22. RED SEA EVAPORITES: A PETROGRAPHIC AND GEOCHEMICAL STUDY

Peter Stoffers, Laboratorium für Sedimentforschung, Universität Heidelberg, Germany
and
Robert Kühn, Kali-Forschungs-Institut der Kali u. Salz A.G., Hannover, Germany

INTRODUCTION

One of the major shipboard findings during Leg 23 drilling in the Red Sea was the presence of late Miocene evaporites at Sites 225, 227, and 228 (Figure 1).

Figure 1. Leg 23 drill sites in the Red Sea.

The top of the evaporite sequence correlates with a strong reflector (Reflector S) which has been mapped over much of the Red Sea (Ross et al., 1969, Phillips and Ross, 1970). This indicates that the Red Sea appears to be

underlain by late Miocene evaporites throughout most of its extent.

Miocene sediments, including evaporites, are known from a few outcrops along the coastal plains of the Gulf of Suez to lat 14°N (Sadek, 1959, cited in Friedman, 1972; Heybroek, 1965; Friedman, 1972). Along the length of the Red Sea, the presence of Miocene salt is indicated by seismic reflection studies (Lowell and Genik, 1972) and confirmed by drilling. The recently published data from deep exploratory wells (Ahmed, 1972) demonstrate the great thickness of clastics and evaporites which were deposited in the Red Sea depression during Miocene time.

The Red Sea evaporites are of the same age as the evaporites found by deep sea drilling (DSDP Leg 13) in the Mediterranean Sea. Therefore, Reflector S in the Red Sea is comparable to Reflector M in the Mediterranean. It is assumed that during Miocene time a connection between these two basins was established (Coleman, this volume) resulting in a similar origin for the evaporites deposited in the Red Sea and in the Mediterranean Sea. The origin of the Mediterranean evaporites has been discussed in great detail (Hsü et al., 1973; Nesteroff, 1973; Friedman, 1973).

The formation of evaporites may be interpreted by three different hypotheses.

1) Evaporation of a shallow restricted shelf sea or lagoon which receives inflows from the open ocean.
2) Evaporation of a deep-water basin which is separated from the open ocean by a shallow sill (Schmalz, 1969).
3) Evaporation of playas or salt lakes which are situated in desiccated deep basins isolated from the open ocean (Hsü et al., 1973).

The purpose of this study is to show whether one of these models might apply to the formation and deposition of the Red Sea evaporites. Therefore, a detailed petrographic and geochemical investigation was carried out.

PETROGRAPHIC DESCRIPTION

The stratigraphy of the evaporite sequence encountered at Sites 225, 227, and 228 is shown in Figure 2, which also indicates the mineralogy of the interbedded black shales. The photographed evaporite cores can be seen in the site report chapters for Sites 225, 227, and 228. Forty selected samples of the evaporite rocks from the Red Sea boreholes were studied in thin section under the petrographic microscope. All anhydrite samples were X-rayed in order to clarify the presence of gypsum, dolomite, calcite, and magnesite. The structure and texture of the rocks were described according to Maiklem et al. (1969).
Figure 2. Stratigraphy of the evaporites encountered at Sites 225, 227, 228 and the mineralogy of the interbedded black shales.
Site 225

Site 225 is situated 10 miles east of the Atlantic II Deep in a water depth of 1228 meters. Two hundred and thirty meters of sediment was continuously cored, including 53 meters of evaporites.

At the base of this hole (Samples 29-3, 0-150 cm; 29-2, 90-150 cm) white to gray rock salt was recovered. Bedding (3-7 cm apart) of cloudy and clear white halite occurred. Thin anhydrite layers (0.2 to 1 cm) are intercalated within these beds. The dip of the anhydrite layers ranges up to 15°. Thin section examination (29-3, 100-103 cm; Figure 3) reveals medium to coarse crystalline halite (2-3 mm in diameter). Fine polyhalite needles (2\(\text{CaSO}_4 \cdot \text{MgSO}_4 \cdot \text{K}_2\text{SO}_4 \cdot 2\text{H}_2\text{O}\) (0.75 mm in length, 0.05 mm in width) are stringlike interspersed in the pore spaces of halite crystals. Twinning of the polyhalite crystals is found quite frequently. In larger pore spaces the polyhalite occurs as densely felted needles (29-3, 100-103 cm; Figure 4). The cloudy appearance of the gray halite is due to scattered brine inclusions. The linear orientation of the inclusions often produced crystal structures known as "hopper" and "chevron" which indicate a primary origin (Dellwig, 1953; Wardlaw and Schwertner, 1966). These structures disappear on recrystallization resulting in more transparent larger crystals clear of opaque inclusions.

The layering of clear and cloudy salt with intercalated thin anhydrite bands is typical of the halite recovered on Leg 23 drilling in the Red Sea. No obvious cyclic relationship is observed between these features. The absence of a statistical correlation between clear salt, cloudy salt, and anhydrite layers is reported by Fuller and Porter (1969) from the Devonian and Mississippian salt deposits of Alberta and North Dakota and by Dellwig (1968). These authors explain the alteration of clear and cloudy salt by changes in the rate of evaporation which reflect changes in climatic conditions. The thickness of the halite laminae is determined by seasonal dilution (removal and recrystallization), whereas the thickness of the anhydrite laminae is determined by seasonal recombination.

The halite sequence is topped by a small interval of slightly layered anhydrite (29-2, 87-90 cm) followed by typically nodular anhydrite (29-2, 6-81 cm). Black shales are intercalated within the anhydrite (29-2, 81-87 cm; 29-1, 60-67 cm). They are predominantly composed of montmorenilte and smaller amounts of cristobalite. These shales are thought to have originated from interbedded volcanic material. The anhydrite structure is described best as distorted nodular mosaic (29-2, 10-20 cm; Figure 5). The white nodules (up to 3 cm in diameter) are separated by thin greenish seams. Under the petrographic microscope the nodules appear to be a randomly arranged combination of microcrystalline and small lath-shaped crystals (0.05-0.1 mm; Figure 6). The nodules are bound by dark brownish streaks of opaque, probably organic-rich, material and very fine grained dolomite (~0.003 mm) which were separated from the matrix during growth or deformation of the anhydrite nodules.

Core 28 and the lower part of Core 27, Section 2 consist of bedded clear white and dusty gray halite (about 2 to 4 cm apart) which is intercalated by thin anhydrite layers (0.5-1 cm). The anhydrite layers show dips between 10-25°. The thin sections (28-3, 95-105 cm; 27-2, 110-112 cm) show medium to coarsely crystalline halite (2 to 4 mm) with partly mosaic structure (Figures 7, 8). Within the halite there are thin layers of needle-like anhydrite which locally forms rounded aggregates. In the dusty gray halite-oriented liquid, inclusions are very abundant.

The halite is topped by a small interval of light gray, slightly laminated anhydrite (27-2, 69-87 cm) followed by distorted nodular to bedded nodular mosaic anhydrite (27-2, 0-60 cm; 27-1, 90-150 cm; Figure 9). Petrographic microscope examination (27-1, 119-122 cm) shows that the matrix is a fine crystalline anhydrite. Besides fine-grained dolomite and streaks of opaque, probably organic-rich, material, clusters of coarse crystalline anhydrite occur which sometimes display sheath-like to stellate patterns (Figure 10). The top of Core 27, Section 1 consists of a more laminated anhydrite. A thin layer of greenish gray fine-grained dolomite (27-2, 60-69 cm) is intercalated in the anhydrite, indicating a regressive stage of evaporation.

Core 26 is mainly composed of white anhydrite with bedded nodular mosaic structure (26-2, 133-136 cm; Figure 11). The nodules are separated by black opaque, probably organic-rich, matter and small amounts of carbonate material. The matrix again is a very fine crystalline anhydrite of aligned felted texture (~0.005 mm). Within this ground mass there are porphyrotopes of coarse crystalline anhydrite (26-1, 140-144 cm; Figure 12). These aggregates are thought to have formed by diagenetic metasblaste.

The anhydrite is topped by a black shale sequence (26-1, 1-34 cm). The thin section (26-1, 13-16 cm) reveals the presence of slightly laminated algal oncolites (Figure 13 a, b, c). The oncolites consist of very fine-grained dolomite (0.005-0.01 mm) impregnated with opaque material, probably rich in organic matter, and some pyrite. The cavities within the oncolites are often filled with secondary anhydrite, sometimes with chaledony.

A finely laminated massive anhydrite of alternating medium gray and white layers is found in the lower section of Core 25 (137-150 cm). The thickness of the individual layers is about 1-1.5 mm. The upper part of the core consists of dark shales. X-ray analyses show that the shales are mainly composed of dolomite (up to 60%), layer silicates, pyargorskite, analcite, quartz, feldspar, and some pyrite. The dominant clay mineral is an illite-montmorillonite mixed layer. The formation of the black dolomite shales is related to regressive evaporation caused by a strong dilution of the brine.

The top 10 cm of Core 25 and the major part of Core 24 (67-150 cm) consist of wavy laminated massive anhydrite (Figure 14). The laminations are described best as stromatolites. They are composed of alternating white 2-4 mm and medium gray 0.5-1 mm thick layers. The thin section (24-1, 70-74 cm; Figure 15) shows very uniform fine crystalline lath-shaped anhydrite (~0.015 mm) in subparallel orientation. The darker layers consist of scattered micron sized opaque, probably organic, material and very fine grained dolomite.

A dark gray to black semilithified to lithified dolomitic silty claystone (Core 24-1, 0-70 cm; Cores 23, 22, 21) tops
the evaporite section of Site 225. The increasing marine fauna found in the upper cores of this sequence indicates the change to open marine conditions.

Site 227

Site 227 is situated about two miles east of the edge of the Atlantis II Deep in a water depth of 1795 meters and 359 meters of Quaternary to late Miocene sediments were cored, including 133 meters of late Miocene and probably older anhydrite and halite.

The lowermost core (45-1, -2) consists of bedded alternating white and light gray rock salt. In general, the individual layers vary between 2 and 5 cm. Anhydrite layers 0.2 to 10 mm thick (2 to 6 cm apart) are intercalated within this halite sequence. The thin section (45-1, 112-117 cm; Figure 16) shows coarse crystalline halite (5-8 mm) interspersed with polyhalite needles (0.05-0.08 mm). Liquid inclusions are abundant within the gray halite forming typical chevron and hopper structure.

Anhydrite is found in Core 44, Sections 1 and 2. In the lower part of Section 2 (70-150 cm), the anhydrite is of distorted bedded nodular structure. The white nodules are surrounded by greenish gray dolomite seams. The top of Section 2 and the entire Section 1 consist of thin layers (1-2 mm) of massive whitish gray anhydrite interbedded with layers of medium gray material (~0.03 mm; Figure 17). Petrographic microscope examination (44-2, 65-70 cm; Figure 18) reveals very fine laminated anhydrite of aligned felted texture (0.03-0.06 mm). The lath-shaped crystals are oriented parallel to the bedding planes. The darker layers are composed of dolomite impregnated with organic matter and very fine grained pyrite.

The entire Core 43 (Sections 1 to 5) and Core 42 (Sections 1 to 3) are represented by irregularly bedded clear white and gray halite. Very distinct layers of anhydrite (~1.5 cm) interbedded within the halite occur in Sections 3, 4, and 5, whereas the anhydrite layers in the upper section of Core 43 and in Core 42 are only slightly indicated. Clear white halite dominates this upper sequence. The thin section (43-2, 20-25 cm; Figure 19) shows medium coarse crystallized halite (1-2 mm) of generally mosaic structure. The halite is interbedded by a 1.5 cm thick anhydrite layer. The subfreted anhydrite crystals (~0.07 mm) are in random orientation. Within the anhydrite there are dark stringers of opaque, probably organic-rich, material.

The lowermost 4 cm of Core 41, Section 1 is composed of bedded massive anhydrite topped by alternating white and gray halite (4-8 cm apart) with intercalated 2-10 mm thick anhydrite layers. Under the microscope (41-1, 105-110 cm; Figure 20), the halite is coarse crystalline and shows a mosaic structure. The microcrystalline anhydrite (0.02-0.05 mm) is often aligned with the halite particles. This concentration of anhydrite along the halite rims is caused by anhedral halite growth which seems to replace euhedral halite crystals.

Core 40, Sections 1 and 2 consist of black shales which were completely fractured by drilling. X-ray diffraction shows that the shales are composed, in general, of montmorillonite and smaller amounts of calcite, feldspar, quartz, clinoptilolite, analcite, pyrite, dolomite, gypsum, and anhydrite. The abundant montmorillonite and the presence of zeolites suggest a strong volcanic influence.

Core 39, Section 1 and Core 38, Section 1 are represented by massive anhydrite with a slight indication of bedding. The thin section (38-1; Figure 21) shows a lath-shaped anhydrite matrix interspersed with large fibrous or sheath-like anhydrite aggregates. The recrystallization of the anhydrite has caused the scattered dolomite and opaque material to migrate to the boundaries of the aggregates.

Anhydrite of bedded mosaic structure is found in Core 37, Sections 1 and 2. The thin section (37, CC; Figure 22) shows fine crystalline dolomite layers (0.005-0.02 mm) impregnated with opaque material. The dolomite layers (0.05-0.1 mm thick) are intercalated within the lath-shaped anhydrite matrix.
Figure 5. Nodular anhydrite (225-29-2, 10-20 cm). Scale bar represents 1 cm.

Figure 6. Nodules consisting of microcrystalline and small lath-shaped anhydrite (225-29-2, 10-20 cm). Scale bar represents 4 mm. Oriented thin section. Crossed nicols.

Figure 7. Medium to coarsely crystalline halite intercalated with thin anhydrite layers (225-28-3, 95-105 cm). Scale bar represents 4 mm.

Figure 8. Medium to coarsely crystalline halite with rounded anhydrite aggregates (225-27-2, 110-112 cm). Scale bar represents 4 mm.
Figure 9. Anhydrite of distorted nodular mosaic structure (225-27-1, 119-122 cm). Scale bar represents 2 cm.

Figure 10. Fine crystalline anhydrite matrix, dark gray semi-opaque, probably organic-rich, streaks, fine-grained dolomite, clusters of coarse crystalline anhydrite (225-27-1, 119-122 cm). Scale bar represents 4 mm. Crossed nicols.

Figure 11. Anhydrite of bedded nodular mosaic structure (225-26-2, 133-136 cm). Scale bar represents 2 cm.

Figure 12. Fine crystalline anhydrite of aligned felted texture, streaks of opaque material, aggregates of coarse crystalline anhydrite (225-26-1, 140-144 cm). Scale bar represents 4 mm. Thin section is oriented. Crossed nicols.
Figure 13. Oncolite structure (225-26-1): (a) scale bar represents 4 mm; partially crossed nicols; (b) partially crossed nicols; some cavities filled with anhydrite and chalcedony; scale bar represent 0.5 mm; and (c) fine grained dolomite impregnated with opaque, probably organic rich material; some anhydrite in the cavities; scale bar represents 0.1 mm.
Figure 14. Wavy laminated anhydrite (stromatolites) of alternating white anhydrite and dark layers composed of dolomite and opaque material (225-24-1, 80-83 cm). Scale bar represents 2 cm.

Figure 15. Fine crystalline anhydrite of aligned felted texture with alternating layers of opaque material interspersed with dolomite (225-24-1, 70-74 cm). Crossed nicols. Scale bar represents 4 mm.

Figure 16. Coarse crystalline halite interspersed with polyhalite needles, oriented liquid inclusions (227-43-1, 112-117 cm). Partially crossed nicols. Thin section is oriented. Scale bar represents 4 mm.

Figure 17. Bedded mosaic anhydrite with intercalated dark layers (227-44-2, 65-70 cm). Scale bar represents ~2 cm.
Figure 18. Fine laminated anhydrite of aligned felted texture. Dark layers consist of dolomite impregnated with opaque material (227-44-2, 65-70 cm). Crossed nicols. Thin section is oriented. Scale bar represents 4 mm.

Figure 19. Coarse crystalline halite with interbedded anhydrite layer (227-43-2, 20-25 cm). Partially crossed nicols. Thin section is oriented. Scale bar represents 4 mm.

Figure 20. Coarse crystalline halite with microcrystalline anhydrite aligned with the halite particles (227-41-1, 105-110 cm). Partially crossed nicols. Thin section is oriented. Scale bar represents 4 mm.

Figure 21. Lath-shaped anhydrite matrix interspersed with large fibrous or sheath-like anhydrite aggregates (227-38-1, 130-134 cm). Crossed nicols. Thin section is oriented. Scale bar represents 4 mm.
Interebed black shales occur within the anhydrites of Core 36 (36-3, 70-150 cm; 36-2, 95-150 cm). The shales were completely broken up by the drilling operation. X-ray examination reveals disordered cristobalite as the predominant constituent (>80%) besides traces of pyrolusite, zeolites, dolomite, quartz, feldspars, and pyrite. Nannofossils (Discosaster quinqueramus) and benthonic foraminifera are present in the core catcher of Core 36. Under the petrographic microscope, the microcrystalline cristobalite is finely laminated by opaque material probably rich in organic constituents (Figure 23). Scanning micrographs show typical silica spherules (5-10 µ) of disordered cristobalite. The spherules consist of irregularly oriented tubes and needles (Figures 24 a, b) described as lussatite (Berger and von Rad, 1972). The mineralogy of the black shales suggests volcanism as a source of the silica.

The anhydrite in Core 36, Section 3 has a massive structure with slight indications of bedding. Under the petrographic microscope (36-3; Figure 25), the anhydrite is of microcrystalline texture (~0.05 mm). Intergrown clusters of coarse recrystallized anhydrite and authigenic quartz crystals (up to 3 mm) are found within this fine-grained matrix. The quartz crystals often show twinning and frequently contain inclusions of anhydrite. These inclusions are oriented parallel to the crystal edges due to the growth of the quartz crystal (36-3; Figure 26). Quartz is generally common in saline deposits. It occurs partly as clastic grains and partly as an authigenic mineral with well-developed idiomorphic forms. According to Grimm (1962), quartz forms in saline sediments during an early stage of diagenesis. Idiomorphous quartz crystals with inclusions of anhydrite demonstrate that this anhydrite rock probably was of primary origin and not derived from gypsum by diagenesis. Due to the volume increase accompanying the transition to anhydrite, the inclusions of primary gypsum would have produced some fracturing. However, microscopic investigations show that the inclusions of anhydrite in the quartz completely fill the available space. No fractures could be observed.

Anhydrite of streaky structure is found in Core 36, Section 2 (Figure 27). Tails and pieces of greenish gray carbonate material are intercalated with dark gray anhydrite beds. The carbonate material was identified as magnesite by X-ray diffraction. Besides magnesite, there are traces of pyrite. The thin section (36-2; Figure 28) shows microcrystalline anhydrite (~0.05 mm) with slightly aligned orientation. The very fine crystalline magnesite (0.003-0.007 mm) occurs as clusters or streaks within the anhydrite. Core 36, Section 1 is also composed of streaky anhydrite. Petrographic microscope examination (36-1; Figure 29) reveals anhydrite crystals of aligned felted texture. Lumps of anhydrite (up to 10 mm) with sharp edges or straight sides are separated from one another by a magnesite matrix (0.007-0.02 mm). Some of these features might be deformed gypsum pseudomorphs (Borchert and Baier, 1953). The microcrystalline magnesite occurs as hypidiotopic to xenotopic grains. Frequently, magnesite forms ringlike aggregates which surround a single magnesite or pyrite grain. Ringlike magnesite aggregates within Zechstein anhydrites of Northern Germany have been described by Langbein (1961) and Kosmahl (1969). It is thought that these ringlike structures originated from recrystallization due to diagenesis.

A study of the available analyses of carbonate in salt rocks of Central Europe (Kühn, 1968) shows that along the margins of the basin, calcite, and more commonly, dolomite occurs, derived from relatively diluted brines; whereas, magnesite occurs exclusively in the sediments of the more highly concentrated brines of the interior of the basin. The distribution of carbonates is evidently related to the process of progressive concentration during which the heavier magnesium chloride solution accumulated in the deepest part of the basin. According to Link (1938, 1942), the dolomite and the magnesite have to be considered as early diagenetic.

The bottom of Core 35 consists of bedded massive light gray anhydrite interbedded with thin dark gray layers about 1 mm apart (Figure 30). Petrographic microscopic examination (35-5, 115-123 cm; Figure 31) shows microcrystalline anhydrite (~0.05 mm) interspersed with sheath-like anhydrite (1-2 mm). Dark layers (0.02-0.1 mm) are intercalated in the fine crystalline matrix and are composed of opaque, probably organic-rich, material and small dolomite grains (~0.004 mm). The anhydrite is topped by bedded alternating clear white and medium gray halite (Sections 1-5). The thickness of the individual beds, generally, is about 3 to 7 cm with the exception of the lower part of Section 5 (60-120 cm), which is represented by a sequence of medium gray halite. Thin brownish green anhydrite layers (0.5-1 cm) are intercalated in the halite (Figure 32) at intervals of 1 to 7 cm. In the medium gray halite of Section 5, these laminations are only 0.5 cm apart. The thin anhydrite layers show dips up to 60°. The thin section (35-3; 53-60 cm; Figure 33) reveals medium to coarse crystalline halite (~1 mm) with clusters and streaks of lath-shaped anhydrite (0.1-0.2 mm). Recrystallization of the halite has caused the anhydrite to concentrate along the crystal boundaries of the halite.

Core 34 (Sections 1 to 4), Core 33 (Sections 1 to 4), and Core 32 (Sections 1 to 5) consist of alternating clear white and medium gray halite with interbedded thin anhydrite layers (0.2 mm). In Cores 32 and 33, the anhydrite layers are only slightly indicated. Clear white halite predominates this sequence. Under the petrographic microscope (33-1, 80-84 cm; Figure 34), the medium to coarse crystalline halite (>1 mm) of mosaic structure is oriented parallel to the bedding plane. Polyhalite needles are found in clusters or stringers interspersed within the halite.

The halite sequence is topped by dark brown to black shales (Core 31, Section 1, 0-70 cm) which are broken up by the drilling operation. The extremely hard finely laminated black shale pieces found in the core catcher of Core 31 are brecciated and fractured (Figure 35). The cracks are filled with halite. According to X-ray diffraction, this black shale is predominantly composed of disordered cristobalite with traces of pyrite and quartz. Under the petrographic microscope (31, CC; Figures 36 a, b, c), angular fragments of cristobalite containing laminated streaks of organic material are cemented by halite. Silica of colloform texture occurs within the halite matrix and forms chains and rings which show undulatory extinction. The texture clearly indicates a colloidal origin. The 2-3 mm thick
Figure 22. Bedded mosaic anhydrite with thin layers of dolomite impregnated with opaque material (227-37, CC). Scale bar represents 0.1 mm.

Figure 23. Microcrystalline cristobalite finely laminated by opaque, probably organic rich, material and pyrite (227-36-2). Scale bar represents 0.1 mm.
Figure 24. Scanning micrographs of cristobalite spherules (227-36-2): (a) scale bar represents 10µ (micron), and (b) scale bar represents 2µ.

Figure 25. Microcrystalline anhydrite matrix, clusters of coarse crystalline anhydrite and authigenic quartz (227-36-3, 5-7 cm). Crossed nicols. Scale bar represents 4 mm.
Figure 26. Authigenic quartz with oriented anhydrite inclusions in fine crystalline anhydrite matrix (227-36-3). Scale bar represents 1 mm. Crossed nicols.

Figure 27. Streaky laminated anhydrite (227-36-2, 73-76 cm). Scale bar represents 2 cm.
Figure 28. Microcrystalline anhydrite with slightly aligned orientation intercalated with dark fine crystalline magnesite (227-36-2, 73-76 cm). Crossed nicols. Thin section is oriented. Scale bar represents 4 mm.

Figure 29. Anhydrite of aligned felted texture. The sharp edges of the anhydrite lumps suggest deformed gypsum pseudomorphs. Dark material is microcrystalline magnesite (227-36-1). Crossed nicols. Scale bar represents 4 mm.

Figure 30. Bedded massive anhydrite (227-35, CC). Scale bar represents 1 cm.

Figure 31. Microcrystalline anhydrite interbedded with dark layers composed of opaque material and small dolomite grains (227-35-5, 115-123 cm). Scale bar represents 4 mm.
fractured vein, which crosses perpendicular to the bedding plane of the shale, is filled with sheave-like partly radiating bundles of length slow fibers of quartzine and some megaquartz. The silica of this black shale might be originated from devitrification of volcanic glass or from the dissolution of opaline skeletons. The absence of volcanic alteration products such as clinoptilolite and montmorillonite suggests a biogenic source for the silica. Redeposition of the silica formed disordered cristobalite. Quartzine was precipitated in veins. Due to some kind of tectonic movement, probably caused by salt flowage, the black shale was intensely brecciated. The halite, which filled the cracks, precipitated from the pore solution. Some of the silica was probably dissolved again because of the hypersaline environment and was redeposited within the halite in typically colloform texture. The thin section might demonstrate a diagenetic process leading from probably opaline skeletons via disordered cristobalite and quartzine to megaquartz. The significance of length slow chaledony in sedimentary rocks is discussed in great detail by Folk and Pittman (1971). According to these authors, the occurrence of quartzine is characteristic of a silification in highly basic environments. Zeolites, and palygorskite. The white chips found in the shale sequence are completely composed of well ordered palygorskite. Abundant nannofossils (Discocysta quinqueramus) and rare neritic benthic foraminifera, which occur in the black shale of Core 31, suggest fluctuations from evaporitic to marine conditions.

X-ray diffraction shows that the dark brown shale fragments in Core 31 consist mainly of dolomite with smaller amounts of layer silicates, quartz, feldspar, pyrite, zeolites, and palygorskite. The white chips found in the shale sequence are completely composed of well ordered palygorskite. Abundant nannofossils (Discocysta quinqueramus) and rare neritic benthic foraminifera, which occur in the black shale of Core 31, suggest fluctuations from evaporitic to marine conditions.

The lower part of Core 30, Section 2 (100-150 cm) is represented by white halite with thin grayish green anhydrite layers. The overlying rock is bedded massive anhydrite of alternating white (0.2-0.6 mm) and dark layers (0.02-0.05 mm). Under the petrographic microscope (30-2, 43-48 cm; Figure 37), the anhydrite is of aligned felted texture (0.015-0.2 mm). The dark brown layers consist of dolomite and micron-sized opaque material.

The anhydrite is topped by dark gray to black semi-lithified to lithified dolomitic silty claystone (Cores 26-29). The occurrence of nannofossils and foraminifera in Core 26 indicates the change to open marine conditions.

Site 228

Site 228 lies close to the axial valley of the Red Sea about 19°N in a water depth of 1038 meters. Three hundred and fifteen meters of sediment was continuously cored, including 35 meters of brecciated anhydrite.

Core 39, Section 1 consists of intensely brecciated anhydrite in a black siltstone matrix (Figure 38). According to X-ray analyses, the siltstone is composed of layer silicates (chlorite and illite, with smaller amounts of mixed layer illite-montmorillonite and kaolinite), anhydrite, quartz, plagioclase, microcline, pyrite, and traces of gypsum. Under the petrographic microscope (39-1, 120-122 cm; 39-1; Figure 39 a,b), the felted anhydrite crystals (0.1-0.3 mm) are in random orientation. Dolomite crystals (0.1-0.3 mm) and authigenic quartz with anhydrite inclusions are scattered in the anhydrite. This siltstone anhydrite breccia is characterized by a high content of heavy metals, especially zinc (Manheim, this volume). The thin section (39-1; Figure 40), in general, shows euhedral sphalerite crystals aligned with the bedding plane. The sphalerite crystals only occur in the anhydrite layers. The brownish, probably organic-rich, semi-opaque layers are free of sphalerite inclusions.

Core 38 is represented by dark greenish gray to dark brown siltstone. Fragments of anhydrite were found in the core catcher. X-ray analyses of siltstone reveal a mineralogy similar to that found in Core 39. No pyrite and gypsum were detected. Palygorskite is abundant in the greenish material. The dark brown color of the sediments is probably due to amorphous iron. The thin section of the anhydrite (38, CC; Figure 41), in general, shows rounded anhydrite grains (0.1-0.5 mm) with abundant liquid inclusions. The grains are cemented by pyrite impregnated with carbonate material. The relatively uniform grain size and the scattered occurrence of detrital quartz grains suggest reworked and resedimented anhydrite (Ogniben, 1955).

Core 37, Sections 1 and 2 consist of brecciated anhydrite in a greenish dark siltstone matrix. According to X-ray analyses, the siltstone is composed of layer silicates (illite, illite-montmorillonite and chlorite-montmorillonite mixed layer minerals, kaolinite, and chlorite), quartz, feldspars, and traces of dolomite. In Section 2, palygorskite is locally very abundant. Distorted bedded mosaic anhydrite is found in Section 2 (20-35 cm). The thin section (37-1, 145-147 cm; Figure 42) shows felted anhydrite (~0.05-0.5 mm). The large aggregates surrounded by opake, probably organic-rich, material seem to indicate gypsum pseudomorphs.

Partly nodular and partly bedded anhydrite with intercalated siltstone (36-1, 108-120 cm) is represented in Core 36, Section 1.

White bedded massive anhydrite (35-2, 125-150 cm) topped by brecciated anhydrite in a black siltstone matrix and black siltstone (35-2, 0-125 cm) is found in Core 35. Under the petrographic microscope, the lath-shaped anhydrite crystals (~1-1.5 mm) are oriented parallel to the bedding plane (35-2, 120-125 cm; Figure 43). Salt tectonics is suggested by the fractured and bent anhydrite layers (35-2, 116-117 cm; Figure 44). The thin section (35-2, 116-117 cm; Figure 45) shows anhydrite of microcrystalline texture (~0.01 mm). The thin dark layers, which are about 0.5 mm apart, consist of opake, probably organic-rich, material, pyrite, and some tiny dolomite crystals. Bedded massive anhydrite is found at the top of Core 35, Section 2 (35-2, 14-17 cm; Figure 46). Darker layers of course-grained material are intercalated in the anhydrite. Under the petrographic microscope (35-2, 14-17 cm; Figure 47), the anhydrite is of microcrystalline texture (~0.01 mm) and shows bedding due to the intercalated thin opaque layers. Abundant elastic material is interpersed in the anhydrite. The elastic material is generally rounded to angular and consists of quartz, feldspar, and heavy minerals (green hornblende and epidote). In the upper part of the section, the elastic material is fine grained (~0.1 mm),
Figure 32. Alternating clear white and gray halite with intercalated thin anhydrite layers (227-35-4).

Figure 33. Coarse crystalline halite with clusters and streaks of lath-shaped anhydrite (227-35-3, 53-60 cm). Scale bar represents 4 mm.

Figure 34. Coarse crystalline mosaic halite with interspersed polyhalite (227-33-1, 80-84 cm). Partially crossed nicols. Scale bar represents 4 mm.

Figure 35. Fractured laminated black shale with a chalcedony filled vein (227-31, CC). Scale bar represents 1.25 cm.
Figure 36. Fractured cristobalitic shale, cracks are filled with halite, colloform silica is within the halite. (227-31, CC): (a) scale bar represents 4 mm; partially crossed nicols, (b) scale bar represents ~1 mm; partially crossed nicols, and (c) fractured length-slow chalcedony vein with megaquartz; scale bar represents ~1 mm; partially crossed nicols.

Figure 37. Anhydrite of aligned felted texture. Dark brown layers consist of fine-grained dolomite and micron-sized opaque material (227-30-2, 43-45 cm). Thin section is oriented. Scale bar represents 4 mm.
Figure 38. Intensely brecciated anhydrite in a black siltstone matrix.

Figure 39. Felted anhydrite crystals with dolomite and authigenic quartz (228-39-1, 120-122 cm): (a) scale bar 4 mm; crossed nicols; and (b) scale bar 0.5 mm.
Figure 40. Sphalerite and pyrite crystals in anhydrite (228-39-1). Scale bar represents 0.4 mm.

Figure 41. Rounded anhydrite grains cemented by pyrite and some dolomite grains (228-38, CC). Scale bar represents 4 mm.

Figure 42. Felted anhydrite. Large aggregates seem to indicate gypsum pseudo-morphs (228-37-1, 145-147 cm). Crossed nicols. Thin section is oriented. Scale bar represents 4 mm.
Figure 43. Lath-shaped anhydrite crystals oriented parallel to the bedding plane (228-35-2, 120-125 cm). Crossed nicols. Thin section is oriented. Scale bar represents 4 mm.

Figure 44. Fractured and bent massive anhydrite layers (228-35-2, 116-117 cm). Scale bar represents 4 mm.

Figure 45. Microcrystalline anhydrite. Dark layers consist of opaque material, pyrite and small dolomite crystals (228-35-2, 116-117 cm). Crossed nicols. Scale bar represents 4 mm.
whereas, the grain size in the lower section is about 0.5 m which suggests graded bedding. Medium gray distorted bedded to nodular mosaic anhydrite occurs at the bottom of Core 35, Section 1. Frequently, the anhydrite masses are separated by a dark matrix. The thin section (35-1, 133-135 cm; Figure 48) shows anhydrite of aligned felted texture. The bedded character is demonstrated by intercalated layers of opaque and elastic material (quartz, feldspar (~0.07 mm). According to X-ray analyses, the shales of Section 1 are composed of layer silicates, quartz, feldspar, clinozoisite, pyrite, and traces of dolomite. The clay fraction is dominated by montmorillonite and smaller amounts of chlorite and illite. Palygorskite is locally abundant.

As in Sites 225 and 227, the evaporites of Site 228 are topped by a semilithified black dolomitic silty claystone. Marine fossils first occur in Core 33 and increase towards the top of the hole, thus indicating the change to open marine conditions.

GEOCHEMICAL STUDIES

Introduction

Basic research concerning the geochemistry of evaporites was carried out by Van't Hoff and coworkers (1912). Since then, geochemical studies, of trace elements in particular, have given increasing significance. Bromine, strontium, and rubidium are of special interest for the study of halite, anhydrite/gypsum, and potash deposits, respectively.

Methods

Bromine (Br) and Rubidium (Rb)

For the Br determination, the analytical method described by D'Ans and Höfer (1934) was used. Approximately 10 g of sample was dissolved and buffered with phosphate solution, oxidized with NaOCl, and the bromate titrated with n/100 thiosulfate solution after the addition of K solution. No other chloridic mineral was detected under the petrographic microscope. CI was determined according to the Mohr method (AgNO₃) and the Br data could be referred to pure halite.

Potassium was detected flame photometrically and Rb by atomic absorption spectrophotometer. Under the petrographic microscope, polyhalite was determined as the only potassium salt. Therefore, the K content was referred to polyhalite. Rb was detected in two samples which have a relatively high polyhalite content. The Rb values were referred to pure polyhalite. Results are given in Table 1.

Strontium (Sr) and Calcium (Ca)

Analyses were carried out to determine the Sr and Ca content of the anhydrite samples as well as the anhydrite obtained as “water insoluble residue” from one of the halite samples. 1 g of sample was dissolved in 1 liter of water with the addition of HCl and Li₂O₃ and the Sr measured by atomic absorption spectrophotometry.

Ca was determined titrimetrically by means of Titriplex III (C₉H₁₂N₄S₂O₈ 2H₂O). In most samples, anhydrite was the only Ca mineral detected under the petrographic microscope; thus, the Ca could be considered as anhydrite and the Sr be referred to it. Carbonate minerals (dolomite and magnesite) were found in four samples. SO₄ was determined gravimetrically as BaSO₄ in these samples.
Figure 46. Bedded massive anhydrite with layers of clastic material (228-35-2, 14-17 cm). Scale bar represents 2 cm.

Figure 47. Microcrystalline anhydrite intercalated with angular clastic material. Clastic material shows grading (228-35-2, 14-17 cm). Crossed nicols. Thin section is oriented. Scale bar represents 4 mm.

Figure 48. Anhydrite of aligned felted texture with intercalated opaque, probably organic-rich, material and clastic detritus (228-35-1, 133-135 cm). Crossed nicols. Thin section is oriented. Scale bar represents 4 mm.
The Br values of Site 227 show an evaporation profile ranging from a first halite up to a beginning polyhalite region. The Br values occasionally vary up to a first halite precipitation (227-43-5; 227-35-5). This might be indicative of brine fluctuations and dilutions caused by influx of seawater into the basin. The low Br content of Sample 227-30-2, 97-101 cm suggests a very strong dilution. This salt seems to be recrystallized completely.

In general, the bromine content found in the clear white halite is higher than that of the medium gray halite intervals. This was also observed in the prairie evaporite formation (Saskatchewan) by Wardlaw and Schwerdtner (1966). Mineralogical studies show that the clear white halite seems to be younger and to replace the medium gray halite. This observation is difficult to reconcile with the bromine data. According to Schmalz (1969), the marked density (or salinity) stratification is responsible for the difference in the bromine content. The gray halite ("hopper salt") is thought to have formed at the surface, whereas, the clear halite is assumed to have precipitated in situ on the basin floor.

The bromine content in the halite is a useful means of assessing the brine depth. Bromine may be accommodated in the halite crystal lattice replacing chlorine in solid solution. Progressive evaporation causes a concentration of bromine in the residual brine. Consequently, the halite precipitated from such a residual brine is characterized by a high Br content. Therefore, the Br content of the residual brine in a shallow basin should exceed that in a deep-water basin. This principle was applied by Kühn (1953) to develop an equation to calculate the brine depth from which the halite had precipitated. The samples were taken from a distinct evaporation stage within an "annual layer." The values are given in the following table.

<table>
<thead>
<tr>
<th>Sample (m)</th>
<th>Interval (m)</th>
<th>Calculated Depth in meters (T)</th>
</tr>
</thead>
<tbody>
<tr>
<td>225-29-3</td>
<td>0.0128</td>
<td>0.0137</td>
</tr>
<tr>
<td>227-35-1</td>
<td>0.093</td>
<td>0.097</td>
</tr>
<tr>
<td>227-43-2</td>
<td>0.069</td>
<td>0.072</td>
</tr>
</tbody>
</table>

In this case, factor f = 12.3 for the polyhalite region has been applied. Using factor f = 8 for the anhydrite region, the depth values decrease even more.

The table above shows that the brine basin was relatively shallow. It is emphasized that the applicability of this method is limited due to the influx and reflux into or out of the basin. Nevertheless, these values may serve as a rough estimate.

\[
T = \frac{b_2}{b_2-b_1} \cdot \frac{f}{s} = \frac{2.17 \cdot 100}{s \cdot c}
\]

where

- \(T\) = calculated depth in meters
- \(b_1\) = lower sample in meters
- \(b_2\) = upper sample in meters
- \(f\) = factor determined from specific gravity of NaCl
- \(s\) = specific gravity of the brine
- \(c\) = medium NaCl concentration in oceanic salt systems in weight%
Rubidium (Rb)

Rubidium, which can replace potassium diadically in potassium minerals, is also useful in studying natural salt solutions.

The Rb content (0.0014 weight-%) found in Samples 225-29-3, 40-45 cm and 227-45-1, 75-80 cm is the highest Rb value described for polyhalites up to now (cf. Kühn, 1972). Only a polyhalite from Hallstatt, Austria has a comparable Rb content. The high Rb values in the Red Sea polyhalites cannot be explained by the normal evaporation of seawater. It is assumed that the brine was relatively enriched in Rb due to local conditions such as active volcanism accompanied by Rb rich thermal springs or exhalations. Volcanism is indicated by the abundant volcanic material which is intercalated within the evaporite and thus explains the high Rb values. For the Hallstatt salts, melaphyre and fine tuffitic layers give evidence of volcanism (Schauberger, 1960) and may be responsible for the high Rb content.

Strontium (Sr)

The close geochemical relationship between Sr and Ca favors the ionic substitution of Ca by Sr in calcium sulphate minerals. The Sr data furnish valuable information on the genetic character of the salts. The Sr values (referred to pure anhydrite) and the factor Sr/Ca ? 1000 were used to discuss the Red Sea anhydrite deposits. Abundant data from European evaporite deposits concerning the Sr/Ca ? 1000 factor are given by German Muller (1962) and Gerhard Müller (1964). According to Gerhard Müller, Sr/Ca ? 1000 data from 1 to 3 are indicative of isomorphous Sr incorporation; Sr minerals (mainly strontianite) may form already. Strontium minerals are found at values >7, while only celestine should occur in the anhydrite at Sr values >20. Sr values varying less than 20 percent suggest uniform evaporation; whereas, scatter greater than 30 per cent is indicative of irregular changes of the brine, perhaps due to freshwater influx. The Sr/Ca ? 1000 data from the Red Sea samples range between 3.5 and 24.9. Excluding the high value of 24.91 (227-36-2, 73-76 cm), an average value of 7.8 ± 2.7 (scattering of 35 percent) is found. This value corresponds to the oceanic anhydrite region in which a beginning precipitation of Sr minerals is expected. The high scattering may be explained by irregular sedimentation caused by effluents or brine fluctuations.

According to Usdowski (1973), the Sr content of gypsum at the beginning of the CaSO4 precipitation varies between 0.11 and 0.12 weight-%. At the beginning and at the end of the NaCl phase, the Sr content is 0.21 to 0.23 and 0.89 to 0.975, respectively. The data at the end of the NaCl precipitation are somewhat uncertain because the isomorphous replacement of Ca with Sr takes place simultaneously with the formation of celestine.

If gypsum changes early diagenetically into anhydrite due to a relatively high temperature in the CaSO4 phase of the evaporation, this anhydrite contains 0.229-0.202 weight-% Sr according to Usdowski; whereas, primary anhydrite should already have a Sr value of 0.229 weight-%.

In general, the Red Sea anhydrite samples are of massive character which suggests their deposition in the CaSO4 phase. On the basis of Usdowski’s data, it might be concluded that part of the anhydrite (anhydrite samples having a Sr content of <0.229) originated diagenetically from gypsum. The part of the anhydrite having a higher Sr content (>0.229 weight-%) must be considered to be of primary origin.

The Sr values of the anhydrites which occur together with carbonate material were calculated according to Usdowski with the distribution factor b = 1200 for anhydrite and b = 100 for CaCO3. The high value of Site 225-24-1 is probably indicative of the presence of Sr minerals. The two samples of the halite phase (225-28-3, 95-105 cm; 0.05 mm φ and 0.2 mm φ) show a relatively low Sr content; therefore, the thin anhydrite layers within the halite might be conceived as short recurrences back into the CaSO4 phase of evaporation. The Sr values varying with the grain size might correspond to different generations of CaSO4 growth.

Origin of the Evaporites

It is assumed that separate great uplifts on either side of the Red Sea during late Eocene and early Oligocene time produced the Ethiopian and Arabian swells with the proto-Red Sea depression separating them (Coleman, this volume). The Miocene transgression of the Mediterranean Sea into this closed depression led to thick evaporite deposits.

Two main problems occur concerning the depositional environment of evaporites. The first problem is whether the evaporites were precipitated under shallow- or deep-water conditions; the second is the question of the depth of the basin in which the evaporites were formed.

The investigation of recent sabkha environments (Tunisia, Libya, Baja California, etc.), especially at Abu Dhabi on the Arabian side of the Persian Gulf (Trucial Coast) (Shearman, 1963; Kinsman, 1966, 1969; Evans, 1966; Kendall and Skipwith, 1969; Butler et al., 1965; Butler, 1969; and others), have provided new models for the interpretation of ancient evaporite deposits. The recent sabkha or tidal flat sediments are characterized by distinct structures such as stromatolites, nodular gypsum and anhydrite, planar and ripple cross-lamination, flat pebble conglomerates, polygonal shrinkage cracks, to mention the most important features. Using recent sabkhas as a guide, much evidence for the presence of former sabkha environments is found in the salt deposits of the geological past (Fuller and Porter, 1969; West, Brandon, and Smith, 1968; Hardie and Eugster, 1971; Shearman and Fuller, 1969; Friedman, 1973).

The petrographic study of the Red Sea evaporites revealed abundant evidence for a sabkha deposit. Undulating laminations composed of semi-opaque or opaque, probably organic-rich, material impregnated with very fine grained dolomite are found within the anhydrites at Site 225 (Cores 24, 25; Figure 15) and Site 227. Finely laminated spheroidal forms consisting of dolomite, organic material, and traces of pyrite are abundant at Site 225, Core 26 (Figure 13). These features are interpreted as algal stromatolites and oncolitcs which accumulated in shallow water.
The nodular anhydrites found in the Red Sea evaporite sequence (especially at Site 225, Core 29; Figure 5) are similar to the supratidal anhydrite nodules of recent sabkha deposits. The nodules grow in carbonate sediments above the ground water table and displace the overlying sediments. At a more advanced stage the nodules coalesce. The original boundaries become distorted which results in a mosaic or distorted mosaic structure. Due to continuous growth, the nodules finally merge into one another (enterolithic structure, Shearman, 1963). These different structures can be seen in the Red Sea anhydrites.

The frequent alternations of halite, anhydrite, and dolomitic black shales reflect strong dilutions and reconcentrations of the brine, which seems most likely to be related to a shallow-water environment.

Further support for a salina model for the Red Sea evaporites is given by geochemical studies. Depth calculation by means of the bromine content revealed a very shallow brine depth in the range of some meters. The data obtained by isotope studies (Friedman and Hardcastle, this volume) show a strong influence of fresh water which can only be explained by a salina model. The great variations in the pore water data (Manheim, this volume) also agree with a shallow-water deposit.

Both the petrographical and the geochemical data suggest that the Red Sea evaporites recovered during Leg 23 originated from very shallow water with occasional subaerial exposure.

This assumption, however, does not permit drawing any conclusions on the depth of the basin. The main difficulty with the explanation of ancient salt deposits is the discrepancy between the rate of salt deposition in comparison to the possible rate of basin floor subsidence. These problems have been discussed very carefully by Schmalz (1969). Schmalz showed that this discrepancy can be solved with a deep-water–deep basin model. The salts were deposited in basins several hundreds to thousands of meters deep. (The discovery of salts under the Gulf of Mexico (DSDP Leg 1, Ewing et al., 1969) is in excellent agreement with this model). Indeed, the paleogeographic reconstruction of the Red Sea suggests that the proto-Red Sea depression represented a topographic depression of considerable depth. The stratigraphic and structural record indicates that the NW-SE trending depression developed as a trough between the Arabian and African swells rather than as a down-faulted block (Brown and Coleman, 1972; Coleman, this volume). During Miocene time, a great thickness of clastics (2-3 km) and evaporites (3-4 km) was deposited in this proto-Red Sea depression as demonstrated by deep exploratory wells (Ahmed, 1972). It is concluded that a "shallow-water-shallow basin" model for the evaporite deposits does not agree with this geological information which evidently favors a deposition in a deep basin.

The question which remains to be answered is whether the salts were precipitated in deep water or in very shallow water on the floor of a desiccated deep basin. This problem has been discussed in detail by Hsü et al. (1973) for the Mediterranean evaporites (DSDP Leg 13). They consider a desiccated deep basin as the only logical alternative to explain the Mediterranean evaporites. The arguments used by these authors against the Schmalz model are not entirely satisfactory. As in the Red Sea, the evaporites recovered in the Mediterranean penetrate only the terminal part of the evaporite sequence. At this final stage, one can also expect a salina or sabkha deposition according to the deep water-deep basin model (see terminal stage II, Figure 8 [nearly filled basin] Schmalz, 1969).

The sedimentological and geochemical studies of the evaporites recovered on Leg 23 reveal convincing evidence of a shallow-water evaporite deposit. However, this evidence should be used with caution. It cannot, necessarily, be taken as an explicit indicator for the total amount of Red Sea evaporites (3-4 km) to be of salina or sabkha origin. Further evidence of the deep strata is necessary and could be provided by the data obtained from deep exploratory wells. Unfortunately, most of these data have not been released for publication by the oil companies.

Leg 23 has proven that Reflector S in the Red Sea is the top of a Miocene evaporite deposit comparable to Reflector M in the Mediterranean. Assuming a connection between the two basins, the "waterfall" descending from the Strait of Gibraltar must be held responsible not only for the up to two kilometer thick evaporite sequence in the Mediterranean but also for the three to four kilometer thick evaporites in the Red Sea. Hsiu et al. (1973) concluded that eight or ten marine invasions could have been sufficient to account for all the salts found under the Mediterranean abyssal plain. Considering the idea of a repeated refilling of the basin during Miocene time, it seems that a desiccated deep-basin hypothesis is not in contrast to the deep water-deep basin model postulated by Schmalz (1969). The desiccated deep-basin model may be regarded as a specific stage in a cycle of repeated refilling and evaporation. Therefore, it is concluded that the origin of the Mediterranean and Red Sea evaporites could be accounted for by Schmalz's "Deep-Water Model" modified by periods of desiccation caused by occasional isolation of the Mediterranean from the Atlantic.

ACKNOWLEDGMENTS

The senior author is indebted to the entire Leg 23 scientific and technical party for their cooperation aboard the Glomar Challenger. Special thanks are due D. A. Ross, P. R. Supko, and O.E. Weser. The assistance given by F.T. Manheim leading to improvement of the manuscript is greatly appreciated. Part of the work was carried out at Woods Hole Oceanographic Institution; financial support came from NATO. We thank K. H. Ritter for doing the chemical analyses and W. Kücking for preparing the thin sections.

REFERENCES


Borchert, H. and Baier, E., 1953. Zur Metamorphose
ozeanischer Gipsablagerungen: Neues Jahrh. Min.,
Abhandl., v. 86, p. 103-154.

Boeke, H.E., 1908. Über das Kristallisationsschema der
Chloride, Bromide, Iodide von Natrum, Kalium, und
Magnesium, sowie über das Vorkommen des Broms und
das Fehlen von iod in den Kalsalzlagerstätten: Zeitschr.
Krist., v. 45A, p. 346-391.

Braitsh, O. and Herrmann, A. G., 1962. Zur Bromver-
teilung in salinaren Salzsystemen bei 25°C: Naturwiss.,
v. 49, p. 336.

Brönnim, W. D., 1962. Idiomorphe Quarze als Leitmineralien
Folk, R. L. and Pittman, J. S., 1971. Length-slow chal-
Evans, G., 1966. The recent sedimentary facies of the
Borchert, H. and Baier, E., 1953. Zur Metamorphose
for halite precipitated from modern sea water under
Heybroek, F., 1965; The Red Sea evaporite basin. In Salt
Second Symposium on Salt: Cleveland, Ohio (N. Ohio

1966b. Diagenetic polyhalite in recent salt from
Hsii, K. J., Cita, M. B., and Ryan, W. B. F., 1973. The
origin of the Mediterranean evaporites. In Ryan, W. B. F.,
Hsii, K. J., et al., Initial Reports of the Deep Sea Drilling
Project, Volume XIII: Washington (U.S. Gov-
ernment Printing Office).

Holocene shallow-water carbonate and evaporite sedi-
ments of Khor al Bazam, Abu Dhabi, southwest Persian

Kinsman, D. J. J., 1966. Gypsum and anhydrite of Recent
age, Trucial Coast, Persian Gulf. In Raup, J. E. (Ed.)
Second Symposium on Salt: Cleveland, Ohio (N. Ohio

Kosmahl, W., 1969. Zur Stratigraphie, Petrographie, Paläo-
geographie, Genese und Sedimentation des Gebänderten
Anhydrits (Zechstein 2), Grauen Salztones und Hauptan-
yhydräts (Zechstein 3) in Nordwestdeutschland: Beihete
Geol. Jahr., no. 71.

Kühn, R., 1953. Tiefenberechnung des Zechsteinmeeres
Gesell., v. 105, p. 646-663.

1955. Über den Bromgehalt von Salzgesteinen, insbesondere die quantitative Ableitung des Brom-
gehaltes nichtprimärer Hartsalze oder Sylviniten aus
Carnallit: Kali und Steinsalz 19, 9, p. 3-16.

1968. Geochemistry of the German Potash

1972. Zur Kenntnis der Rubidiumgehalte von
Kalsalzen ozeanischer Salzlagerstätten nebst einigen
gerößten kundlichen Ausdeutungen: Geol. Jahrh., v. 90,
p. 127-220.

(Z 3) im Südhharz: Chemie der Erde, 21, 2, p. 248-264.

Link, G., 1938. Bildung des Dolomits und Dolomitisation;

1942. Beobachtungen und ihre Ergebnisse an
Gesteinen des mittleren Zechsteins (Hauptdolomit und
grauer Salzton) in Thüringen: Chemie der Erde, v. 14,
p. 315-357.

Lowell, J. D. and Genik, G. J., 1972. Sea-floor spreading
and structural evolution of southern Red Sea: Am.

Classification of anhydrite—a practical approach: Bull.

Müller, Gerhard, 1964. Ein Beitrag zur Geochemie des
Strontiums in Ca-Sulfat-Gesteinen nebst einigen
Gesteinen des mittleren Zechsteins (Hauptdolomit und
grauer Salzton) in Thüringen: Chemie der Erde, v. 14,
p. 315-357.

Nesteroff, W. D., 1973. Mineralogy, Petrography, Distribu-
tion, and Origin of the Messinian Mediterranean evap-
orites. In Ryan, W. B. F., Hsii, K. J., et al., Initial Reports


