Origin of marine barite deposits: Sr and S isotope characterization

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ABSTRACT

Barite can precipitate in microenvironments in the water column (marine barite), from supersaturated pore fluids at the oxic-anoxic boundary within marine sediments and where Ba-rich pore fluids are expelled and come into contact with sulfate-rich seawater (diagenetic barite), or from hydrothermal solutions (hydrothermal barite). Barite is relatively resistant to alteration after burial and has been used in paleoceanographic studies to reconstruct seawater chemistry and productivity through time. For such applications it is very important to determine the origin of the barite used, because both diagenetic and hydrothermal barite deposits may not accurately record the open-ocean contemporaneous seawater chemistry and productivity. We show here that it is possible to distinguish between the different types of barite by using Sr and S isotopes along with crystal morphology and size characteristics.

Keywords: barite, sulfur isotopes, strontium isotopes, paleoceanography.

INTRODUCTION

Barite in marine sediments is frequently used as a paleoproductivity proxy (Schmitz, 1987; Dymond et al., 1992; Gingele and Dahmke, 1994; Paytan et al., 1996a; Dean et al., 1997) as well as to reconstruct the seawater Sr isotope curve (Paytan et al., 1993; Martin et al., 1995), to determine the S isotope ratio of marine sulfate (Cecile et al., 1983; Goodfellow and Jonasson, 1984; Strauss, 1997; Paytan et al., 1998), and to characterize Holocene sedimentation rates by using excess ²²⁶Ra decay (Paytan et al., 1996b; van Beek and Reyss, 2001; van Beek et al., 2002). In all these applications it is assumed that the barite crystals analyzed precipitated directly from seawater and thus recorded contemporaneous seawater productivity and chemistry. Barite microcrystals have been observed in the water column (Dehairs et al., 1980; Bishop, 1988), and there are indications that barite precipitates inorganically directly from seawater in microenvironments containing decaying organic matter and other biogenic remains (Bishop, 1988; Dehairs et al., 1990; Ganeshram and Francois, 2002). This authigenic marine barite (as defined here) precipitates as small crystals or aggregates, ranging in size from 0.5 to 5 µm (Bishop, 1988; Dehairs et al., 1980; Paytan et al., 1993).

Barite may also precipitate in association with submarine hydrothermal activity (hydrothermal barite) (Zierenberg et al., 1984; Lonsdale and Becker, 1985; Feely et al., 1987, 1990; Hannington and Scott, 1988; Peter and Scott, 1988; Kusakabe et al., 1990; Moore and Stakes, 1990) as well as diagenetically at the oxic-anoxic boundary within sediments in association with sulfate-reducing conditions (Bolze et al., 1973; Dean and Schreiber, 1977; Brumsack and Gieskes, 1983; Cecile et al., 1983; Breheret and Brumsack, 2000) or when Ba-rich fluids from seeps or along fractures are expelled into seawater (diagenetic barite) (Lonsdale, 1979; Dia et al., 1993; Torres et al., 1996a; Aquilina et al., 1997; Naehr et al., 2000). These nonbiogenic barite deposits do not necessarily reflect the contemporaneous seawater conditions (chemistry or productivity) and are therefore unsuitable for paleoceanographic studies.

Criteria to distinguish between barite crystals that form via those distinct pathways are therefore required in order to utilize this phase for paleoceanographic studies (Church, 1979). A combination of S and Sr isotope analyses and crystal-morphology characterization is suggested here to provide an indicator of the depositional environment of a given barite. Sr and/or S isotope analyses have been previously applied to describe mechanisms and conditions of barite formation; however, all previous studies were restricted to a single deposit or location (Dean and Schreiber, 1977; Hannington and Scott, 1988; Kusakabe et al., 1990; Torres et al., 1996b; Aquilina et al., 1997; Fu and Aharon, 1997; Aharon and Fu, 2000; Naehr et al., 2000; Breheret and Brumsack, 2000). Here we compare the isotope systematics and mineral-habit characteristics of a wide range of barite deposits and establish criteria to distinguish between different modes of barite formation.

ANALYTICAL METHODS

Barite was separated from other minerals in the samples by a sequential leaching procedure (Paytan et al., 1993). All samples were examined for purity by X-ray diffraction (XRD) and scanning electron microscopy (SEM) with energy-dispersive spectrometry (EDS), and only samples that contained >95% barite were used. S isotope analyses were done by continuous-flow mass spectrometry using a Carlo Erba NA 1500 elemental analyzer connected to a Micromass Isoprime mass spectrometer. Samples of 4-8 mg were introduced in tin boats with \sim 5 mg vanadium pentoxide mixed in with each sample. A commercial tank of SO₂ was used as a reference gas for δ^{34} S measurements, and results are reported relative to the Canyon Diablo troilite standard, with a standard deviation (2σ) of $\pm 0.3\%$. Barite samples were dissolved for Sr isotope analysis by using a resin chelation method (Paytan et al., 1993), and the Sr fraction was separated by standard ion-exchange chromatography. Values of ⁸⁷Sr/86Sr were determined on a VG-261 mass spectrometer. The external precision of the analyses is ± 0.00002 . All values were normalized to an ⁸⁶Sr/⁸⁸Sr ratio of 0.1194. During the period of analysis, the mean 86Sr/87Sr ratio of the standard NBS 987 was 0.71024.

RESULTS AND DISCUSSION

The S and Sr isotope ratios of barite samples collected from different oceanic environments are given in Table 1 and Figure 1. Barite

TABLE 1. Sr AND S ISOTOPE COMPOSITIONS OF MARINE BARITE
SAMPLES

Sample name	87Sr/86Sr	$\delta^{34}S$	Site and
		(%0)	sample
		(700)	description*
			description
Core tops#	0.709175	21.1	1
Sediment traps**	0.709170	20.9	2
JDFR##, sediment trap ER 270383	0.709178	20.8	3
JDER ^{##} sediment trap ER 270388	0 706 077	21.1	3
IDER## Clam Bed Site 2463-R7	0 704 460	20.7	4
IDER*** East Wall AI V 1924-6-20	0 705 210	18.2	5
IDER*** Casm Site ALV 2084-24-1	0.706.251	10.2	5
IDER*** ALV 1924-6-20	0.700.201	20.5	5
IDER*** ALV 2045-482 2450	0.704413	10.2	5
DED### ALV 2343-432,2430	0.705243	19.2	5
JDFR , ALV 2234-21	0.703 102	20.9	5
JDFR""", ALV 2234-23-1	0.707 141	19.7	5
JDFR*****, ALV 1461-5	0.704512	21.0	6
JDFR ^{****} , ALV 1459-8	0.705347	20.8	6
JDFR, Southern Site, ALV 1457	0.705518	20.9	/
Mid-Atlantic Ridge####, ALV 2602-3	0.705 310	21.2	8
Mid-Atlantic Ridge####, ALV 2606-3	0.705971	20.9	8
Mid-Atlantic Ridge####, ALV 2604-5	0.704 432	21.0	8
Mariana backarc 1830-R-1	0.704 172	21.2	9
Mariana backarc 1831-R1-C	0.705 460	21.0	9
Mariana backarc 1831-R2	0.703 991	21.1	9
Mariana backarc 1832-R2-C	0.704762	20.9	9
East Pacific Rise 21°N, 914-R1-A	0.704275	20.8	10
ODP Site 765C 34-1, 57-60 cm	0.710892	24.2	11
ODP Site 765C 34-2 118–121 cm	0 710 817	23.7	11
Peru Margin SO 78/177	0 709 939	50.6	12
Peru Margin NP 2/34	0 711 031	32.7	12
Peru Margin SO 78/180	0.710.621	45.2	12
Raia California, Ensonada, BC 1	0.710.021	28.0	12
Baja California, Ensenada, BC 2	0.700.001	20.0	13
Paja California, Ensenada, BC-2	0.700701	29.0	10
Daja California, Eriseriaua, DC-3	0.700010	29.0	13
San Clemente Basin, 355 A	0.708487	22.1	14
San Clemente Basin, 355 B	0.708269	23.0	14
San Clemente Basin, 355 C	0.708372	24.6	14
Guaymas Basin 1170-16	0.706037	24.2	15
Guaymas Basin 11/2-6	0.706251	25.7	15
Guaymas Basin 1173-1	0.706 165	26.4	15
Guaymas Basin 1173-11	0.705649	23.2	15
Monterey Canyon, tubeworm slump	0.708 182	26.7	N.D.
Monterey Canyon 36.6°N, 122.4°W	0.708 163	31.3	16
Monterey Canyon 36.6°N, 122.4°W	0.708215	27.1	16
Gulf of Mexico (barite crusts)	0.708 587	28.5	17
Gulf of Mexico (barite crusts)	0.708 421	59.5	17

*References for site and sample description: 1, Paytan et al. (1993, 1998); 2, Dymond and Collier (1988); 3, Dymond and Roth (1988); 4, Moore and Stakes (1990); 5, Reyes et al. (1995); 6, Shanks and Seyfried (1987); 7, Feely et al. (1987); 8, Tivey et al. (1995); 9, Kastner et al. (1987); 10, Zierenberg et al. (1984); 11, Gradstein et al. (1990); 12, Torres et al. (1996a) and Dia et al. (1993); 13, Legg (1980); 14, Lonsdale (1979); 15, Lonsdale and Becker (1985); 16, Naehr et al. (2000); 17, Fu and Aharon (1997).

#Average value of 30 core-top samples.

**Average for 9 sediment-trap samples from the Pacific Ocean.

##Juan de Fuca Ridge (JDFR) Endeavour Segment.

***JDFR Axial Seamount.

###JDFR Middle Valley.

*****JDFR South Explorer Ridge.

####Lucky Strike.

samples cluster into three groups depending on their origin. The first very tight cluster (represented by one green diamond in Fig. 1) includes barite samples separated from core-top sediments in the Pacific, Atlantic, and Indian Oceans (30 samples) and barite crystals separated from sediment-trap samples (9 samples). These barite samples have precipitated from seawater (marine barite) and record present-day Sr (0.709 17) and S (21.1‰) isotope ratios (as determined in our laboratory).

Because the seawater Sr and S isotope ratios have not remained constant over time, the combination of Sr and S isotope ratios representing contemporaneous seawater values is different for barite separated from sediments of different ages. Accordingly, the isotope ratios for any given barite sample of known age (independently derived) should be compared to the well-known seawater Sr and S isotope



Figure 1. Plot of Sr and S isotope values for barite samples separated from different depositional environments. Green diamonds (plotted as one point)—isotope values for all core-top and sedimenttrap samples (excluding Juan de Fuca Ridge [JDFR] trap) (i.e., marine barite). Blue circles—diagenetic barite samples. Red circles hydrothermal barite samples. Black dot within diamond—barite in <8 µm fraction of Juan de Fuca Ridge trap. Dotted lines pass through present-day seawater Sr and S isotope ratios. CDT—Canyon Diablo troilite (S isotope standard).

curves (Burke et al., 1982; Paytan et al., 1998), and deviation from the expected contemporaneous seawater values suggests nonseawater-column origin (not marine barite).

The second category (blue circles in Fig. 1) includes barite samples with S isotope ratios higher than contemporaneous seawater values. For example, a recent massive barite deposit dredged in Baja California (Lonsdale, 1979; Table 1) has a δ^{34} S isotope value of 29.0‰ and an Sr isotope ratio of 0.708701, clearly different from those expected for present-day seawater. These samples have precipitated from fluids that had some degree of sulfate loss due to bacterial sulfate reduction (diagenetic barite). Sulfate reduction leads to enrichment of the heavy S isotope $({}^{34}S)$ in the residual sulfate in these fluids (Harrison and Thode, 1958). Barite precipitation may occur within the sedimentary column when Ba-rich fluids (from barite dissolution by the sulfatereduction process or from continental sources) migrate by diffusion or advection toward sulfate-rich sections in the sediment, typically at oxic-anoxic fronts (Dean and Schreiber, 1977; Kastner et al., 1990; Elderfield et al., 1990). Alternatively, barite could form at the sedimentwater interface where these Ba-rich fluids are discharged through seeps or faults into sulfate-rich seawater, and barite saturation is exceeded (Torres et al., 1996a, 1996b; Naehr et al., 2000).

The Sr isotope ratio of these diagenetic barites depends on the Sr isotope ratio of the water at the site of precipitation. The Sr could have less ⁸⁷Sr than the Sr that is typical of contemporaneous seawater if the pore fluids have been modified by Sr from less radiogenic sources like older marine sediments or the oceanic crust (as in San Clemente Basin and Monterey Canyon); the Sr could have more ⁸⁷Sr relative to contemporaneous seawater if the pore-fluid Sr was derived from alteration of radiogenic terrigenous material in the sediment or meteoric water (Torres et al., 1996a; Kastner et al., 1990; Elderfield et al., 1990) (as in Peru Margin and Ocean Drilling Program Site 765C samples).

A third group is composed of barite samples (red circles in Fig. 1) characterized by Sr isotope ratios that are between the modern seawater value and the Sr isotope ratio of pure (mantle derived) hydrothermal fluids (estimated to be 0.703 50; Albarède et al., 1981). These hydrothermal barite samples precipitate from fluids influenced by hydrothermal processes. Nonradiogenic Sr as well as Ba are leached from



Figure 2. Scanning electron microscope images of barite crystals from different oceanic settings. A: Marine barite, core-top sediments, Pleiades expedition, core 77, 1.03°N, 119.55°W. B: Marine barite, Deep Sea Drilling Project Site 574C, 29.1 Ma. C: Marine barite, sediment trap, <8 μ m fraction from Juan de Fuca Ridge black smoker. D: Hydrothermal barite, Mariana backarc chimney 1831-R1-C. E: Hydrothermal barite, Mid-Atlantic Ridge, chimney at Lucky Strike, ALV 2602-3. F: Hydrothermal barite, sediment trap, >8 μ m fraction from Juan de Fuca Ridge black smoker. G: Diagenetic barite, Ocean Drilling Program Leg 132, Site 765C 34-1, 47–60 cm, ca. 112 Ma. H: Diagenetic barite, San Clemente Basin, sea cliff 355A.

the oceanic crust (basalt), and when the circulating fluids interact and mix with sulfate-rich seawater, barite may precipitate (Kusakabe et al., 1990). The S isotope ratio of these barite deposits is either equal to, or may be lower than, contemporaneous seawater depending on the relative fraction of sulfur of hydrothermal origin (H_2S oxidation) in the mixture (Hannington and Scott, 1988).

In samples separated from a sediment trap deployed at the base of a black-smoker plume on the Endeavour Segment of the Juan de Fuca Ridge (Dymond and Roth, 1988), two types of barite crystals were identified. About 20% of the particles were $<8 \,\mu\text{m}$, whereas 80% of the particles were $>8 \,\mu\text{m}$. The isotopic signatures of the sizeseparated fractions are distinct; the $<8 \,\mu\text{m}$ fraction recorded presentday seawater S and Sr isotope values, and the $>8 \,\mu\text{m}$ crystals fall into the hydrothermal barite cluster. Thus the respective origins of the barite crystals are indicated (see Table 1).

Barite samples collected from the Guaymas Basin may represent a combination of the two nonbiogenic formation processes (hydrothermal and diagenetic). The Ba and Sr source for these deposits originated most likely from hydrothermal solutions. However, in this sedimentcovered ridge system, the fluids are expelled into marine-sediment sections where sulfate reduction takes place, depleting the pore-water sulfate from the light S isotope and resulting in barite with S isotope ratios slightly greater than seawater (Elsgaard et al., 1994).

In addition to the characteristic isotopic signatures of barite deposited in the different marine settings already described, the size and morphology of barite crystals formed by those different precipitation modes are distinct. Figure 2 shows SEM micrographs of typical barite crystals separated from the different depositional environments. Marine barite crystals precipitated in the water column and extracted from sediment-trap samples (including the $< 8 \mu m$ barite fraction from the Juan de Fuca Ridge black smoker) or from marine sediments that have not undergone extensive sulfate reduction are smaller than 5 µm and are typically ellipsoidal in shape (Fig. 2, A-C). Hydrothermal barite crystals are larger, 20-70 µm, and are typically precipitated as crosscutting tabular crystals commonly forming rosettes (Fig. 2, D-F). The >8 µm barite crystals from the sediment trap above the Juan de Fuca Ridge, as expected, have morphological features typical of hydrothermal precipitates. Diagenetic barite crystals are also large (20-700 µm), flat, tabular-shaped crystals and appear as barite beds in the sedimentary column. Diagenetic barite crystals that precipitate at the sedimentwater interface form mounds of highly porous barite with the layered appearance of platy crystals that form diamond-shaped clusters (Fig. 2, G-H).

CONCLUSIONS

On the basis of crystal size and morphology and the Sr and S isotope ratios of barite deposits, it is possible to distinguish between the depositional environments and thus the origin of this mineral if the age of the barite sample is independently determined. Careful examination of barite samples used for paleoceanographic studies would eliminate any questions with respect to the authenticity of marine barite origin and its fidelity in recording the seawater characteristics (productivity and/or chemistry).

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