Barite Precipitation on Suspended Organic Matter in the Mesopelagic Zone

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Mechanisms underlying barite precipitation in seawater and the precise depths of barite precipitation in the water column have been debated for decades. Here we present a detailed study of water column barite distribution in the mesopelagic zone at diverse stations in the open ocean by analyzing samples collected using multiple unit large volume in-situ filtration systems in the Pacific, Atlantic and Indian oceans. Our results demonstrate that barite is an organo-mineral particularly abundant at intermediate depths throughout the world’s ocean regardless of saturation state with respect to barite. This is confirming the notion of precipitation at depths of intense organic matter mineralization. Our observations further support the link between barite formation and microbial activity, demonstrated by the association of barite particles with organic matter aggregates and with extracellular polymeric substances. Evidence for microbial mediation is consistent with previous experimental work showing that in bacterial biofilms Ba binds to phosphate groups on cell surfaces and within extracellular polymeric substances. This organo-accumulation promotes high concentrations of Ba leading to saturated microenvironments and nucleation sites favoring precipitation. The distribution of Ba isotopes in the water column and in particulate matter is also consistent with the proposed precipitation mechanism.

Keywords: pelagic barite, organo-mineralization, barite saturation state, extracellular polymeric substances, bioaccumulation

INTRODUCTION

Barium and barite are routinely used for reconstructing past export production in the ocean yet the processes linking barite formation to export production are still elusive. Since the early work of Chow and Goldberg (1960) who reported high Ba concentrations in marine sediments underlying regions of high biological productivity, a link between organic matter fluxes and Ba abundance has been broadly demonstrated (e.g., Dehairs et al., 1980; Dymond et al., 1992; Francois et al., 1995; Paytan et al., 1996; Paytan and Griffith 2007; Griffith and Paytan, 2012; Carter et al., 2020 and references therein). Studies using sediment traps have provided further evidence on the association of particulate Ba and particulate organic carbon (POC). It has been proposed that barite
precipitates in close association with aggregates of organic matter and sinking biological debris (e.g., Bishop, 1988; Dehairs et al., 1991). Different algorithms have been suggested to correlate export production and excess Ba (total Ba concentration corrected for the lithogenic phase) or barite accumulation (e.g., Dymond et al., 1992; Francois et al., 1995; Paytan et al., 1996; Eagle et al., 2003), allowing the reconstruction of past ocean productivity (e.g., Dymond et al., 1992; Ginele and Dahmke, 1994; Nürnberg et al., 1997; Eagle et al., 2003; Ma et al., 2015). Nevertheless, quantification of export production from Ba proxies is still hindered by poor understanding of the mechanisms and processes leading to barite (the main phase carrying excess Ba) formation in the oceanic water column. Moreover, barite distribution in the oceanic water column is variable in space, time, and depth and such variability is not yet fully understood (e.g., Hernandez-Sanchez et al., 2011; Bates et al., 2017).

Over decades of research, several hypotheses have been proposed to explain barite precipitation in the oceanic water column given that most of the world’s ocean mesopelagic zone (200–1,000 m depth, Sutton et al., 2017) is undersaturated with respect to barite (Monnin et al., 1999; Rushdi et al., 2000). Hypotheses included precipitation in microenvironments formed within sinking biogenic particulate matter (e.g., Dehairs et al., 1980; Bishop, 1988), precipitation as a result of celestine (SrSO₄) dissolution from Acantharian tests (e.g., Bernstein et al., 1992; Bernstein et al., 1998) and by way of microbially mediated precipitation processes (Gonzalez-Muñoz et al., 2003; Gonzalez-Muñoz et al., 2012; Torres-Crespo et al., 2015). Specifically, Gonzalez-Muñoz et al. (2003) demonstrated in laboratory culture experiments the ability of soil bacteria to induce precipitation of barite, and later also highlighted the potential role of bacteria in barite precipitation in the ocean by using diverse marine strains in culture experiments (Gonzalez-Muñoz et al., 2012; Torres-Crespo et al., 2015). Bacterially mediated precipitation of barite is consistent with studies demonstrating the positive correlation between mesopelagic particulate Ba abundance and enhanced bacterial production in the North Pacific and the Southern Ocean as well as the relation between particulate Ba abundance and microbial oxygen consumption (Dehairs et al., 2008; Jacquet et al., 2011; Planchon et al., 2013). Moreover, mesocosm experiments inducing the decay of various phytoplankton species in the dark demonstrated that Ba is released during the decomposition of the phytoplankton, leading to barite precipitation (Ganeshram et al., 2003). Overall, several lines of evidence have supported the suggestion that the nutrient-like behavior of Ba is due to biological processes mediating barite precipitation. In particular, analyses of Ba isotopes of both water column and particulate matter at various sites in the ocean (e.g., Horner et al., 2015; Bates et al., 2017; Bridgestock et al., 2018) are also consistent with the formation of barite which preferentially incorporates the light Ba isotope at mesopelagic depths.

Additional experimental work (Martinez-Ruiz et al., 2018) demonstrated that bacterial biofilms, specifically, extracellular polymeric substances (EPS) may play a major role in barite precipitation by providing nucleation sites to locally enhance Ba concentration leading to barite precipitation. This work also showed that an amorphous P-rich phase is formed at the initial stages of Ba bioaccumulation eventually being replaced by sulfate and leading to the formation of barite crystals. The capacity of EPS to bind metal ions to negatively charged functional groups has been broadly demonstrated (e.g., Braissant et al., 2007; Tourney and Ngwenya, 2014) and the role of phospholipids acting as nucleation sites to incorporate diverse cations has been demonstrated for diverse metals such as U (e.g., Morcillo et al., 2014). Moreover, P-rich precursors have been described in the precipitation of several minerals such as aragonite apatite and iron oxides, both under experimental conditions (Rivadeneyra et al., 2010), and in the geological record (e.g., Sanchez-Navas and Martin-Algarra, 2001; Miot et al., 2009). In general, microbial precipitation through an amorphous precursor is a widespread process in natural environments (e.g., Weiner et al., 2005; Enyedi et al., 2020). Such crystallization path through an amorphous precursor phase is also associated with the inorganic precipitation of barite. For example, a metastable amorphous Ba sulfate phase that precedes barite formation has been recognized during the early stages of barium sulfate crystallization from aqueous solutions (Ruiz-Agudo et al., 2020), however, the binding of Ba to EPS in the open ocean has not been thoroughly investigated.

Mineralogical and crystallographic analyses of marine barite collected using multiple unit large volume in-situ filtration systems (MUL-VFS) at two stations in the North Atlantic and the Atlantic sector of the Southern Ocean demonstrated that barite formation in the open ocean also involves an amorphous precursor (Martinez-Ruiz et al., 2019). This initial amorphous precursor is a phosphorus-rich phase that evolves into barite when phosphate groups are substituted by sulfate. These crystallization pathways are similar to those reported to form within biofilms in laboratory based experiments (Martinez-Ruiz et al., 2018). These findings support the role of EPS in the precipitation of barite in the oceanic water column and the correlation between bacterial production and the abundance of Ba-rich particles. As our previous study (Martinez-Ruiz et al., 2019) focused only on two stations, from the North Atlantic and the Atlantic Sector of the Southern Ocean, the present work aims at providing new insights into the mechanisms of barite formation in the ocean water column by analyzing barite, also collected by MUL-VFS, at new stations representing additional ocean sectors with diverse barite saturation conditions. Stations within each sector were selected in high productivity settings. Barite crystallography, mineralogy and abundance in the water column have been investigated at different depths within the mesopelagic zone at the locations and sites described below.

SAMPLES AND METHODS

Oceanographic Setting

Locations for this study have been selected to represent four different ocean sectors (Figure 1; Table 1) where productivity is relatively high and hence barite is abundant in the upper water column however these sites represent different saturation states within the mesopelagic zone.
North Pacific: OC1608A, C-SNOW cruise, California current system, coastal North Pacific Station 1 (1,929 m water depth), and OC1608A, C-SNOW cruise, California current system, Santa Barbara basin, Station 2 (4,880 m water depth). These stations are at the edge of the North Pacific oligotrophic gyre. The North Pacific Subtropical Gyre harbors one of the largest biomes on Earth, it is a relatively stable oligotrophic environment, with low surface concentrations of nitrogen and phosphorus. Nutrients derived from advective transport from depth into the surface ocean stimulates primary production in this region (Karl and Church, 2017; Robidart et al., 2019). The two stations in the Santa Barbara basin are on the edge of this gyre and are characterized by relatively high productivity and high phytoplankton biomass that supports a productive pelagic ecosystem (Letelier, et al., 2019). The productivity is fueled by intensive coastal upwelling induced by northerly winds along the California margin (e.g., Brzezinski and Washburn, 2011; Abella-Gutiérrez and Herguera, 2016).

Atlantic: MV1101, Great Calcite Belt (GCB) 1, South Atlantic, Station 117 (Rosengard et al., 2015; Balch et al., 2016). During the GCB1 cruise the R/V Melville crossed the Atlantic sector from Punta Arenas, Chile, to Cape Town, South Africa, sampling between 39° S and 59° S. Station 92 from this cruise has been previously analyzed for marine barite (Martinez-Ruiz et al., 2018) and station 117 has been selected for this study (5,048 m water depth). Station GCB1-117 is located in the subtropical region of the South Atlantic Ocean and is dominated by seasonal coccolithophores and diatoms blooms (Smith et al., 2017).

Indian ocean: RR1202, Great Calcite Belt (GCB) 2, South Indian Ocean, Station 63 (Rosengard et al., 2015; Balch et al., 2016; Smith et al., 2017). During the GCB2 cruise, the R/V Revelle crossed the Indian sector from Durban, South Africa, to Perth, Australia, sampling between 37° S and 60° S. Station 63 has been selected for this study (1,310 m water depth). The region is characterized by elevated surface reflectance that is thought to result from high seasonal concentrations of coccolithophores. Data for multiple parameters sampled during the GCB cruises including chlorophyll, particulate inorganic carbon (PIC), POC, biogenic silica (BSi), coccolithophore concentration, calcification, photosynthesis, dissolved inorganic carbon (DIC), total alkalinity, iron limitation of phytoplankton and 234Th-based vertical flux rates are available for these stations (Rosengard et al., 2015; Balch et al., 2016). Shipboard scientists reported dense coccolithophore populations that exported small, highly degraded, and compact particles out of the euphotic zone. Coccolithophore blooms are considered very efficient in transferring POC to the base of the mesopelagic zone, although the magnitude of exported POC is not as high as in diatom-rich regions (e.g., Henson et al., 2012).

South Pacific: NBP1101, Seafarers cruise, Ross Sea, Station 14. This station is located off the Ross Sea Shelf in the Pacific (1,887 m). Samples were collected between January 17, and February 13, 2011 aboard the R/V Nathaniel B (Hatta et al., 2017). The Ross Sea continental shelf is one of the most productive areas in the Southern Ocean (e.g., Smith, Jr. et al., 2014). Here a significant supply of dissolved Fe to surface waters is required to sustain high productivity (Sedwick et al., 2011; Hatta et al., 2017), and include dust, sea-ice, icebergs and upwelling of deeper waters as some of the main inputs (e.g., Measures et al., 2012; Marsay et al., 2014).

**Particulate Ba Sampling, Particulate Organic Carbon Analyses, and Barite Saturation State (Ω**$_{\text{barite}}$**)**

Size-fractionated particulate material has been collected using multiple MUL-VFS (Bishop et al., 1985) and battery-operated McLane in-situ pumps (LV-WTS) (Rosengard et al., 2015). Analyzed samples and corresponding depths are indicated in Table 1. Particulate Ba concentrations (pBa) have been

![Figure 1](https://svs.gsfc.nasa.gov/30801)
determined at the South Atlantic (Great Calcite Belt) and South Pacific Ocean (Antarctic sector) stations, but these data are not available at other stations. In situ deployed filters were processed using the protocol described in Bishop and Wood (2008). Samples for p[Ba] were collected on PES filters to ensure low blank and Ba concentrations in the particulate leachate was analyzed using an iCAP RQ inductively-coupled plasma mass spectrometer. Quantification was achieved via comparison of blank- and indium-normalized ion beam intensities in samples against those measured in a serially diluted multi-element standard that was prepared in house. Precision is generally better than ±3% relative standard deviation. POC samples were collected on pre-combusted QMA filters and concentrations were measured using a CHN elemental analyzer immediately on the ship during the cruises as described in Rosengard et al. (2015). Sampling details and complication associated with particles collected by MUL-VFS as well as retention efficiency are discussed in detail in Bishop et al. (2012). In the South Indian ocean station, both large (>51 μm) and small (1–51 μm) size particulates were analyzed. For the rest of stations, only filters retaining the 1–51 μm fraction were analyzed. The MUL-VFS sampling was found to be highly suitable for barite particles retention and QMA filters were ideal for barite microscopic detection and observation.

To place the p[Ba] data in context, we calculated the barite saturation state of seawater with respect to barite (Ω\textsubscript{barite}) at the depths of sample collection. Ω\textsubscript{barite} is the ratio between the barium and sulfate ion activity product and the barite solubility product. Values of Ω <1, =1, and >1 indicate undersaturation, perfect saturation, and super-saturation, respectively. For consistency with the literature, we consider water samples with Ω between 0.9 and 1.1 as being in saturation equilibrium (e.g., Monnin et al., 1999). Since co-located

| Ocean (bottom depth) | Coordinates | Sample | Depth (m) | p[Ba] (pM) | POC (μM) |
|----------------------|-------------|--------|-----------|------------|----------|
| **East Pacific (C-SNOW cruise)** | | | | | |
| St 1 — Santa Barbara basin (1,929 m) | | CS 1 | 30 | | |
| | | CS 2 | 107 | | |
| | | CS 3 | 160 | | |
| | | CS 4 | 267 | | |
| | | CS 5 | 535 | | |
| | | CS 6 | 59 | | |
| | | CS 7 | 70 | | |
| | | CS 8 | 99 | | |
| | | CS 9 | 149 | | |
| | | CS 10 | 248 | | |
| | | CS 11 | | | |
| (1,927 m) | | CS 12 | | | |
| | | CS 13 | 55 | | |
| | | CS 14 | 101 | | |
| | | CS 15 | 151 | | |
| | | CS 16 | 251 | | |
| | | CS 17 | 503 | | |
| | | CS 18 | | | |
| | | CS 19 | 25 | | |
| | | CS 20 | 151 | | |
| | | CS 21 | 251 | | |
| | | CS 22 | 402 | | |
| | | CS 23 | 603 | | |
| **South Atlantic (Great calcite belt 1, MV1101) (5,048 m)** | | GCM120 | 25 | | |
| | | GCM121 | 62 | 75 | 3.12 |
| | | GCM122 | 112 | 247 | 0.66 |
| | | GCM123 | 162 | 266 | 0.46 |
| | | GCM124 | 300 | 362 | 0.47 |
| | | GCM125 | 500 | 276 | 0.21 |
| | | GCM126 | 750 | | | |
| | | GCM127 | 1,000 | 276 | 0.11 |
| **South Indian Ocean (great calcite belt 2, RR1202) (1,310 m)** | | GCM 199 | 20 | | |
| | | GCM 198 | 90 | | |
| | | GCM 197 | 125 | | |
| | | GCM 196 | 160 | | |
| | | GCM 195 | 200 | | |
| | | GCM 194 | 300 | | |
| | | GCM 193 | 500 | | |
| | | GCM 192 | 800 | | |
| **South Pacific Ocean (Antarctic sector, SEAFARERS) (1,887 m)** | | NBP 1016 | 50 | 26 | 3.66 |
| | | NBP 1017 | 100 | 99 | 0.61 |
| | | NBP 1018 | 150 | 197 | 0.29 |
| | | NBP 1019 | 250 | 279 | 0.18 |
| | | NBP 1020 | 400 | 256 | 0.13 |
| | | NBP 1021 | 600 | 291 | 0.10 |

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TABLE 1 | Analyzed samples for this study.

| Ocean (bottom depth) | Coordinates | Sample | Depth (m) | p[Ba] (pM) | POC (μM) |
|----------------------|-------------|--------|-----------|------------|----------|
| East Pacific (C-SNOW cruise) | | | | | |
| St 1 — Santa Barbara basin (1,929 m) | | CS 1 | 30 | | |
| | | CS 2 | 107 | | |
| | | CS 3 | 160 | | |
| | | CS 4 | 267 | | |
| | | CS 5 | 535 | | |
| | | CS 6 | 59 | | |
| | | CS 7 | 70 | | |
| | | CS 8 | 99 | | |
| | | CS 9 | 149 | | |
| | | CS 10 | 248 | | |
| (1,927 m) | | CS 12 | | | |
| | | CS 13 | 55 | | |
| | | CS 14 | 101 | | |
| | | CS 15 | 151 | | |
| | | CS 16 | 251 | | |
| | | CS 17 | 503 | | |
| | | CS 18 | | | |
| | | CS 19 | 25 | | |
| | | CS 20 | 151 | | |
| | | CS 21 | 251 | | |
| | | CS 22 | 402 | | |
| | | CS 23 | 603 | | |
| South Atlantic (Great calcite belt 1, MV1101) (5,048 m) | | GCM120 | 25 | | |
| | | GCM121 | 62 | 75 | 3.12 |
| | | GCM122 | 112 | 247 | 0.66 |
| | | GCM123 | 162 | 266 | 0.46 |
| | | GCM124 | 300 | 362 | 0.47 |
| | | GCM125 | 500 | 276 | 0.21 |
| | | GCM126 | 750 | | | |
| | | GCM127 | 1,000 | 276 | 0.11 |
| South Indian Ocean (great calcite belt 2, RR1202) (1,310 m) | | GCM 199 | 20 | | |
| | | GCM 198 | 90 | | |
| | | GCM 197 | 125 | | |
| | | GCM 196 | 160 | | |
| | | GCM 195 | 200 | | |
| | | GCM 194 | 300 | | |
| | | GCM 193 | 500 | | |
| | | GCM 192 | 800 | | |
| South Pacific Ocean (Antarctic sector, SEAFARERS) (1,887 m) | | NBP 1016 | 50 | 26 | 3.66 |
| | | NBP 1017 | 100 | 99 | 0.61 |
| | | NBP 1018 | 150 | 197 | 0.29 |
| | | NBP 1019 | 250 | 279 | 0.18 |
| | | NBP 1020 | 400 | 256 | 0.13 |
| | | NBP 1021 | 600 | 291 | 0.10 |
### TABLE 2 | Dissolved Ba, temperature, and calculated profiles of dissolved $\Omega_{\text{barite}}$ across depth in nearby stations to those analyzed in this work (co-located dissolved samples were generally not available for our study).

#### East Pacific (close to C-SNOW St. 1)

| Cruise   | GEOSECS Test | Station  | Test | Location  | 28.483°N, 121.633°W | Collected | September 1969 | Citation  | Wolgemuth and Broecker, 1970 |
|----------|--------------|----------|------|-----------|---------------------|-----------|----------------|-----------|-------------------------------|
| Depth (m) | [Ba] (nM)    | Temperature (°C) | $\Omega_{\text{barite}}$ |          |
| 1        | 50           | 18.07    | 0.29 |          |
| 30       | 43           | 17.85    | 0.25 |          |
| 150      | 50           | 11.66    | 0.39 |          |
| 255      | 57           | 8.71     | 0.50 |          |
| 400      | 67           | 6.9      | 0.65 |          |
| 700      | 85           | 5.07     | 0.89 |          |
| 1,000    | 93           | 4.01     | 1.02 |          |

#### Northeast Pacific (close to C-SNOW St. 2)

| Cruise   | KN195-08 | Station  | SAFe | Location  | 30°N, 140°W | Collected | May 2009 | Citation  | Geyman et al., 2019 |
|----------|----------|----------|------|-----------|-------------|-----------|----------|-----------|---------------------|
| Depth (m) | [Ba] (nM) | Temperature (°C) | $\Omega_{\text{barite}}$ |          |
| 25       | 35.0      | 19.01    | 0.20 |          |
| 75       | 37.8      | 18.20    | 0.22 |          |
| 110      | 35.0      | 18.18    | 0.20 |          |
| 150      | 35.3      | 16.76    | 0.22 |          |
| 200      | 37.1      | 12.81    | 0.27 |          |
| 250      | 37.5      | 11.23    | 0.30 |          |
| 300      | 39.4      | 9.92     | 0.33 |          |
| 350      | 43.5      | 8.85     | 0.38 |          |
| 400      | 48.9      | 7.95     | 0.45 |          |
| 500      | 59.4      | 6.32     | 0.59 |          |
| 600      | 68.3      | 5.19     | 0.71 |          |
| 700      | 77.9      | 4.64     | 0.83 |          |
| 850      | 89.4      | 4.08     | 0.98 |          |
| 1,000    | 98.1      | 3.72     | 1.10 |          |

#### South Atlantic (close to MV1101 St. 117)

| Cruise   | D357 (GA10E) | Station  | 3 | Location  | 36°27.6′S, 13°23.4′E | Collected | October 2010 | Citation  | Bates et al., 2017 |
|----------|--------------|----------|---|-----------|---------------------|-----------|----------------|-----------|---------------------|
| Depth (m) | [Ba] (nM)    | Temperature (°C) | $\Omega_{\text{barite}}$ |          |
| 5        | 43.0         | 12.08    | 0.32 |          |
| 23       | 42.4         | 12.07    | 0.32 |          |
| 47       | 42.6         | 12.06    | 0.32 |          |
| 97       | 44.5         | 11.19    | 0.36 |          |
| 196      | 43.3         | 10.8     | 0.34 |          |
| 395      | 48.0         | 8.17     | 0.44 |          |
| 594      | 56.1         | 5.22     | 0.59 |          |
| 999      | 70.6         | 3.64     | 0.79 |          |

(Continued in next column)

#### South Indian Ocean (close to RR1202 St. 63)

| Cruise   | INDIGO 1 | Station  | 18 | Location  | 45°09′S, 72°20′E | Collected | March 1985 | Citation  | Jeandel et al., 1996 |
|----------|----------|----------|----|-----------|-----------------|-----------|------------|-----------|---------------------|
| Depth (m) | [Ba] (nM) | Temperature (°C) | $\Omega_{\text{barite}}$ |          |
| 99       | 52.7      | 8.84     | 0.47 |          |
| 124      | 53.3      | 7.78     | 0.49 |          |
| 152      | 54.1      | 7.24     | 0.51 |          |
| 197      | 55.9      | 6.32     | 0.55 |          |
| 295      | 54.9      | 6.12     | 0.55 |          |
| 397      | 58.5      | 5.00     | 0.62 |          |
| 792      | 65.8      | 3.07     | 0.76 |          |
| 1,039    | 68.6      | 2.58     | 0.81 |          |

#### South Indian Ocean (close to RR1202 St. 63)

| Cruise   | INDIGO 3 | Station  | 90 | Location  | 55°01′S, 31°13′E | Collected | June 1987 | Citation  | Jeandel et al., 1996 |
|----------|----------|----------|----|-----------|-----------------|-----------|------------|-----------|---------------------|
| Depth (m) | [Ba] (nM) | Temperature (°C) | $\Omega_{\text{barite}}$ |          |
| 52       | 76.6      | 2.06     | 0.93 |          |
| 76       | 76.1      | 1.44     | 0.96 |          |
| 100      | 77.7      | 1.22     | 0.98 |          |
| 151      | 77.6      | 0.95     | 0.98 |          |
| 198      | 77.6      | 1.10     | 0.98 |          |
| 303      | 80.8      | 1.91     | 0.98 |          |
| 400      | 82.2      | 1.70     | 1.00 |          |
| 496      | 81.6      | 1.93     | 1.00 |          |
| 745      | 88.1      | 1.82     | 1.12 |          |
| 891      | 87.2      | 1.79     | 1.07 |          |
| 1,289    | 91.0      | 1.45     | 1.15 |          |

#### Southern Ocean (close to NBP1101 St. 14)

| Cruise   | GEOSECS | Station  | 287 | Location  | −69.3°S, 186.5°E | Collected | February 1974 | Citation  | Ku et al., 1976 |
|----------|---------|----------|-----|-----------|-----------------|-----------|---------------|-----------|---------------|
| Depth (m) | [Ba] (nM) | Temperature (°C) | $\Omega_{\text{barite}}$ |          |
| 2        | 77.8      | −1.07    | 1.10 |          |
| 21       | 77        | −1.49    | 1.02 |          |
| 41       | 78.2      | −1.40    | 1.12 |          |
| 81       | 78.1      | −1.65    | 1.12 |          |
| 151      | 79.8      | −0.31    | 1.07 |          |
| 201      | 83.1      | 0.87     | 1.07 |          |
| 272      | 84.7      | 1.43     | 1.05 |          |
| 352      | 85.7      | 1.43     | 1.07 |          |
| 449      | 92.6      | 1.37     | 1.15 |          |
| 598      | 89.8      | 1.27     | 1.12 |          |
| 797      | 92        | 1.12     | 1.18 |          |
| 996      | 94.5      | 0.97     | 1.20 |          |

(Continued in next column)
samples for analysis of dissolved Ba and sulfate were generally not available for our study, we estimated \( \Omega_{\text{barite}} \) from nearby stations with reliable published Ba concentration depth profiles (see Table 2 for details and Figure 1 for locations). Calculations were performed using PHREEQC version 3 (Parkhurst and Appelo, 2013). Values of \( \Omega_{\text{barite}} \) were computed for each sample based on input parameters of in situ temperature, d [Ba], pressure (estimated from depth). Both pH and salinity were prescribed in all calculations at 8.1 and 35, respectively. The major ion composition of seawater in the calculations was based on that reported by Kester et al. (1967). We believe that using a fixed salinity in our calculations is a reasonable assumption given the relatively minor effect this property has on \( \Omega_{\text{barite}} \) over the range of salinities encountered in open ocean seawater.

**Electron Microscopy Observations**

Quartz fiber filters (Whatman QMA) have been used for scanning electron microscopy (SEM) observation and analyses. Representative filter pieces were coated with

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**FIGURE 2** | SEM photographs showing representative examples of barite from the Pacific sector. Samples are indicated in Table 1. Both secondary electron and in backscattered electron (BSE) mode at 30 kV are shown in each sample. (A) CS4 (267 m), (B) CS9 (99 m), (C) CS10 (149 m), (D) CS20 (151 m), (E) CS 17 (603 m), (F) (CS 23) and (G) (CS 5) correspond to the deepest filter samples obtained at this ocean sector, 603 and 535 m, respectively.
carbon for observation under the SEM using an AURIGA FIB-FESEM Carl Zeiss SMT microscope equipped with an energy dispersive X-ray (EDX) detector system (Centre for Scientific Instrumentation, University of Granada). Filter pieces were grounded in an agate mortar and then dispersed in ethanol by sonication for approximately 3 min. Particulate matter released from the filter was deposited on carbon-film-coated copper grids for high-resolution transmission electron microscopy (HRTEM) observation by using a FEI TITAN G2 60–300 microscope with a high brightness electron gun (X-FEG) operated at 300 kV and equipped with a Cs image corrector CEOS (Centre for Scientific Instrumentation, University of Granada). For analytical electron microscopy (AEM), a SUPER-X silicon-drift windowless EDX detector was used. EDX maps and selected area electron diffraction (SAED) patterns were also collected on barite particles for crystallographic characterization and for determining major constituents composition.

RESULTS
Barite particles were observed in all the analyzed samples from all the locations and water depths (Figures 2–6). Figure 2 shows representative examples of barite particles from the Coastal East Pacific water column. Barite shows typical rounded to oval morphologies ranging in size from nanometers to a few microns, and it is always associated with organic material, which in many cases has EPS-like morphology (Figures 2A,C,E). Aggregates of barite grains are commonly observed with grains of different sizes ranging from less than 100 nm to a few hundred nm (Figure 2B). Barite is also observed as individual barite grains of micron size (Figure 2D) present at all depth analyzed. Figures 2F,G show examples of the deepest samples analyzed at 603 and 535 m water depth at these stations.

Figure 3 shows examples from the South Atlantic sector demonstrating barite associated with organic aggregates, which in some cases show EPS-like morphologies (Figure 2D). SEM
observations confirm the abundance of barite throughout this water column including the deepest sample at 1,000 m water depth (Figure 3G) and demonstrate that barite is particularly abundant in the intermediate mesopelagic zone (Figures 3E,F; sample GCM 125, 500 m). In the Indian sector of the Southern Ocean, a profile down to 800 m also shows higher barite abundance at depths corresponding to the upper mesopelagic zone (Figure 7). Crystals of different sizes are observed (Figure 4) as well as barite in organic aggregates (Figures 4D,E) at all depths. In the Antarctic sector of the South Pacific Ocean, barite shows similar morphologies and organic association as at the other sites (Figure 5). The particulate barite abundance at this station is slightly lower than at the studied station in the South Atlantic sector (Figure 7).

Electron microscopy observations demonstrate that morphology and composition of the barite particles are similar at all the studied ocean sectors and across depth in each station. The composition of the analyzed barite grains is similar to that previously reported in Martinez Ruiz et al. (2019), hence the EDX spectra are not shown in this work. EDX analyses show the expected barite composition, and in some cases, some barite grains also contain appreciable amounts of Sr and P. Although some variability among sites and depths in the P and Sr content is seen, a clear quantitative pattern of vertical or spatial variability in the content of P and Sr cannot be established with the EDX available data. However, qualitatively, at the South Atlantic and South Indian ocean stations, the number of barite grains enriched in Sr and P generally decreased with depth, suggesting that barite grains are more enriched in these elements at shallow depths. Acantharia shells have also been observed at the shallow depths in several stations (South Atlantic and South Indian oceans and East Pacific). It is important to note that the large Si peaks in EDX analyses from the quartz filter substrate overlap with the Sr L alpha (1.806 keV) peak, which may mask Sr when it is not in high enough abundance. However, at high concentrations Sr is easily detectable by the Sr K alpha 1 (14.165 keV) peak, and this has been carefully checked in SEM and HRTEM EDX spectra. As reported by Martinez Ruiz et al. (2019), Sr is clearly present in some of the analyzed barite grains but in other cases it is not detectable likely because of relatively low concentrations resulting in peaks.

FIGURE 4 | SEM photographs showing representative examples of barite from the Indian sector of the Southern Ocean. Samples are indicated in Table 1. Both secondary electron and in backscattered electron (BSE) mode at 30 kV are shown in each sample. (A) GCM 199 (20 m), (B) GCM 197 (125 m), (C) GCM 193 (500 m). (D) and (E) GCM 194 (300 m) show a detail of the different size of barite grains formed within organic aggregates.
obscured by the EDX Si spectra. HRTEM analyses also allow for Sr detection and indeed it is observed in many of the analyzed particles as shown in Figure 6. EDX maps also show that Sr content is variable at the nanometer scale and may vary even within the same particle (Martinez Ruiz et al., 2019).

Composition and crystallographic characteristics have been obtained by HRTEM. Representative examples are included in Figure 6. The analyses of organic aggregates indicate that barite crystallization starts with nanometer-sized amorphous precursors either P-rich (Martinez Ruiz et al., 2019) or with a barite-like composition and that the composition is variable even at this nano scale (Figures 6A–F). The amorphous nature of some of these particles is supported by SAED images and HRTEM diffraction data (Figure 6A). These analyses show crystals in which d-spaces corresponding to barite are clearly recognized (Figure 6G) as well as particles without a clear crystalline organization. High Angle Annular Dark Field (HAADF) STEM images and corresponding EDX map also demonstrate the high concentration of P and Sr in some of the barite grains (Figures 6C–E).

Particulate organic carbon (POC) and particulate Ba (pBa) profiles from the South Atlantic (Great Calcite Belt) Station 117 and the South Pacific Ocean (Antarctic sector) are shown in Figure 7. POC profiles are similar at both sites and show the typical POC profile with export out of the euphotic zone. At the Great Calcite Belt, Rosengard et al. (2015) argued that in this ocean region diatom-rich communities produce large and labile POC aggregates, which result in intense mineralization in the mesopelagic zone. The pBa profiles also support a significant increase in barite abundance at about 300 m, decreasing above and below that depth. At the Antarctic sector, the pBa profile show barite down to the deepest sample (600 m) analyzed at this station. Samples recovered from the top 1,000 m at low latitude sub/tropical locations exhibit undersaturation ($\Omega_{\text{barite}} < 1$) in the epipelagic and upper mesopelagic zones (e.g., east Pacific, southeast Atlantic). In contrast, water samples from the high-latitude Southern Ocean are generally close to saturation ($\Omega_{\text{barite}} = 1$) or even slightly supersaturated ($\Omega_{\text{barite}} > 1$), consistent with previous studies (e.g., Monnin et al., 1999; Rushdi et al., 2000).
FIGURE 6 | (A) High Angle Annular Dark Field (HAADF) STEM image and corresponding EDX maps showing the distribution of S, Ba and Ca in an organic aggregate from sample GCM 195 (Indian sector of the Southern Ocean). In the upper left corner, a selected area electron diffraction (SAED) image obtained by HRTEM is included to show the poor crystallinity of the nanometer-sized particles with barite composition; (B) representative spectrum (indicated in the HAADF image as square 2) showing the barite composition of nanometer-sized particles; (C), (D), and (E) show HAADF-STEM images and corresponding EDX maps of P (D) and Sr, S, and Ba (E) in a barite particle from NBP 1019 sample; (F) representative EDX spectrum from the barite particle shown in photograph c. Analyzed area corresponds to square 2 in that photograph, the Sr peak shows its enrichment in this grain; (G) shows a lattice-fringe image from sample GCM 124 in which d-spaces characteristic of barite are indicated: 3.77 Å (201), 3.57 Å (002), 2.83 Å (112), and 2.72 Å (020). In upper left side, a HAADF-STEM image and corresponding EDX maps (S and Ba) of the analyzed particle are included.
DISCUSSION

Barite Distribution in the Open Ocean

To date, a large body of work from multiple oceanographic expeditions and sampling location has provided ample datasets of dBa and non-lithogenic pBa distribution in the ocean water column. In general, the GEOTRACES-era datasets exhibit similar depth-dependent patterns in pBa to those shown here; however, this study also adds novel results regarding the mechanisms behind these distributions. It has been demonstrated that barite abundance shows significant spatial differences mostly related to productivity. Also, significant differences are recognized with depth in the water column since processes involved in precipitation are occurring at certain depths and the barite may dissolve deeper in the water column. Since pBa is closely correlated with the flux of organic carbon, it is enriched in the mesopelagic zone and typically shows a maximum abundance at intermediate depths (200–600 m) (e.g., Dehairs et al., 1980; Bishop, 1988; Dehairs et al., 1991; Dymond and Collier, 1996; Dehairs et al., 1997; Dehairs et al., 2008; Stenberg et al., 2008; Jacquet et al., 2011; Planchon et al., 2013; Lemaître et al., 2018; Conte et al., 2019). Overall, vertical pBa profiles are similar to those of calculated oxygen consumption rates, which supports the link between organic matter degradation and barite formation (e.g., Dehairs et al., 1997). Importantly, pBa has been correlated with rates of microbial degradation of organic matter, which further support the link to oxygen consumption and carbon respiration. Barium isotopes also support barite formation at mesopelagic depths as demonstrated by enrichment in the isotopically-heavy Ba in seawater (138Ba) and depletion of the lighter Ba (134Ba) due to the preferential incorporation of the lighter Ba isotopes in barite (Horner et al., 2015; Hsieh and Henderson, 2017; Bates et al., 2017; Bridgestock et al., 2018). Indeed a local maximum in δ138Ba at depths between 200 and 600 m in diverse ocean basins indicates that barite precipitation mostly occurs at these depths (Horner et al., 2015; Bates et al., 2017). Our pBa data from the two analyzed stations at the South Atlantic (Great Calcite Belt) and South Pacific (Antarctic sector) stations are also consistent with the idea of enhanced barite formation at this depth the mesopelagic zone. Profiles show higher pBa concentrations below 200 m and down to 600 m with a maximum in at about 200–400 m. Though qualitative, SEM observations from all the analyzed stations similarly show higher barite abundance at these intermediate depths.

As discussed above, barite formation is linked to organic carbon mineralization and export production, however a notable spatial variability in the Ba:Corg ratio is found over ocean regions. For instance, sediment trap samples from the Western Atlantic have significantly lower Ba/Corg values than samples from the Pacific (e.g., Dymond and Collier, 1996). The causes for this spatial variability are still poorly understood. Differences in the Ba:Corg ratio have been related to the efficiency of mineralization of POC in the mesopelagic zone relative to the exported amount (e.g., Francois et al., 1995). Thus, using algorithms that relate Ba to carbon export may not be appropriate in regions of highly variable carbon flux. Moreover, barite formation may be affected by the rate at which particles sink, given that particles that are quickly removed from the water column by rapid sinking may reduce the likelihood for precipitation of particulate Ba phases (McManus et al., 2002).
This has also added uncertainty to sedimentary Ba interpretations and paleoproductivity reconstructions, particularly to the use of Ba content as a quantitative proxy for reconstructing productivity. In general, the poor understanding of barite distribution in the water column stems from our limited knowledge of the processes leading to barite saturation. A better knowledge of such processes may improve our ability to assess changes in past productivity. Results from experimental work and from the analyses of pBa phases in the water column have recently shed light into potential mechanisms leading to saturation and precipitation of barite in microenvironments within sinking particulate matter in the mesopelagic zone (Martinez-Ruiz et al., 2018; Martinez-Ruiz et al., 2019). Collectively this work emphasizes the role of biofilms as Ba-concentrating agents in a process that could be termed organo-mineralization. According to the Encyclopedia of Geobiology, organo-mineralization (Défarge, 2011) is a process of mineral formation mediated by organic matter (OM), independent of the living organisms which the OM derives from. The organic compounds may be excretion products or detached parts of living organisms, or relics and by-products of dead organisms that have been released into waters or incorporated into soils, sediments, or rocks. Our observations from suspended marine particulate matter agree with this process and further support previous findings. Our SEM and TEM observations at multiple stations also demonstrate that barite forms through a P-rich amorphous precursor phase, seen in the pBa composition, ranging from Ba-phosphate to Ba-sulfate consistent with the pBa composition, size, distribution and association to organic material, prominently showing association with EPS-like morphologies, which further supports EPS production is a major factor in promoting barite formation (Figures 2–5) as well as organo-mineralization as a common process for barite formation throughout the ocean. Considering that the relation of pBa with export production depends on microbial processes related to organic matter degradation, barite formation is therefore linked to the suite of complex processes involved in the ocean biological carbon pump. Specifically, the fraction of primary production that leaves the upper ocean and is exported to depth, defined as export production is the fraction that “fuels” barite formation. Export production depends on diverse factors such as phytoplankton and zooplankton community structures, the formation of aggregates, sinking by ballasting, and bacterial mineralization rates, the formation of aggregates, sinking by ballasting, and bacterial mineralization rates (e.g., Francois et al., 2002; Cavan et al., 2015; Belcher et al., 2016; Le Moigne et al., 2016) and these factors change in space and time. For example, Henson et al. (2019) demonstrated that low primary production and high export efficiency regimes tend to occur when macro-zooplankton and bacterial abundances are low in the surface ocean. Thus, a large fraction of primary production is exported, likely as intact cells or phytoplankton-based aggregates. In contrast, when macro-zooplankton and bacterial abundances in the surface ocean are high, the export efficiency decreases. These results support that the whole ecosystem structure, rather than just the phytoplankton community, play a major role in export efficiency (Dehairs et al., 1992). All these factors not only depend on seasonality but can also be very different at regional and global scales. Hence, appropriate knowledge of the processes involved in carbon export fluxes, the formation of organic aggregates, and particle sinking is required for assessing the relationship between productivity and Ba proxies, and proper interpretative care and caution are required for using Ba as a proxy for export production. As the relationship between export production and barite largely depends on microbial processes and EPS production, temperature, oxygen abundance, and the type of organisms involved may impact this relationship. Although the reason for Ba accumulation in bacteria and EPS or in other living organisms is not yet well understood, it is known that bioaccumulation of Ba occurs throughout the ocean. The nucleation and crystallization of barite results in the formation of a highly stable mineral that is hard to dissolve under oxygenated conditions, consequently a relatively large fraction of the particulate barite that forms reaches the sediments (Paytan and Kastner, 1996). Barite accumulation in the sediment would therefore represent a record of the combination of diverse processes including export productivity, organic matter degradation, bacterial activity, and EPS production. This complexity should be considered when interpreting temporal and spatial variability in the Ba:Corg ratios and in barite accumulation in marine sediments.

Role of Extracellular Polymeric Substances in Barite Precipitation

Understanding the microbial processes leading to the formation of the mineral barite in the oceanic water column is crucial to determining the utility of Ba proxies for paleo-productivity and paleo-chemistry reconstructions. How and why Ba associates with organic matter in microenvironments and how the rates of organic matter decomposition affect barite production are key questions that link primary productivity or export productivity to barite abundance in marine sediments. The important role that EPS and microbial cells may play in nucleation and crystallization in the ocean is still far from being well understood at the molecular scale. Even though the EPS in the ocean have been widely investigated, their role in mediating mineral precipitation remains mostly unknown, in particular as it pertains to barite formation. In seawater, these secretions facilitate attachment to surfaces leading to the formation of biofilms, organic colloids, and larger aggregations of cells (marine snow). Though difficult to measure accurately, EPS represent a significant portion of the bioavailable carbon pool in the ocean. These substances occur in a range of molecular sizes, with diverse physical and chemical properties, and their composition includes polysaccharides, proteins, lipids, and nucleic acids (Decho and Gutierrez, 2017). In general, the attachment of microbes to surfaces, or
to each other, provides higher environmental stability than being a free-living cell and may be favorable in the ocean (e.g., Flemming et al., 2016). The EPS matrix of biofilms provides a three-dimensional architecture framework (Decho, 2000) that is the building block of the aggregates suspended in the water column. These organic aggregates are known to be very rich in microbial communities with abundances up to two orders of magnitude higher than in the surrounding seawater environment (e.g., Alldredge et al., 1986; Herndl, 1988).

While both experimental work and observations in diverse natural environments have demonstrated that functional groups associated with EPS are able to bind different metal ions (e.g., Braissant et al., 2007; Tourney and Ngwenya, 2014), their precise role for the binding, trapping and concentrating metals in the open ocean has not been sufficiently investigated. For instance, in bioremediation, it has been demonstrated that the polyanionic nature of the EPS promotes the binding of heavy and toxic metal ions, and EPS use for detoxification of heavy metals is well known. Many examples have been described in the literature such as in the remediation of Cd, Cr, Pb, Ni, Cu, Al, and U (e.g., Beech and Cheung, 1993; Iyer et al., 2005; Bhaskar and Bhosle, 2006; Gerber et al., 2018). However, as many of these studies had commercial purposes, very few have addressed the ecological implications of marine EPS in metal biogeochemical cycles. Loaec et al. (1997, 1998) reported the heavy metal-binding capacity of EPS produced by hydrothermal vent bacteria and suggested that this could represent a survival strategy for the bacteria by reducing their exposure to toxic metals released from the hydrothermal vents. Major elements such as Na, Mg, Ca, K, Sr, and Si, have also been shown to be adsorbed by marine bacterial EPS (Gutierrez et al., 2008). Likewise, Fe uptake by EPS in eukaryotic phytoplankton has been investigated (Hassler et al., 2011; Gutierrez et al., 2012), and binding of Th to carboxylate, phosphate and sulfate groups in marine EPS has also been shown (Alvarado Quiroz et al., 2006). Nevertheless, the ecological implications of these binding processes are not well understood, and Ba has not been investigated yet in this regard. Thus, our data open an unexplored field and support the crucial role that EPS play not only for Ba bioaccumulation but also for mineral precipitation in the ocean, with important implications for paleo-oceanographic reconstructions. Although further investigations are required to elucidate precise nucleation and crystal growth processes, the available data strongly support that biofilm matrix is crucial for metal precipitation in the ocean. In this dynamic environment, with abundant microbial cells, polysaccharides and water, together with excreted cellular products (e.g., Sutherland, 2001), functional groups including sulfate and phosphate would contribute to the overall negative charge of the EPS, and these functional groups would interact with metals promoting precipitation.

**The Microbial Pump and Future Perspectives**

The use of barite as a proxy to gain insights into past microbial processes is a promising tool in paleoceanographic research. It is broadly known that microbial communities play a major role in biogeochemical cycles since they play a role both at the base of the oceanic food web and as decomposers (e.g., Falkowski et al., 2008; Robidart et al., 2019). Further knowledge of microbial productivity and structure of communities is required for predicting future marine ecosystem functions, and the impact of increasing environmental effects on ocean ecosystems. This is challenging at present because biogeochemical processes and microbial communities are very complex, but it is far more complex for the past because a record of microbial processes is not usually well preserved in marine sediments beyond the preservation of biomarkers and some minerals that form through direct or indirect association with microorganisms. Accordingly, although barite accumulation rates are closely correlated with carbon export to the deep ocean (Carter et al., 2020 and references therein), the occurrence of barite may also reflect Ba utilization in the surface ocean through microbial processes, EPS production and organic matter mineralization. Except in sulfate reducing environments, barite is well preserved in marine sediments, thus the presence of pelagic barite particles is an indication of past bacterial respiration processes. In fact, barite has been proposed as a good proxy reflecting average mineralization processes (e.g., Cardinal et al., 2005), which are in turn a major control in the global carbon cycle and atmospheric carbon sequestration (e.g., Cavan et al., 2017). Our observations at diverse ocean sectors and depths showing formation within organic aggregates commonly rich in EPS, further support that mineralization due to microbial respiration is responsible for barite formation and consequently barite is a bioindicator for such processes.

The occurrence of barite in marine particles may have additional effects that have so far not been thoroughly investigated. For example, the potential role of barite particles within marine snow in the ballasting and mineralization controls of carbon sedimentation. It has been demonstrated that particulate minerals, for instance eolian dust, can be incorporated into organic aggregates and act as ballast enhancing the marine carbon export hence the significant increase in the sinking velocities of aggregates (Van der Jagt et al., 2018). Although biological processes affecting the fragmentation and mineralization of large particles are the most important factors determining the POC profiles (e.g., Lam and Bishop, 2007), barite is a high-density mineral that could also affect export processes. Another important aspect that still requires further investigation is the distribution of barite at greater depths than the mesopelagic zone. To date most of the studies on barite distribution have focused on mesopelagic depths and little is known about distribution and potential precipitation or dissolution with depth since only very few data from deeper samples are available (Conte et al., 2019). Furthermore, in the bathypelagic ocean (depth >1,000 m), Archaea and Viruses are particularly important in the microbial loop, but they remain largely unexplored in deep waters, and interactions between microbes and minerals beyond bacterial precipitation is almost unknown, except some recent work on the role of viruses in carbonate precipitation (e.g., Lan et al., 2020; White III et al., 2020).
CONCLUSIONS

Comparisons of suspended marine particulate matter obtained from diverse ocean sectors indicate barite formation at intermediate depths (200–600 m) in the mesopelagic zone independent of barite saturation state. The formation of barite within organic aggregates in close association with EPS is a common process recognized in all the studied regions. Our results further confirm that microbial processes are mediating barite precipitation (organomineralization) within such aggregates as previously supported by experimental work showing that Ba binds to phosphate groups on cell surfaces and EPS in bacterial biofilms. Mineralogical and crystallographic characteristics of suspended barite particles in the ocean support the same crystallization path, from an amorphous P-rich phase to mineral barite. EPS play a crucial role in locally concentrating Ba and providing nucleation sites leading to saturation. The binding capacity of the functional groups associated with EPS, including phosphate groups, has been widely demonstrated in experimental conditions and diverse natural environments, and the interactions between Ba and the EPS is similarly occurring in the ocean leading to barite precipitation. The distribution of particulate Ba and Ba isotopes in the water column are consistent with this precipitation mechanism. Many processes are involved in barite precipitation including primary production, export production, organic matter degradation, bacterial respiration, EPS formation, aggregation and sinking, and all should be further investigated and taken into account when interpreting temporal and spatial variability in the Ba:Corg ratios and barite accumulation in sediments. In addition, EPS production by organisms other than bacteria, such as phytoplankton, may also play an important role in barite production. However, the ecological implications of these processes and interactions between diverse organisms have been poorly investigated. The strong link between organomineralization of pelagic barite and microbial processes could be used to gain insights into past microbial processes and the mineralization of pelagic barite and microbial processes could also be investigated. The binding capacity of the functional groups associated with EPS, including phosphate groups, has been widely demonstrated in experimental conditions and diverse natural environments, and the interactions between Ba and the EPS is similarly occurring in the ocean leading to barite precipitation. The distribution of particulate Ba and Ba isotopes in the water column are consistent with this precipitation mechanism. Many processes are involved in barite precipitation including primary production, export production, organic matter degradation, bacterial respiration, EPS formation, aggregation and sinking, and all should be further investigated and taken into account when interpreting temporal and spatial variability in the Ba:Corg ratios and barite accumulation in sediments. In addition, EPS production by organisms other than bacteria, such as phytoplankton, may also play an important role in barite production. However, the ecological implications of these processes and interactions between diverse organisms have been poorly investigated. The strong link between organomineralization of pelagic barite and microbial processes could be used to gain insights into past microbial processes and the mineralization of the microbial pump. This is of importance for reconstructing mineralization and microbial respiration, and their link to export production, which are key processes in the global carbon cycle and the ocean carbon sink.

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DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

All authors listed have made a substantial contribution to the work and approved it for publication. FM conceived and led this project and has written the manuscript with an active contribution by discussing the results and writing from AP, MK, and MG. MG and FJ also contributed to perform related culture experiments and to discuss microbiological aspects. PL provided the studied filter samples and corresponding information, and TH led the pBa analyses, and both contributed to the discussion. MA contributed TEM analytical data and corresponding discussion.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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