Lugli et al., 2010) in the Vena del Gesso Basin (Figure 2b) and (2) decimeter-high stromatolitic build-ups made of gypsum (sensu Rouchy & Monty, 2000; Allwood et al., 2013) in Nijar, Crete, and Cyprus (Figure 2c,d). Both types of gypsum show an internal lamination, characterized by the alternation of mm thick turbid and limpid laminae (Figure 3a, b), possibly reflecting seasonal climate variability (Orti, 2011; Dela Pierre et al., 2015; Reghizzi et al., 2018). In the limpid laminae, solid inclusions are scarce or even absent. In contrast, the turbid laminae entrap densely interwoven filamentous structures (Figures 3 and 4). In particular, in the twinned selenite crystals of the Monte Tondo and Monticino quarries, the filaments occur both in the re-entrant angles of the twins and along the vertical growth bands (Figure 3a), with their maximum lengths aligned to the main crystal faces. Clay-rich aggregates and sparse terrigenous grains are also observed in the turbid laminae. In gypsum build-ups (Nijar, Cyprus, Crete), the turbid laminae (light-colored in hand samples; Figure 2d) are 0.5–1 cm thick and contain dense aggregates (felts) of filaments aligned parallel to the laminae, while the limpid laminae (dark-colored in hand samples) contain much less filaments (Figure 3b). Accessory minerals like dolomite, celestine, and bassanite were detected in all the studied samples (Table 1).

The filaments are characterized by rounded ends and are up to 2 mm long and 60 to 110 µm across, with uniform width along their length (Figures 4 and 5a). Under transmitted light, the filaments appear as opaque tubular features (Figure 4b-g); some filaments are partially hollow (Figure 4c,d). The surface of the filaments is fluorescent when exposed to UV light (Figure 4e). Locally, the filaments contain small dolomite (e.g., in Monte Tondo and Nijar samples) and calcite (e.g., in Monticino, Crete, and Cyprus samples) crystals up to 15 µm in size (Figure 5b). Most filaments include aggregates of µm-sized opaque minerals (Figure 4e) identified as iron sulfide by SEM-EDAX.
Lipid biomarker inventory

The lipid biomarker inventory is largely similar among the various studied samples (see Table 2), although some differences can be recognized. Generally, the western and northern Mediterranean samples (Nijar and Vena del Gesso) show lower total molecular fossil contents, ranging from 1,594 ng/g rock (Monticino) to 3,348 ng/g rock (Nijar). In contrast, the samples from the eastern Mediterranean (Crete and Cyprus) are typified by higher contents (up to 5,179 ng/g rock in the Crete sample; Table 2; Figure 6a). Carboxylic acids are the most abundant molecular fossils, representing more than 60 wt% of the total amount of lipids (Figure 6a).

4.2.1 Hydrocarbons

The average contents of hydrocarbons range from ca.120 ng/g rock in the Monte Tondo and Monticino samples to ca. 420 ng/g rock in Nijar, Crete, and Cyprus. The n-alkanes are the dominant compounds in the hydrocarbon fractions, peaking either at n-C_{18} or n-C_{31}. Some variations in the distribution of short-chain (n-C_{17-22}) and long-chain (n-C_{26-33}) n-alkanes are observed. While Nijar, Crete, and Cyprus samples are characterized by prevailing short-chain n-alkanes, the Vena del Gesso samples show equal contents of short- and long-chain n-alkanes (Table 2). Among the short-chain n-alkanes, the n-C_{17} alkane is not enriched compared to neighboring n-alkanes, representing 8 to 11% of all hydrocarbons. The head-to-tail linked C_{19} isoprenoid pristane and the head-to-tail linked C_{20} isoprenoid phytane are the only isoprenoid hydrocarbons detected.

4.2.2 Alcohols

Gypsum from all study sites contains n-alcohols, although in different proportions; short-chain n-alcohols comprise 16% of the alcohol fraction in Crete and Cyprus samples and about 57 and 55% in Nijar and Monte Tondo samples, respectively. Among all fractions, the molecular fossil inventories vary most from site to site for the alcohol fractions. While in the Nijar sample, the n-alcohols peak at n-C_{16}, n-C_{18} is most abundant in the Monte Tondo sample. In the Cyprus sample, n-C_{24} is most prominent. Apart from n-alcohols, isoprenoid alcohols account for 20% (Nijar) to 63% (Crete) of the compound inventory of the alcohol fractions, also including diphytanyl glycerol diethers (DGDs), glycerol dibiphytanyl glycerol tetraethers (GDGTs),
the C20 isoprenoid phytanol, and sn2- and sn3-phytanylglycerol monooethers (ph-monoether; Figure 6b). Among the DGDs, the C20–20 isoprenoid archaeol is the major isoprenoid alcohol in all sites with contents varying between 18 ng/g rock (Monticino) and 289 ng/g rock (Crete). In the Crete sample, C20–20 archaeol is the most abundant compound representing 26% of all alcohols. Except for the Monticino sample, C20–20 archaeol is always accompanied by C20–25 archaeol (extended archaeol), yielding contents as high as 21 ng/g rock (Crete). In the Crete sample, C20–20 archaeol is the most abundant isoprenoid alcohol in all sites with contents varying between 18 ng/g rock (Monticino) and 289 ng/g rock (Crete). All samples contain GDGTs with a very uniform distribution (Figure 6c). The GDGT with 4 cyclopentane and one cyclohexane moiety (GDGT-5, crenarchaeol; Table 2; Figure 6) predominates in all samples, comprising 40–50% of all GDGTs (Table 2). This compound is accompanied by minor amounts of the crenarchaeol isomer (crenarchaeol; Liu et al., 2016; Sinninghe Damsté et al., 2018). The acyclic GDGT (GDGT-0, caldarchaeol) varies between 15 and 31% of all GDGTs. GDGT-0 is accompanied by almost equal amounts of GDGTs with 1–2 cyclopentane moieties (GDGT-1; GDGT-2) and minor amounts of GDGT with 3 cyclopentane moieties (GDGT-3; Figure 6c).

Phytanol accounts for 2 to 5% of all the isoprenoid alcohols in samples from Nijar, Monte Tondo, and Cyprus, but was absent in the Monticino sample. Phytanol accounts for 20% in Crete and is accompanied by sn2- and sn3-phantanoylglycerol monooethers accounting for of 10 and 2% of all isoprenoid alcohols, respectively. Traces of these phytanyl monooethers are also present in Cyprus (ca. 2%). Further compounds in the alcohol fractions are cholesterol and dinosterol in all samples. Dinosterol is particularly abundant in the Nijar sample (5% of all alcohols) and in the Crete sample (9% of all alcohols). Traces of the pentacyclic triterpenoid tetrahymanol as well as cholestanol were found for few compounds. As for the n-alcohols, the n-fatty acids yielded about 4% higher values than the n-alkanes; only in the Crete sample, δ13C values of n-fatty acids are in the same range like n-alkanes.

4.3 | Compound-specific carbon stable isotopes

A synthesis of the compound-specific δ13C values obtained from the studied gypsum samples is shown in Figure 7. The δ13C hydrocarbon values vary from −33 to −25‰. Long-chain n-alkanes are slightly more 13C-depleted (−33 to −28‰) than short-chain n-alkanes (−29 to −24‰). Besides long-chain n-alkanes, pristane and phytane are among the most 13C-depleted compounds with δ13C values as low as −33‰. The n-alcohols show in general about 4% higher values than the n-alkanes. The isoprenoidal alcohols (phytanol, phytanoylglycerol monooethers, and DGDs) yielded the highest δ13C values ranging from −22 to −13‰. Among them, the DGDs show a narrower range of values between −20 and −14‰. Relatively high values were also obtained for tetrahymanol and the MAGE in the Cyprus (−12 and −18‰, respectively) and Crete samples (−18 and −17‰). The δ13C values of sterols range from −24 to −17‰, with the highest value found for dinosterol in Cyprus and Monticino samples (−17‰).

Among the carboxylic acids, δ13C values were only obtained for few compounds. As for the n-alcohols, the n-fatty acids yielded about 4‰ higher values than the n-alkanes; only in the Crete sample, δ13C values of n-fatty acids are in the same range like n-alkanes. The δ13C values of the predominant short-chain n-alkanes range from −30 to −22‰. Only few data were obtained for monounsaturated n-C16:1 (−26‰; Nijar) and n-C18:1 fatty acids (−25 and −28‰ for Monticino and Crete, respectively), the terminally branched isodecanoic n-C18:1 fatty acid (−15‰; Crete), and phytanic acid (−16‰; Crete). The terminally branched iso-C17:0 fatty acid and phytanic acid yielded similarly high values like the isoprenoidal alcohols.

5 | DISCUSSION

Modern marine gypsum deposits typically form in shallow-water hypersaline environments, including coastal mudflats (sabkhas), lagoons, and human-made salterns. In such settings, biological diversity is limited by high salinity (>110 PSU), which is lethal for most eukaryotes; consequently, these environments are inhabited by highly specialized prokaryotes, such as cyanobacteria and halophilic...
### Table 2: Molecular fossil inventory and compound-specific carbon isotopes of the Messinian gypsum with entrapped filaments

|                | Nijar      | Monte Tondo | Monticino | Crete      | Cyprus     |
|----------------|------------|-------------|-----------|------------|------------|
|                | ng/g rock  | δ¹³C [%o]   | ng/g rock | δ¹³C [%o]  | ng/g rock  | δ¹³C [%o]  | ng/g rock  | δ¹³C [%o]  | ng/g rock  | δ¹³C [%o]  |
| **Hydrocarbons**|            |             |           |            |            |            |            |            |            |            |
| n-C₁₇          | 44         | -25         | 13        | -28        | 8          | -27        | 30         | -25        | 35         | -26        |
| n-C₁₈          | 63         | -27         | 17        | -29        | 12         | -28        | 38         | -28        | 39         | -28        |
| Σ sc-alkanes (n-C₁₇–₂₁) | 231       | -28 to -25  | 56        | -30 to -28 | 40         | -29 to -27 | 250        | -29 to -27 | 159        | -28 to -24 |
| Σ lc-alkanes (n-C₂₆–₃₃) | 169       | -30 to -28  | 60        | -32 to -31 | 32         | -32 to -31 | 182        | -32 to -30 | 110        | -30 to -28 |
| Σ n-alkanes    | 400        |             | 116       |             | 72         |             | 432        |             | 269        |             |
| Pristane       | 40         | -29         | 16        | -30        | 7          | -31        | 26         | n.d.       | 18         | n.d.       |
| Phytane        | 60         | -30         | 17        | -33        | 8          | -33        | 4          | -31        | 22         | -30        |
| Σ hydrocarbons | 500        | 149         | 87        |             | 462        |             |            |            | 309        |             |
| **Alcohols**   |            |             |           |            |            |            |            |            |            |            |
| n-C₁₆          | 132        | -27         | 88        | -29        | n.d.       | n.d.       | 115        | -23        | 54         | -22        |
| n-C₁₈          | 87         | -29         | 259       | -29        | n.d.       | n.d.       | 62         | -29        | 49         | -26        |
| Σ sc-alcohols (n-C₁₄–₂₀) | 350       | -29 to -21  | 375       | n.d.       | n.d.       | 248        | n.d.       | 181        | n.d.       |
| Σ lc-alcohols (n-C₂₆–₂₈) | 35        | -27 to -20  | 46        | n.d.       | 35         | n.d.       | 66         | n.d.       | 126        | n.d.       |
| Σ n-alcohols   | 385        | 421         | 35        |             | 314        | 307        |
| MAGE n-C₁₆:₀   | 0          | n.d.        | 0         | n.d.        | 0          | n.d.       | 28         | -17        | 23         | -18        |
| Phytanol       | 20         | -14         | 14        | -21         | 0          | n.d.       | 108        | -17        | 16         | -22        |
| sn2-phytanyl monoether | 0          | n.d.        | 0         | n.d.        | 0          | n.d.       | 108        | -17        | 16         | -22        |
| sn3-phytanyl monoether | 0          | n.d.        | 0         | n.d.        | 0          | n.d.       | 108        | -17        | 16         | -22        |
| Archaeol       | 54         | -16         | 91        | -20         | 18         | -17        | 289        | -18        | 124        | -14        |
| Extended archaeol | 10       | -15         | 18        | -18         | 0          | n.d.       | 21         | -19        | 19         | -15        |
| Tetrahymanol   | 0          | n.d.        | 0         | n.d.        | 0          | n.d.       | 5          | -18        | 11         | -12        |
| Cholesterol    | 12         | -19         | 56        | -23         | 58         | -22        | 26         | -21        | 0          | -24        |
| Cholestanol    | 0          | n.d.        | 0         | n.d.        | 0          | n.d.       | 37         | -19        | 26         | -21        |
| Dinosterol     | 26         | -19         | 0         | -23         | 5          | -17        | 9          | -24        | 61         | -17        |
| GDGT-0 (calarchaeol) | 4          | 12          | 5         |             | 5          |             | 5          |             | 11         |             |
| GDGT-1         | 2          | 6           | 2         |             | 6          |             | 3          |             |             |             |

(Continues)
| Nijar | Monte Tondo | Monticino | Crete | Cyprus |
|-------|-------------|-----------|-------|--------|
| GDGT-2 | ng/g rock | δ¹³C [%o] | ng/g rock | δ¹³C [%o] | ng/g rock | δ¹³C [%o] | ng/g rock | δ¹³C [%o] | ng/g rock | δ¹³C [%o] |
| 4 | 7 | 2 | 4 | 4 | 2 |
| GDGT-3 | 1 | 4 | 0.5 | 4 | 2 |
| Crenarchaeol | 9 | 34 | 10 | 14 | 15 |
| Crenarchaeol isomer | 1 | 5 | 0.5 | 1 | 1 |
| Sum GDGTs | 21 | 68 | 19 | 34 | 36 |
| Σ isoprenoid alcohols | 105 | 191 | 37 | 707 | 247 |
| Σ alcohols | 528 | 668 | 135 | 1,126 | 675 |

Carboxylic acids

| n-C15:0 | 120 | -24 | 40 | -28 | 52 | -23 | 129 | -25 | 156 | -22 |
| n-C16:0 | 839 | -26 | 715 | n.d. | 725 | -25 | 1,229 | -30 | 1,137 | -25 |
| n-C17:0 | 65 | -22 | 0 | n.d. | 48 | n.d. | 166 | -25 | 219 | -22 |
| n-C18:0 | 301 | -24 | 258 | n.d. | 229 | -26 | 434 | -26 | 505 | -23 |
| n-C26:0 | 151 | n.d. | 307 | n.d. | 27 | -28 | 438 | -28 | 317 | -28 |
| sc-fatty acids (n-C14-19) | 1,795 | -26 to -22 | 1,065 | n.d. | 1,226 | n.d. | 2,574 | n.d. | 2,629 | n.d. |
| lc-fatty acids (n-C26-30) | 172 | n.d. | 332 | n.d. | 49 | n.d. | 485 | n.d. | 356 | n.d. |
| Σ n-fatty acids | 1,967 | 1,397 | 1,275 | 3,059 | 2,985 |

| i-C15:0 | 33 | n.d. | 0 | n.d. | 0 | 47 | 39 | n.d. |
| ai-C15:0 | 36 | n.d. | 0 | n.d. | 0 | 47 | n.d. | 30 | n.d. |
| i-C17:0 | 25 | n.d. | 0 | n.d. | 0 | 95 | 47 | n.d. |
| ai-C17:0 | 15 | n.d. | 0 | -28 | 0 | n.d. | 37 | n.d. | 24 | n.d. |
| n-C16:1 | 114 | -26 | 0 | -30 | 0 | n.d. | 0 | n.d. | 0 | n.d. |
| n-C18:1 | 130 | n.d. | 0 | n.d. | 97 | -25 | 109 | -28 | 120 | n.d. |
| Phytanic acid | 0 | n.d. | 0 | n.d. | 0 | 197 | -16 | 57 | n.d. |
| 17β(H),21β(H)-C₃₂ hopanoic acid | 0 | n.d. | 0 | n.d. | 0 | n.d. | 0 | n.d. | 1 | n.d. |
| Σ carboxylic acids | 2,320 | 1,397 | 1,372 | 3,591 | 3,303 |
| Σ lipids | 3,348 | 2,214 | 1,594 | 5,179 | 4,287 |

Abbreviations: ai, anteiso; GDGT, glycerol dialkyl glycerol tetraethers; i, iso; lc, long chain; n.d., not determined; sc, short chain.
Euryarchaeota (Jebbar et al., 2020; Oren, 2002; Ventosa et al., 2014), and few eukaryotes including the brine shrimp *Artemia salina* and the green alga *Dunaliella salina* (Oren, 2005). Some prokaryotes also live as endoliths in gypsum deposits (Oren et al., 1995; Oren et al., 2009; Stivaletta & Barbieri, 2009; Stivaletta et al., 2010; Jahnke & Des Marais, 2019; Huang et al., 2020). The low biological diversity of modern hypersaline environments is also mirrored by molecular fossil inventories, mostly reflecting input of lipids from halophilic archaea such as DGDs (Dawson et al., 2012; Grice et al., 1998; Turich & Freeman, 2011) and cyanobacteria such as saturated and monounsaturated heptadecanes and heptadecenes (Jahnke et al., 2014). Compared to most modern marine gypsum occurrences (e.g., Jahnke et al., 2014), molecular fossils of the studied Messinian gypsum reflect different ecologies. This difference is probably best explained by the respective depositional environments. While modern gypsum forms in salinas and hypersaline lagoons with water depth at the cm- to m-scale, the primary, bottom-grown Messinian gypsum formed at the seafloor of a marine basin (e.g., Reghizzi et al., 2018).
A significant input of lipids from planktic biota is not to be expected in case of salterns, salinas, and lagoons.

5.1 | DGDs and GDGTs indicate a diverse aquatic archaeal community

The high contents of C_{20:0} DGD (archaeol) co-occurring with C_{20:5} DGD (extended archaeol)—a compound diagnostic of halophilic archaea (Dawson et al., 2012; Teixidor et al., 1993)—in all samples except for Monticino (Figure 6) agree with the presence of a community of halophilic archaea. Archaeol is the major membrane lipid of halophiles (Dawson et al., 2012; Kates, 1993), but is also a common membrane lipid of other archaea, particularly methanogenic and methanotrophic euryarchae (e.g., Hinrichs et al., 2000; Koga et al., 1998). Other archaeal-producing archaea are marine planktic, mesophilic euryarchaeal groups (e.g., Sollai et al., 2019). Among the potential source organisms of the DGDs extracted from Messinian gypsum, methanotrophic archaea can be excluded, since archaeol (average $\delta^{13}C = -17\%$) does not show the typical $\delta^{13}C$ values of lipids of this group of archaea (as low as $-130\%$; e.g., Niemann & Elvert, 2008; Himmel et al., 2015) in any of the Mesinian gypsum samples. The lowest $\delta^{13}C$ value of archaeol was found for the Monte Tondo sample (−20%). The fact that this $\delta^{13}C$ value is similar to values of compounds of presumed planktic algal origin (i.e., sterols: $-21\%$ on average; Figure 6) and values of planktic archaeal origin (i.e., bicyclic and tricyclic biphytanes; $-20\%$ on average; see below for further discussion) agrees with mesopholic planktic euryarchae as source of archaeol. The $\delta^{13}C$ values of archaeol from the other sites reflect even less $^{13}C$ depletion ($\delta^{13}C$ values as high as $-14\%$, Cyprus). Since extended archaeol exhibits the same $\delta^{13}C$ value like archaeol for all study sites, halophilic euryarchae, probably planktic, are the most likely producers of both DGDs. Apart from Monte Tondo, $\Delta\delta^{13}C_{\text{DGD-sterol}}$ is as high as $+7\%$. A similar offset between the lipids of a halophilic archaeal community (DGD producer) and algal phytoplankton (sterol producers) has been described for the late Miocene evaporite-bearing deposits of the Dead Sea Basin (Grice et al., 1998). The observed variability of $\delta^{13}C$ values between halophilic archaea (average $\delta^{13}C = -17\%$) and phytoplankton (−21% on average) is in accord with water column stratification, reflecting variations in salinity across the water column. However, the high $\delta^{13}C$ values of the lipids of halophilic archaea in such settings are not yet fully understood (Birgel et al., 2014; Grice et al., 1998). The large $\Delta\delta^{13}C_{\text{DGD-sterol}}$ could be possibly explained by the heterotrophic lifestyle of halophilic archaea, involving the uptake of relatively $^{13}C$-enriched carbon from carbohydrates and proteins by halophilic archaea in contrast to photoautotrophic algae assimilating dissolved carbon dioxide (Oren, 1994). Carbohydrates and proteins are generally less $^{13}C$-depleted than other organic substrates (e.g., lipids; De Niro and Epstein, 1977, Grice et al., 1998).

Phytanol and the two phytanylglycerol monoethers extracted from Crete and Cyprus gypsum revealed similar $\delta^{13}C$ values (−22 to −13%) to archaeol and extended archaeol (cf. Birgel et al., 2014; Ziegenbalg et al., 2012); most likely the former compounds represent degradation products of DGDs (cf. Liu et al., 2018). The isoprenoid hydrocarbon phytane, however, is more $^{13}C$-depleted ($\delta^{13}C$: −33 to −30%) than other isoprenoids with phytanyl moieties (Figure 7). Phytane probably derives either from other heterotrophic archael communities (Wakeham et al., 2003) or from a secondary, possibly terrigenous source, and is unrelated to the other compounds with phytanyl moieties. To sum up, the lipid biomarker inventory of Messinian gypsum confirms the prominent presence of halophilic archaea in the depositional paleoenvironment—not a surprising result if the gypsum is interpreted as a product of fully hypersaline conditions. Halophilic archaea are commonly thriving in high salinity environments like in some modern shallow-water microbial mats (Bühring et al., 2009; Jahnke et al., 2014). However, uncultured members of the genus Halobacteria have also been recognized in aquatic systems with low salt concentrations such as at sulfur-rich springs (Elshahed et al., 2004), estuaries (Purdy et al., 2004; Singh et al., 2010), and the Black Sea, where the salinity of bottom waters is ca. 20 PSU (Jessen et al., 2016). In the latter environment, Halobacteria were found in sediments covering Beggiatoa-dominated microbial mats, occurring where the chemocline intercepts the sea floor and steep gradients of molecular oxygen and hydrogen sulfide exist. According to this example, sulfur-rich microenvironments may have also favored the proliferation of halophilic archaea (cf. Elshahed et al., 2004) in low salinity waters (i.e., with low contents of Na⁺ and Cl⁻ ions) during Messinian gypsum formation (cf. Natalicchio et al., 2014; Evans et al., 2015; see paragraph 5.3).
The observed GDGT assemblages are much different from those reported for modern shallow-water hypersaline settings (Petrick et al., 2019). Halophilic archaea are not known to produce GDGTs (Schouten et al., 2013; Teixidor et al., 1993). Although methanogens isolated from hypersaline lakes have been shown to produce GDGT-0 and GDGT-1, they do not produce extended archaeol (Bale et al., 2019). Most of the GDGTs found in marine sediments are thought to be produced by planktic, ammonia-oxidizing Thaumarchaeota (Brochier-Armanet et al., 2008; Könneke et al., 2005; Schouten et al., 2013; Spang et al., 2010; Wuchter et al., 2006). Crenarchaeol, which is the most abundant GDGT in the studied Messinian gypsum (Figure 5c), is also the predominant GDGT of Thaumarchaeota in the marine epipelagic realm (Schouten et al., 2013; Turich et al., 2007). It is also produced by soil-dwelling Thaumarchaeota (Sinninghe Damsté et al., 2012) and occurs in marine hydrothermal settings (Pearson et al., 2004; De La Torre et al., 2008). In contrast to soil-dwelling Thaumarchaeota, marine Thaumarchaeota produce not only crenarchaeol, but also the acyclic GDGT-0 (caldarchaeol) and minor amounts of cyclopentane ring containing GDGTs (GDGT-1 to GDGT-4). Interestingly, the Messinian gypsum shows a GDGT-distribution similar to that of marine Thaumarchaeota from the modern Mediterranean Sea (Besseling et al., 2019). A derivation of GDGTs from marine Thaumarchaeota also agrees with the GDGT-0/crenarchaeol ratios ranging from 0.4 and 0.7, similar to ratios reported for modern open marine environments (0.5 to 1; Zhang et al., 2016; Petrick et al., 2019). In modern (Stowakiewicz et al., 2016) and ancient (Cheng et al., 2017) hypersaline settings, on the other hand GDGTs are scarce and mostly dominated by GDGT-0.

The δ\(^{13}\)C values of biphytanes found in the Monte Tondo and Cyprus samples help to constrain the sources of (1) caldarchaeol, mostly yielding acyclic biphytanes (BP0) after ether cleavage, and (2) crenarchaeol, the probably source of tricyclic biphytane (BP3) after ether cleavage. The monocyclic (BP1) and bicyclic (BP2) biphytanes derive from GDGTs 1 to 3 and crenarchaeol (BP2 only). The δ\(^{13}\)C values of BP3 (ca. −20‰ on average) agree with crenarchaeol deriving from marine autotrophic Thaumarchaeota, an isotopic composition close to that found in modern oceans (ca. −21‰; Schouten et al., 2013). The negative δ\(^{13}\)C values of BP0 (as low as −37‰), on the other hand, point to a different source of caldarchaeol, including heterotrophic, possibly benthic archaeal communities (cf. Hoffmann-Sell et al., 2011; Pearson et al., 2016; Wakeham et al., 2003). The former presence of benthic archaea in addition to planktic archaea is supported by relatively high contents of GDGT-1 (up to 18%, Crete) and GDGT-2 (up to 18%, Nijar) compared to modern marine GDGT distributions; these compounds are less abundant in marine planktic archaea (<10% of total GDGTs; Lipp & Hinrichs, 2009). The phylogenetic affiliation of benthic archaea cannot be determined.
with certainty, yet methanotrophic archaea can be excluded since this clade reveals different GDGT patterns with the predominance of GDGT-2, $^{13}$C depletion, and other isoprenoids not observed in the Messinian gypsum (Birgel et al., 2006, 2008; Natalicchio et al., 2012). A possible group from which the $^{13}$C-depleted biphytanes could derive are methanogenic euryarchaeota (Hoffmann-Sell et al., 2011; Koga et al., 1998). Among other factors, the isotope composition of lipids synthesized by methanogens depends largely on the metabolized substrate (Londry et al., 2008). Unfortunately, the isotopic composition of GDGT-derived biphytanes has not yet been determined in culture experiments with methanogens. Since archaeol, interpreted to be derived from halophilic archaea, yielded $\delta^{13}$C values that are 15 to 20% higher than the values of acyclic and monocyclic biphytanes, an origin of the latter compounds from halophiles can be excluded—an interpretation in accord with the known compound inventories of halophiles.

To sum up, the lipid inventory of Messinian gypsum suggests the former presence of three archaeal communities including (1) halophiles typified by relatively $^{13}$C-enriched DGDs and phytanyl monoothers, (2) planktic thaumarchaeota yielding crenarchaeol ($\delta^{13}$C: ca. $-20\%a$), and (3) a group of probably benthic archaea producing $^{13}$C-depleted GDGT-0 and the cyclic GDGTs 1 and 2. Such high archaeal diversity is not observed in modern shallow-water hypersaline environments (Słowakiewicz et al., 2016; Petrick et al., 2019) and rather agrees with a relatively deep stratified marine basin (Christelet et al., 2015; García-Keigas et al., 2018). The planktic thaumarchaeal community is believed to have lived in the upper water column, probably typified by diluted marine conditions (Grothe et al., 2020; Natalicchio et al., 2014; Reghizzi et al., 2018), whereas the halophiles inhabited the deeper, more saline part of the water column. Such interpretation is supported by the presence of tetrahymanol in the Cyprus and Crete samples, a proxy of water column stratification (Sinninghe-Damsté et al., 1995; Natalicchio et al., 2017; Sabino et al., 2021).

5.2 The affiliation of filamentous microfossils

The affiliation of the superabundant filamentous microfossils preserved in Messinian gypsum has not yet been resolved with certainty. At first, they were interpreted to represent fecal pellets of brine shrimps (Schreiber and Decima, 1976), and later, they were assigned to algae (Vai & Ricci Lucchi, 1977), cyanobacteria (Panieri et al., 2010; Rouchy and Monty, 2000) or sulfide-oxidizing bacteria (Dela Pierre et al., 2015; Schopf et al., 2012). An attribution to fecal pellets does not agree with the partial hollowness of the studied filaments (Figure 4e). Hollowness is not expected in fecal pellets, but is strong evidence of a microbial origin of the filamentous microfossils (Andreottato et al., 2019, and reference therein). Such origin is in accord with the curved shape of filaments and their internal segmentation (Dela Pierre et al., 2015). The most striking argument for an assignment to cyanobacteria was the extraction of 16S rRNA from samples of the Vena del Gesso gypsum, matching with several cyanobacterial clones and in particular with those of the genus *Geitlerinema*, typical of coastal shallow marine environments (Panieri et al., 2010). The finding of genetic material in several million-year-old geological samples requires excellent preservation of organic matter, supposedly caused by early permineralization in gypsum prior to cell decay and disintegration (cf. Schopf et al., 2012). Molecular fossils, which are much more stable than 16S rRNA, should therefore confirm the suggested cyanobacterial origin of filaments. Cyanobacteria synthesize lipid biomarkers (Table 3), including diplopterol (Bühring et al., 2009; Rohmer et al., 1984) and bacteriohopanepolyols (Jahnke et al., 2004; Summons et al., 1999; Talbot et al., 2008). Other diagnostic cyanobacterial lipids are short-chain n-alkanes, but especially heptadecane and heptadecenes, as well as methylated hepta- and octadecanes (Birgel et al., 2015; Hefter et al., 1993; Jahnke et al., 2014; Kozlowski et al., 2018; Wieland et al., 2008). Some of these compounds or their degradation products tend to be preserved for millions of years in the geological record (Heindel et al., 2015; Summons et al., 1999). Among these cyanobacterial biomarkers, only n-C$_{17}$-alkane (heptadecane) is present in the studied Messinian gypsum (Table 3), yet with moderate contents only, accounting on average for 7% of the total hydrocarbon fraction and corresponding to n-C$_{17}$/n-C$_{18}$ ratios below 1. In cyanobacterial mats from modern hypersaline environments, n-C$_{17}$ accounts for approximately 80% of the total hydrocarbon fraction and the n-C$_{17}$/n-C$_{18}$ ratio is significantly above 1 (e.g., Bühring et al., 2009; Jahnke et al., 2004). Despite heptadecanes being prone to degradation (Hefter et al., 1993; Wieland et al., 2008), in cases of 16S rRNA preservation, these cyanobacterial-derived hydrocarbons should be present as well. Although such negative evidence can obviously not exclude the presence of cyanobacteria in the Messinian depositional environment, the scarcity of cyanobacterial biomarkers (see Table 3) renders unlikely an assignment of the superabundant filamentous microfossils to cyanobacteria.

The other previously suggested hypothesis is that the interwoven filaments in the Messinian gypsum represent the remains of colorless sulfide-oxidizing bacteria like *Beggioa* and *Thioploca* (Dela Pierre et al., 2015; Schopf et al., 2012). Microbial mats dominated by these large prokaryotes are found in coastal upwelling areas (Arning et al., 2008; Bailey et al., 2009), in stratified basins (e.g., the Black Sea; Jessen et al., 2016), as well as in sulfide-rich marine sediments associated with methane seepage (Knittel et al., 2003; Zhang et al., 2005). The lipid biomarker inventory of mats of sulfide-oxidizing bacteria is typified by high contents of saturated (C$_{16:0}$ and C$_{18:0}$) and monounsaturated (C$_{16:1}$ and C$_{18:1}$) fatty acids (Arning et al., 2008; Jacq et al., 1989; Jiang et al., 2012; McCaffrey et al., 1989; Zhang et al., 2005). The high abundance of these compounds in the majority of the studied Messinian samples is consistent with this hypothesis (Table 3). Unfortunately, other organisms including many bacteria (Elvert et al., 2003; Londry et al., 2004) and diverse phytoplankton (Viso & Marty, 1993; Wakeham, 1995) are known to produce these largely unspecific fatty acids too.

Given the low specificity of the lipids synthesized by sulfide-oxidizing bacteria (Arning et al., 2008), the origin of the filaments can be assessed further with the overall lipid assemblage and the
carbon stable isotope patterns. Compounds synthesized by sulfate-reducing bacteria such as *iso*- and *anteiso*-*C*<sub>15</sub>:0 and *C*<sub>17</sub>:0 fatty acids (Kaneda, 1991; Perry et al., 1979; Taylor & Parkes, 1983) are present in gypsum from Nijar, Crete, and Cyprus. In the case of Cyprus and Crete gypsum, which are the samples with the highest total lipid abundance (Table 2), even a MAGE—a biomarker of sulfate-reducing bacteria, as suggested by the biomarker inventory of some of the Messinian gypsum, has been reported for different modern environments (e.g., Arning et al., 2008; Fukui et al., 2015), which is a clade diagnostic feature of colorless sulfide-oxidizing bacteria (Schulz & Jørgensen, 2001). Interestingly, a tight association of sulfide-oxidizing bacteria with sulfate-reducing bacteria, as suggested by the biomarker inventory of some of the Messinian gypsum, has been reported for different modern environments (e.g., Arning et al., 2008; Fukui et al., 2019; SamKamaleson et al., 2021).

### 5.3 | Paleoenvironmental implications

Provided that the assignment of the filamentous fossils to benthic sulfide-oxidizing bacteria like *Beggiatoa* and *Thioploca* is correct and given the association of filamentous body fossils with molecular fossils of sulfate-reducing bacteria—especially in the samples with the highest total lipid contents (i.e., Nijar, Cyprus and Crete)—the basin floor where the Messinian gypsum grew was covered by a benthic assemblage of chemotrophic bacteria involved in an intense sulfur cycle. Such interpretation suggests suboxic to anoxic bottom water conditions and production of hydrogen sulfide through intense bacterial sulfate reduction, in turn sustaining sulfide-oxidizing bacteria (cf. Bailey et al., 2009; Teske et al., 2009). Sulfate-reducing bacteria live in suboxic to anoxic environments, using low molecular weight organic compounds or molecular hydrogen as electron donors (Grossi et al., 2015; Vinçon-Laugier et al., 2016). In case of the Messinian depositional environment, the electron donor was probably a mixture of organic substrates deriving from (1) phytoplankton (diatoms, dinoflagellates) and (2) planktic archaea. The circumstance that the δ<sup>13</sup>C values of the MAGE (average ca. −17‰; Crete and Crete) are similar to those of archaeal-derived compounds (archaeol...
and sn2/sn3-phytanyl monoethers) is in favor of sulfate-reducing bacteria metabolizing low molecular weight compounds deriving from archaeal biomass. This hypothesis is in line with recent studies of the Dead Sea, where bacteria have been found to develop strategies involving the recycling of archaeal necromass (Thomas et al., 2019), which represents the most abundant organic carbon source in this saline basin (Oren, 1999).

The precipitation of gypsum probably caused the rapid entrapment of the benthic assemblage of microbial filaments. The paleoenvironmental conditions behind the formation of the Messinian gypsum are still discussed (e.g., García- Veigas et al., 2018; Grothe et al., 2020; Natalicchio et al., 2014). Data from fluid inclusions in bottom-grown selenitic gypsum challenged the idea that gypsum formed from hypersaline brines resulting from seawater evaporation. Instead, these data indicated that gypsum formed from low salinity waters (i.e., with low contents of Na⁺ and Cl⁻ ions) equivalent to 1.6 wt% sodium chloride on average (Costanzo et al., 2019; Evans et al., 2015; Natalicchio et al., 2014). The water masses probably represented a mixture of marine water, freshwater from Mediterranean rivers (Reghizzi et al., 2018), and brackish water from the Paratethys (Grothe et al., 2020). Interestingly, studies on modern gypsum suggest the involvement of prokaryotes in gypsum formation (van Driessche et al., 2019; Lepinay et al., 2018; Mansor et al., 2018; Thompson & Ferris, 1990). In particular, the oxidation of reduced sulfur species by sulfide-oxidizing bacteria may supply, at least in part, the sulfate needed for gypsum precipitation (e.g., Lepinay et al., 2018; Mansor et al., 2018). Such microbial oxidation has been suggested to promote gypsum supersaturation in bottom waters and to favor the formation of gypsiferous thrombolites in a hypersaline lagoon in Venezuela (Petrasch et al., 2012). Since we interpret the fossilized filaments in Messinian gypsum as the remains of sulfide-oxidizing bacteria, a possible microbial involvement in gypsum precipitation should be taken into account in future studies.

6 | CONCLUSIONS

Primary Messinian bottom-grown gypsum deposits, formed during the first stage of the MSC (5.97–5.60 Ma), preserve biosignatures of ancient microbial communities, including diverse molecular fossils and mazes of filamentous microfossils. The molecular fossil assemblages of gypsum from different Mediterranean subbasins are unlike typical assemblage of modern marine gypsum deposits forming in shallow-water hypersaline settings. The abundance of lipids of planktic halophilic archaea, planktic thaumarchaea, and a poorly constrained community of benthic archaea confirms that gypsum formed in a stratified basin typified by a normal marine to diluted upper water column and more saline deeper waters. Petrographic
and mineralogical observations, the overall biomarker inventories, and compound-specific carbon stable isotope patterns agree best with the interpretation that the superabundant filamentous microfossils enclosed in Messinian gypsum are the remains of sulfide-oxidizing bacteria. Sulfide-oxidizing bacteria and sulfate-reducing bacteria were probably the main constituents of chemotrophic microbial mats on the Messinian seafloor.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest regarding the publication of this article.

DATA AVAILABILITY STATEMENT

All data of this article are available in PANGAEA Data Publisher.

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