ABSTRACT

Observations of the physical properties of sediments, the chemistry of interstitial waters and solid phases, the mineralogy of the sediments, and the isotope geochemistry of interstitial waters and sediments have led to the conclusion that two types of hydrothermal systems are operative in the Guaymas Basin of the Gulf of California: (1) hydrothermal activity associated with relatively shallow basaltic sill intrusions in highly porous sediments—generally of relatively short duration and associated with temperatures <200°C—and (2) hydrothermal activity caused by large magmatic intrusions at greater depths. This latter activity involves recharge of the hydrothermal system by basin bottom waters; fractures in low-porosity sediments and faults in the sea floor serve as conduits for fluids heated up to temperatures in excess of 300°C. This type of hydrothermal activity lasts long enough to lead to substantial alteration of sediments into a well-developed greenschist facies: chlorite-quartz-albite-sphene-epidote-Fe-sulfides. The mechanisms of the hydrothermal systems are discussed in detail and are consistent with the extensive heat flow data obtained in this area.

INTRODUCTION

In order to explain heat flow anomalies near oceanic spreading centers, the interaction between upwelling hot magmas and downward-flowing ocean water has been invoked by various authors (Lister, 1972; Sclater and Klitgord, 1973; Williams et al., 1974; Wolery and Sleep, 1976; Anderson et al., 1977, 1979). Subsequently, hydrothermal vents have been discovered in the Galapagos Rift (Corliss et al., 1979), on the East Pacific Rise at 21°N (Francheteau et al., 1979;Spiess et al., 1980), and at other oceanic spreading centers. Chemical analyses of hydrothermal waters emanating from vents at the Galapagos Rift, as well as of those obtained at 21°N, indicate profound changes in chemical composition of these waters when compared to the original sea water (Corliss et al., 1979; Edmond et al., 1979a, b, c), similar to those observed in hydrothermal sea water–igneous rock interaction experiments (Bischoff and Dickson, 1975; Hajash, 1975; Menzies and Seyfried, 1979; Mottl and Holland, 1978; Seyfried and Bischoff, 1977). Associated with such hydrothermal activity are metal-rich deposits of various composition (Bostrom and Peterson, 1965; Francheteau et al., 1979; Haymon and Kastner, 1980; Moore and Vogt, 1976; Natland et al., 1979; Scott et al., 1974).

These observations have been made in environments typical of mature oceanic spreading ridges, that is, in areas of low sedimentation rates, where basalts are exposed to the ocean. In this report, however, we wish to describe hydrothermal activity in the Guaymas Basin of the Gulf of California (Fig. 1), which is more representative of the initial stages of ocean evolution. Here basaltic magma forming new oceanic basement intrudes as sills into rapidly deposited soft sediments, thus creating a basement quite different from the normal ophiolite model (Moore, 1973; Einsele et al., 1980; Einsele, this volume, Pt. 2).

Prior to the drilling program in the Guaymas Basin (Fig. 1), evidence for hydrothermal circulation in this area was obtained from intensive heat flow surveys (Lawver et al., 1975; Williams et al., 1979). The extremely local distribution of areas of high heat flow (Fig. 1) was interpreted in terms of basaltic intrusions. Hydrothermal deposits were discovered in the northern trough of the basin (Fig. 1) and have been described by Lonsdale (1978; Lonsdale et al., 1980). Since the Leg 64 drilling program, extensive hydrothermal deposits have been mapped in August 1980 in the southern trough (Lonsdale, pers. comm.) and sulfide deposits have been sampled in the same area by Alvin dives in January 1982.

Essentially the hydrothermal system in the Guaymas Basin, which we describe in this report, is one of a series of hydrothermal systems associated with the East Pacific Rise and the extension of this spreading system into the Gulf of California and the Imperial Valley of Southern California. Whereas at 21°N on the East Pacific Rise hydrothermal solutions emanate directly from the basalts, in the Guaymas Basin the high-temperature fluids stem from interactions of sea water and sediment interstitial waters with heated sediments and with basaltic lavas. In the Imperial Valley of Southern California and Mexico, magmas penetrate into terrigenous sediments and hydrothermal waters originate by advection of meteoric waters into the system (Elders, 1979).
Although sill intrusions have previously been noticed in deep-sea drilling studies, their effects on surrounding sediments have not received much attention. A notable exception to this, of course, is the study of the organic geochemistry of sediments near a basaltic sill intrusion at Site 268, DSDP Leg 41 (Baker et al., 1978; Simoneit et al., 1981). In the Guaymas Basin, where sill intrusions occur now or have occurred in the recent past (Einsel e et al., 1980; Lonsdale, pers. comm.), there arose an excellent opportunity to study their effect on the physical properties of the sediments, the mineralogy and chemistry of the solid phases, and the chemistry of the connate waters.

In this chapter we discuss the nature of the hydrothermal systems operative in the Guaymas Basin on the basis of our observations during DSDP Leg 64. Detailed background data can be obtained from other chapters by Gieskes, et al., Kastner, Einsel e, Kelts, and Niemitz (this volume, Pt. 2). Here we first summarize these observations and then undertake a general discussion of the nature of the hydrothermal systems. The effects of the basaltic sill intrusions on the organic geochemistry of the sediments at these sites are summarized by Simoneit (this volume, Pt. 2).

**LITHOLOGY OF GUAYMAS BASIN SITES 477, 478, AND 481**

In this chapter we focus mainly upon Sites 477 and 481, both located in the actual spreading troughs, but Site 478 is also of interest, in that it is located near the transform fault connecting the active spreading troughs (Fig. 1). The lithologies of Sites 477, 478, and 481 are given in Figure 2. Note in particular the occurrence of sill intrusions in all three of these sites. Heat flow at Site 477 is higher than 20 HFU (1 HFU [heat flow unit] = 10^{-6} \text{ cal cm}^{-2} \text{ s}^{-1}), whereas at Site 481 a heat flow of \sim 4 HFU and at Site 478 a heat flow of 3.65 HFU were measured.

The sediments in this area are generally characterized by rapidly deposited diatomaceous oozes and clays, tur-
Figure 2. Lithology of Sites 477, 478, and 481, Guaymas Basin.
bidites, and mudflows. Intrusions of dolerite sills have subsequently altered these sediments during and shortly after emplacement of the sills. Large changes in porosity above and below the sills attest to the fact that the layers of dolerite are sill intrusions rather than basalt flows deposited on top of the sediments (Einsele et al., 1980). At both Site 477 and Site 481, contact metamorphism has affected the sediments above and below the sill intrusions. At Site 477, below the upper sill complex, extensive hydrothermal activity has led to the formation of greenschist facies rocks (Fig. 2).

These observations on the thermal alteration of the sediment have led us to postulate that two types of hydrothermal systems are operative in the Guaymas Basin spreading troughs (Kastner and Gieskes, 1981):

1) Hydrothermal activity associated with the emplacement of dolerite sills into highly porous sediments, and
2) Hydrothermal activity associated with large-scale magmatic intrusions under a relatively thick sediment cover.

In the following sections we describe the characteristics of these hydrothermal systems.

PHYSICAL PROPERTIES

Einsele (this volume, Pt. 2) describes careful studies of the physical properties of the sediments in the Guaymas Basin and notes in particular large decreases in porosity near the sill intrusions (Fig. 3). Einsele et al. (1980) and Einsele (this volume, Pt. 2) estimate the volume of pore waters that must have been expelled as a result of the thermally induced convection and suggest that the reduction in sediment thickness is roughly equivalent to the thickness of the sills. The important points for our discussion are that large-scale migration of interstitial waters away from the sill intrusions does occur and also that the zones above the sills over which porosities appear to be affected probably represent the minimum thickness of the sediments present at the time the sill intruded. This implies that sills penetrate into these sediments at depths of at least 10-50 meters below the sediment/water interface. In turn, this means that this water must have been expelled off the sediments and into the basin bottom waters, presumably along faults, rifts, and fissures or along the basin fault scarps.

INTERSTITIAL WATER CHEMISTRY

A detailed description of the interstitial water chemistry of Sites 477, 478, and 481 is presented by Gieskes et al. elsewhere in this volume, Pt. 2. Here we emphasize the observations relevant to high-temperature interactions between the dolerite sills and heated sediments and the interstitial waters of the sediments. Figure 4 presents the chemistry of the interstitial waters for Sites 477 and 481.

At Site 477, where hydrothermal activity still continues, gradual increases with depth in the concentrations of chloride, calcium, lithium, potassium, and rubidium indicate that reactions in the sediment column below ~180 meters are chiefly responsible for the large concentration increases in these constituents. Especially higher than expected are increases in potassium and rubidium when compared with concentrations resulting from hydrothermal interactions between sea waters and basalts (Edmond et al., 1979a). This can be understood in terms of the hydrothermal interactions of sea water and interstitial water, not only with the basaltic intrusions but also with the sediments, which contain much higher concentrations of these elements than do the basalts. Indeed, at Site 477 below 180 meters, potassium has been removed almost quantitatively from the solid sedimentary phases. Increases in chloride concentrations are most likely due to removal of water into hydrated alteration phases.

Site 481 represents a different situation. At this site, the most recent hydrothermal event must have been associated with the intrusions of the sill complex between present sub-bottom depths of 170 and 200 meters. This is clearly indicated by the profiles of dissolved chloride and of dissolved rubidium and lithium, all of which show enhanced levels around the sills. Not enough time has yet elapsed to annihilate such concentration anoma-
lies by diffusion and reaction processes. The sill complex has cooled considerably since its intrusion, whence the low heat flow (4 HFU) and the decrease in dissolved potassium and lithium toward the sills. Below 75°C (Seyfried and Bischoff, 1979) basalt alteration will generally lead to the uptake of potassium, rather than to its release to the pore fluids. The signals in chloride, lithium, and rubidium simply indicate that the effects of the hydrothermal event have not yet decayed away by diffusional transport processes.

CHEMISTRY AND MINERALOGY OF SEDIMENTS

The bulk chemical compositions of the solid phases of the sediments of Sites 477, 478, and 481 are described in detail in the chapters of Niemitz (this volume, Pt. 2) and Gieskes et al. (this volume, Pt. 2). Mineralogical data are discussed by Kastner (this volume, Pt. 2) and Kelts (this volume, Pt. 2).

In general, two types of hydrothermal effects can be distinguished in these holes in the Guaymas Basin:

1) Contact metamorphism of the sediments as a result of basaltic sill intrusions.

2) Greenschist facies metamorphism as a result of substantial hydrothermal alteration of sediments at high temperatures (>200°C).

In Table 1 the relevant mineralogical and chemical observations are summarized.

Contact metamorphism is generally associated with little change in the chemical composition of the sediments, with the exception of noticeable increases in Mg/Al near the sill of Hole 477, and decreases in K/Al, Si/Al, and Li near the upper contact with the sill in Hole 477 and perhaps in Hole 481A. The mineralogical observations described in Table 1 indicate that contact metamorphism is generally associated with temperatures below 200°C.

Table 1. Mineralogy and chemistry of hydrothermally altered sediments.

| Contact Metamorphism: Associated with sill complexes in Site 477 and 481 |
|-----------------------------|-----------------------------|-----------------------------|
| Above sills: Clays recrystallized to smectite; calcite recrystallized (Site 481: ~170°C); Diatoms transformed to quartz; pyrite |
| Below sills: Same as above sills, but to lesser extents |
| In 481: Na-phase is analcime, indicating T <200°C |
| In 477: Some K-feldspar |
| Chemistry (above and below sill) increase Mg/Al (477); decrease Si/Al (477); decrease in K/Al (above sill 477); decrease in Li (477; 481) |

Greenschist Facies: Site 477/477A below sills (Fig. 2)

- 100 m: Contact metamorphism; K-feldspar result of activity below
- 120-130 m: Chlorite appears; K-feldspar, albite; anthophyllite and gypsum; pyrite; quartz
- 130-140 m: Appearance of epidote and spherule; chlorite increasing with depth
- 180 m: Layer with abundant pyrite nodules; co-occurrence pyrite and pyrrhotite below indicates T <300°C
- 190-260 m: Dominant assemblage in greenschist facies: chlorite (iron rich)-quartz-anthophyllite-sphene-epidote; nonsilicates: pyrite and pyrrhotite

Chemistry: disappearance of K; low Zn, V; increased Zn (in Fe-sulfides?)
Greenschist facies metamorphism at Site 477 requires higher temperatures than those prevailing around sill intrusions, probably over longer time periods of hydrothermal activity than in the zones of contact metamorphism. The mineralogical observations at Site 477 present a classical sequence of hydrothermal greenschist facies. In the sediment horizons below ~150 meters, substantial hydrothermal alteration has led to the formation of the assemblage chlorite-quartz-albite-sphene-epidote and to the almost complete removal of potassium from the solid phases. This assemblage, together with the occurrence of pyrite/pyrrhotite, implies temperatures of up to 300°C (Hoagland and Elders, 1978; Elders et al., 1979). The heat flow data obtained at Site 477 (>20 HFU) also suggest temperatures in excess of 200°C at these depths. Kastner (this volume, Pt. 2) estimates a temperature of ∼300°C using oxygen isotope data on the co-occurrence of chlorite-quartz. At depths of ∼180 meters there occurs a layer of pyrite nodules, up to 5 mm in diameter. These well-crystallized pyrites are probably the result of in situ precipitation of sulfides as a result of upward-migrating hydrogen sulfide and iron-rich solutions. In the upper part of the greenschist zone, the high dissolved-potassium concentrations in the upward-advecting interstitial waters lead to the precipitation of K-feldspar between 130 meters and the sill intrusion at 100 meters. Elders et al. (1979) indicate the temperature range of the K-feldspar stability zone to be 150–300°C.

**STABLE ISOTOPES**

Studies of stable isotopes have concentrated on oxygen isotopes in solid silicates and interstitial waters (Kastner, this volume, Pt. 2; Gieskes et al., this volume, Pt. 2), on strontium isotopes in bulk sediments and interstitial waters (Gieskes et al., this volume, Pt. 2) and on sulfur isotopes in the sulfides of Site 477 (Shanks and Niemitz, this volume, Pt. 2).

Oxygen isotopic data are summarized in Figure 5. As discussed by Kastner (this volume, Pt. 2) the δ¹⁸O value of silicates in the vicinity of the sill intrusions drop to values of approximately 10‰ above the sills and 12‰ below the sills. The oxygen isotopic composition of interstitial waters above the sills appears lower than below the sills, at least at Site 481. This can be understood in terms of renewal of water by recharge above the sills, thus leading to lower δ¹⁸O in the alteration phases. The very high δ¹⁸O values of the interstitial waters below the sill at Site 481 indicate that the system below these sill intrusions is essentially closed.

In the zone of greenschist facies at Site 477, δ¹⁸O values of the silicates gradually decrease to ~6.5‰ (SMOW), values similar to those of fresh basaltic waters. Recovery in the deeper sections of the sediments unfortunately was poor, and we were unable to make a precise determination of the interstitial water profile of δ¹⁸O at Site 477. The data in Figure 5, however, suggest that relatively low values occur despite the large decrease in δ¹⁸O of the solid phases. This observation suggests that below the sills at Site 477 the hydrothermal system is an open system, with hydrothermally heated waters being replaced by recharge of bottom waters in the Guaymas Basin.

Data on δ¹⁸O of interstitial waters at Site 478 suggest slightly elevated values as a result of past hydrothermal reactions. However, diffusion processes and low-temperature alteration reactions have almost erased the originally increased δ¹⁸O values.

Data on the distribution of δ⁴⁰Sr/δ⁸⁶Sr in the dissolved strontium of the interstitial waters are summarized in Figure 6. The data suggest that hydrothermal interactions between interstitial waters and the sediments and/or basalts have led to substantial equilibration with the interstitial waters. In the deeper sections of Hole 477, interstitial-water δ⁴⁰Sr/δ⁸⁶Sr ratios are lower than those of the bulk solids (Gieskes et al., this volume, Pt. 2). Complete equilibration with basaltic material would lead to a ratio of 0.703–0.704 in this site (cf. Menzies and Seyfried, 1979; Elderfield and Greaves, 1981; Albarede et al., 1981). Further work will be necessary to determine the relative contribution of sediments and basalts to this isotopic exchange.

Sulfur isotopic studies of sulfides in the sediments of Site 477 suggest a range, for δ³⁴S, of ~11.06 to +7.31‰ (Shanks and Niemitz, this volume, Pt. 2). Below 170 meters, δ³⁴S values of sulfides are in the range of 5–7.31‰, values considered to be representative of...
Hyalocline of the Guaymas Basin has been shown from 34°C to 315°C. That these vents do contribute substantially to the bottom waters of the Guaymas Basin and estimated that waters emanating from the hydrothermal vent associated with this deposit must have reached temperatures of ~280°C. Similarly, during dives of the submersible *Alvin* in January 1982, hydrothermal sulfide deposits and associated vents indicated exit temperatures of hydrothermal waters between 275 and 315°C. That these vents do contribute substantially to the bottom waters of the Guaymas Basin has been shown from 1He anomalies observed by Lupton (1979).

The information gained during Leg 64 has served mainly to clarify the nature of magmatic intrusions previously postulated to occur in the Guaymas Basin (Moore, 1973; Lawver et al., 1975). Two major types of hydrothermal systems can be distinguished; these are represented schematically in Figure 7: (1) hydrothermal activity of relatively short duration associated with sill intrusions into sediments of high porosity (Fig. 7B), and (2) hydrothermal activity of greater longevity associated with proximate magmatic sources (Fig. 7A). In the following we wish to discuss these systems in greater detail.

Silicic intrusions into wet sediments appear to be the main mechanism of new ocean floor creation in the Guaymas Basin of the Gulf of California (Einselle et al., 1980; Einselle, this volume, Pt. 2). Drilling at Sites 477, 478, and 481 suggests that the thicknesses of such sills can be greater than 100 meters, but that generally smaller thicknesses occur, except perhaps near the magmatic sources for these intrusions. Data on porosity decreases (Fig. 3) clearly indicate that large-scale reductions occur in porosity both above and below these sill intrusions. Associated with these porosity decreases are large-scale expulsions of pore fluids, the height of the water column expelled often being in excess of the thickness of the sill intrusion (Einselle et al., 1980; Einselle, this volume, Pt. 2). Sill intrusions generally occur within 50 meters of the sediment/water interface, which is evident not only from the thickness of the porosity-reduction zone, but also from the Deep Tow records in the basin (P. Lonsdale, per. comm.). The porosity reduction does not per se require a recharge of water from the basin waters, though such recharge cannot be disproven, especially above the sill intrusions. Associated with the hydrothermal activity caused by the intrusion of the hot sills are changes in interstitial water chemistry (Figs. 4, 5, and 6), which indicate changes of the same nature as described for hydrothermal fluids associated with open-ocean spreading ridges (e.g., Edmond et al., 1979a). Differences in chemical composition are caused mainly by the involvement of sediments as well as basaltic melts in the hydrothermal interactions. Evidence from oxygen isotopes of bulk silicates suggests moderate temperature alteration (<200°C). The expelled waters would be transported out of the sediments mainly along faults and fissures on the trough floor or at the fault scarps of the troughs (Figs. 7; Einselle et al., 1980; Lonsdale et al., 1980). Exit temperatures of hydrothermal vents associated with these sill intrusions should be moderate, though sulfides may still be associated with possible hydrothermal deposits. When magnesium removal from the heated interstitial waters is incomplete, the expelled fluids, which would contain high concentrations of dissolved silica, could lead to deposition of magnesium silicates. However, the talc deposit described by Lonsdale et al. (1980) implies exit temperatures above 280°C, and should, therefore, be associated with the second type of hydrothermal activity.

In the vicinity of more permanent heat sources, for example near magma chambers or feeder dikes, it is possible to develop hydrothermal systems in which recharge by bottom waters plays an important role. High temperatures in the sediments will be reached and the seat of hydrothermal activity will be located at greater depths than the depth at which sill intrusions generally occur. Site 477 is a typical example of this case. The sills in Holes 477 and 477A are essentially cooled off and are not the cause of the observed high heat flow at this site. Instead, the sill complex of this site serves in many ways as a cap rock for the hydrothermal system, thus causing...
the possible circulation pattern presented in Figure 7A. Site 477 is located between two elevated structures in the southern trough of the basin (Lonsdale, pers. comm.), which are probably due to the large accumulation of basaltic material of relatively young age (Lonsdale, pers. comm.). These hills are probably related to the formation of the sills in Holes 477 and 477A. Lonsdale (pers. comm.) also indicates the presence of fault zones near Hole 477A. We postulate that a magmatic source below Site 477 has been active over a considerable period of time, thus allowing the advanced state of hydrothermal alteration of the sediments. The zone of hydrothermal alteration is at least 100 meters thick. We postulate that the return flux of hydrothermal waters to conduits such as faults occurs mainly in the lower parts of Site 477. The temperatures of 280-315°C for exiting fluids recently observed from the submersible Alvin in this area imply considerably large fluxes of water. At these high temperatures, the emanating water will presumably be enriched in sulfides and trace metals, which then can lead to sulfide deposition. The sources of these sulfides and trace metals may be located both in the basalts (evident from 3He anomalies; Lupton, 1979) and in the sediments. Waters advecting through the sediments at slower rates, that is, at depths less than 180 meters at Site 477, will lead to the deposition of sulfides in the sediments as well as to the formation of K-feldspars near the upper sill complex.

CONCLUSION
Drill sites in the Guaymas Basin in the Gulf of California demonstrate that during the initial stage of opening of some ocean basins, basaltic magma forming new oceanic basement intrudes as sills into rapidly deposited soft sediments, creating a basement quite different from the normal ophiolite model.

Two types of hydrothermal systems can be recognized:
1) Hydrothermal activity associated with basaltic sill intrusions at relatively shallow depths (<50 m) into highly porous sediments. Associated with this activity are contact metamorphism (recrystallization of clays, dissolution of opaline silica; recrystallization of carbonates) and chemical changes in interstitial waters (increases in lithium, rubidium, potassium, and δ18O; decreases in magnesium and 87Sr/86Sr) and in the solid phases (decreases in K/Al, Li, δ18O of silicates; increases in Mg/Al). Isotopic evidence on recrystallized carbonates and relatively high δ18O values of recrystallized silicates indicate that temperatures >100°C and <200°C are involved in this hydrothermal activity.
2) Hydrothermal activity associated with large-scale magmatic intrusions at greater depths in the sediments. In this case hydrothermal activity leads to substantial thermal alteration of sediments (greenschist facies), and mass-balance considerations of oxygen isotopes in bulk silicates and interstitial waters lead to the conclusion that these alteration reactions must occur in an open system at temperatures of approximately 300°C. Recharge of basin bottom waters and discharge of heated fluids occur through fault zones, with sills acting as lids on these hydrothermal systems, thus forcing fluid flow in lateral directions toward faults. At locations of such hydrothermal activity, fluids emanating from faults and fissures will lead to the buildup of hydrothermal deposits recently observed in the southern trough by the submersible Alvin.
REFERENCES


