21. PORE WATERS FROM SERPENTINITE SEAMOUNTS IN THE MARIANA AND IZU-BONIN FOREARCS, LEG 125: EVIDENCE FOR VOLATILES FROM THE SUBDUCTING SLAB

Michael J. Mott

ABSTRACT

Sediment pore fluids sampled by drilling in subduction zones are commonly depleted in chloride relative to seawater and highly enriched in methane. These features have been attributed to processes within the accretionary prism of sediment that is typical of most subduction zones. Sites 778 through 780 in the Mariana forearc yielded low-chlorinity pore fluids rich in methane, ethane, and propane from a subduction zone setting that lacks an accretionary prism. They were drilled on the flanks and summit of Conical Seamount, an active serpentine mud volcano located 80 km behind the Mariana Trench and 120 km in front of the island arc. Alvin dives on this summit in 1987 found chimneys made of aragonite, calcite, and amorphous Mg-silicate from which cold water was seeping. During drilling at the summit Site 780 we recovered these fluids from serpentine silts at depths up to 130 mbsf and demonstrated that the fluids are upwelling through the seamount at velocities greater than a few millimeters per year. The fluids have less than one-half the chloride and bromide of seawater, pH up to 12.6, methane up to 37 mmol/kg along with ethane and propane, H$_2$S up to 2 mmol/kg, and ammonia up to 270 µmol/kg. Relative to seawater, they are enriched in alkalinity (+26), sulfate (+1.7), K (+1.5), Rb (+5.6), and B (+10). They are highly depleted in Li, Mg, Ca, Sr, and $^{34}$S, and have low concentrations of Si, Ba, and Mn. Although they are also depleted in Na, their Na/Cl ratio increases to nearly twice that in seawater.

These pore fluids from the summit of Conical Seamount contrast greatly with those recovered from Sites 783 and 784 on Torishima Seamount, an inactive serpentine seamount in the Izu-Bonin forearc. The Torishima pore fluids were produced by the reaction of seawater with harzburgite at low temperatures. They have pH up to 10, very low Si, Mn, and methane, and no ethane or propane. Relative to seawater, they have low alkalinity, sulfate, Mg, K, and B; slightly lower Li and Rb; little-changed chloride; Br, Na, and NaCl; and high Ca, Sr, Ba, and $^{34}$S. Pore waters from Sites 778 and 779 on the flanks of Conical Seamount are complex mixtures of these two types of fluids.

The absence of an accretionary prism in the Mariana forearc severely constrains the origin of the fluids upwelling through Conical Seamount. Their freshening relative to seawater does not result from uptake of chloride into solids during serpentinization, but it requires instead a source of H$_2$O deeper than was drilled. Whereas the presence of gas hydrates cannot be excluded, there is no direct evidence of them, and they probably are not a major source of H$_2$O. The fluids at Conical Seamount probably originate at the top of the downgoing slab, 30 km below the seafloor, by heating of the sediments and basalt of the subducted oceanic crust. The water driven off serpentinizes the overlying mantle wedge. Serpentine seamounts subsequently act as conduits through which H$_2$O, CO$_2$, hydrocarbons, and sulfur pass from the slab into the oceans.

INTRODUCTION

Because of their potential role in transferring material from the surface of the earth back into its deep interior, processes of mass transfer in subduction zones are of great interest to geochemists. While the bulk of the downgoing lithospheric plate reenters the asthenosphere during subduction, scientists have long suspected that some material, particularly from the upper part of the oceanic crust, makes its way back to the surface via processes that are poorly known at present (Peacock, 1990). To identify and quantify the various processes that occur in subduction zones, scientists of the Deep Sea Drilling Project (DSDP) and Ocean Drilling Program (ODP) have drilled in several such zones, including the Japan Trench during DSDP Leg 57, the Middle America Trench off Mexico during DSDP Leg 66 and off Guatemala during Legs 67 and 84, the Northern Barbados Ridge during ODP Leg 110, and the Peru continental margin during Leg 112.

Pore waters collected from sediments in all of these subduction zones share an unusual feature, compared with waters from marine sediments elsewhere: highly variable chloride. Whereas large increases in chloride relative to seawater were found in some of these subduction zones, large decreases of one-quarter to one-half occurred in all of them, i.e., the pore waters commonly are much fresher than seawater. As the chloride of the pore waters was typically still decreasing with depth at the maximum depth drilled, the ultimate origin of the freshened water has remained speculative. This freshening has been attributed to the presence of fossil freshwater aquifers, the formation and melting of hydrocarbon gas hydrates, membrane filtration by clays, removal of interlayer water from clay minerals andopal, and higher temperature removal of structural water from hydrous alteration minerals. Most of these explanations require the presence of a thick sedimentary sequence to be viable, and such a sequence is indeed present at all of the above subduction zones in the form of an accretionary prism, accumulated by offscraping of sediment from the top of the downgoing plate and by volcanism and erosion in the associated volcanic arc.

Here we report the discovery of greatly freshened pore waters of unusual composition from a subduction zone that lacks an accretionary prism of sediment. These pore waters were collected during Leg 125 from Conical Seamount, an active serpentine "mud volcano" in the Mariana forearc (Fig. 1). The Mariana subduction zone is nonaccretionary, and tectonic erosion has been suggested to occur in the forearc. The pore waters were sampled from silts that consist almost entirely of serpentine and contain suspended clasts of serpentinized harzburgite. Geologic evidence indicates that Conical Seamount is an active mud volcano, with a central conduit of low-density serpentine that rises buoyantly through the crust and produces periodic, unconsolidated, cold serpentine mud and debris flows ("sedimentary serpentinite") that move downslope to form the flanks of the seamount (Fryer et al., 1990). The low-chlorinity pore waters are flowing upward through the core of the seamount and exit at the summit to form chimneys of Ca-carbonate and amorphous Mg-silicate (Haggerty, 1987a; Fryer et al., 1990). We postulate that these freshened waters originate by dehydration at the top of the subducting slab 30 km below the seafloor. As they rise, they serpentinize the overlying mantle wedge and eventually mix at shallow levels with...
M. J. MOTTLE

40°N

130° E

140°

150°

Figure 1. Active plate boundaries in the Philippine Sea region. Barbed lines locate subduction zones. Basins and ridges are outlined by the 4-km bathymetric contour, except for the Izu-Bonin, West Mariana, and Mariana arcs, which are outlined by the 3-km contour. Hachured boxes show locations of Figures 2 and 3.

seawater that has reacted with ultramafic rock. To identify and characterize the reacted fluids from these two sources, we present evidence from pore waters that were collected by drilling on a second serpentinite seamount, Torishima Seamount in the Izu-Bonin forearc, that shows no sign of activity and contains only seawater-derived fluids that have reacted with peridotite.

GEOLOGIC SETTING

Sites 778 through 780 were drilled on Conical Seamount, which lies in a broad zone of forearc seamounts situated along the Mariana outer-arc high between 50 and 120 km behind the Mariana Trench axis. Conical Seamount is 20 km in diameter, 1500 m high, and nearly circular in plan view (Fig. 2). It is located about 80 km behind the trench axis and 120 km in front of the active island arc. Sites 778 and 779 were drilled on the south and southeast flanks at 3914 and 3947 mbsl, respectively. Site 780 consists of four holes drilled at the summit at 3083 to 3094 mbsl, 4.6 km from the flank sites. Drilling at all three sites penetrated mainly unconsolidated clay- to sand-sized serpentinite with suspended clasts of variously serpentinized harzburgite, dunite, and scarce metabasalt. Clasts vary in size from sand to boulders. The serpentine matrix material is sheared and foliated at the flank sites, indicating that they penetrated sedimentary serpentinite flows, but not at the summit site.

Sites 783 and 784 were drilled on Torishima Seamount on the north and west flanks at 4649 and 4901 mbsl, respectively. Torishima Seamount is 20 km wide, more than 600 m high, and semicircular in plan view, perched as it is about halfway up the flank of the outer-arc high (Fig. 3). The seamount is located about 30 km behind the axis of the Izu-Bonin Trench and 170 km in front of the island arc. The summit lies above 4100 mbsl. Drilling at both sites penetrated a glass-rich layer of silty clay to claystone and then bottomed in phacoidal sheared serpentinite silt containing clasts of serpentinized
The composition of pore waters from all five sites drilled on both serpentine seamounts is plotted vs. depth in Figures 4 through 9. These data are tabulated and the sampling and analytical methods are given by the Shipboard Scientific Party (1990a-f) and by Mottl and Alt (this volume). Relative to seawater, Mg, Ca, Sr, and Br and have low concentrations of Si, Ba, and Mn. They are enriched in the light hydrocarbons methane, ethane, and propane. The pH increases (Mottl and Alt, this volume). Relative to seawater, Mg, Ca, Sr, and Br are nearly constant with depth. Among the hydrocarbons, only methane is detectable, at very low concentrations (less than 3 mmol/kg). Many species that are elevated relative to seawater at Conical Seamount are depressed in the Torishima serpentinite, including alkalinity, sulfate, K, and B. Conversely, species that are depressed at Conical are elevated at Torishima, including Ca, Sr, Ba, and $S_{32}^4SO_4$. Species that are greatly changed at Conical, but not at Torishima, include Na (and Na/Cl), Rb, and, relative to concentrations in pore waters of the overlying claystone, Li and ammonia, as well as Cl and Br. The only elements that behave similarly at the two localities are Si, Mg, and Mn, all of which are present at very low concentrations. Pore waters from the flanks of Conical Seamount at Sites 778 and 779 are generally intermediate in composition between the summit Site 780 and the Torishima Sites 783 and 784. Site 778 resembles the Torishima sites for all species in Figures 4 through 9 except for lower chlorinity and higher Si, Br, Na/Cl, and Sr. Site 779 is intermediate for all species, except that it has the lowest ammonia and the highest Li, Na, methane, ethane, and propane of any site, and it resembles the Torishima sites in Br.

### EVIDENCE FOR UPWELLING OF PORE WATERS THROUGH CONICAL SEAMOUNT

In 1987, divers in the submersible Alvin discovered chimneys at the summit of Conical Seamount, up to 3.5 m tall, composed either of aragonite, calcite, and minor amounts of amorphous Mg-silicate, or of amorphous Mg-silicate with trace amounts of carbonate (Fryer et al., 1987; Haggerty, 1987a; Fryer et al., 1990). On sampling, one of the latter type began to emit a slow flow of cold water high in pH (9.3), bicarbonate, methane, sulfate, and H$_2$S relative to seawater. Relative to carbonates of seawater origin that form in fracture zone serpentinites, the chimney carbonates were found to be depleted in Sr and S relative to seawater. Chlorinity and Br are nearly constant with depth. Among the hydrocarbons, only methane is detectable, at very low concentrations (less than 3 mmol/kg). Many species that are elevated relative to seawater at Conical Seamount are depressed in the Torishima serpentinite, including alkalinity, sulfate, K, and B. Conversely, species that are depressed at Conical are elevated at Torishima, including Ca, Sr, Ba, and $S_{32}^4SO_4$. Species that are greatly changed at Conical, but not at Torishima, include Na (and Na/Cl), Rb, and, relative to concentrations in pore waters of the overlying claystone, Li and ammonia, as well as Cl and Br. The only elements that behave similarly at the two localities are Si, Mg, and Mn, all of which are present at very low concentrations. Pore waters from the flanks of Conical Seamount at Sites 778 and 779 are generally intermediate in composition between the summit Site 780 and the Torishima Sites 783 and 784. Site 778 resembles the Torishima sites for all species in Figures 4 through 9 except for lower chlorinity and higher Si, Br, Na/Cl, and Sr. Site 779 is intermediate for all species, except that it has the lowest ammonia and the highest Li, Na, methane, ethane, and propane of any site, and it resembles the Torishima sites in Br.

### COMPOSITION OF PORE WATERS FROM CONICAL AND TORISHIMA SEAMOUNTS

The composition of pore waters from all five sites drilled on both serpentine seamounts is plotted vs. depth in Figures 4 through 9. These data are tabulated and the sampling and analytical methods are given by the Shipboard Scientific Party (1990a-f) and by Mottl and Alt (this volume).

Pore waters from Site 780 on the summit of Conical Seamount are among the most unusual ever sampled in the deep sea. They have the highest pH ever measured in a deep-sea sediment, up to 12.6. They have lower chlorinity than at any other site drilled in a subduction zone, up to 57% lower than that in seawater. They show a large increase in alkalinity with depth, to 26 times the value in seawater, and are highly enriched in the light hydrocarbons methane, ethane, and propane. The sulfate concentration increases to 1.7 times that in seawater, and one sample from 2.7 mbsf in Hole 780D yielded 2 mmol/kg H$_2$S. $S_{32}^4SO_4$ decreases from the seawater value of +20.5 to about 13.5$S_{32}^4SO_4$ as the sulfate concentration increases (Mottl and Alt, this volume). Relative to seawater, the pore waters at Conical Seamount summit are also enriched in ammonia, K (× 1.5), Rb (×5.6), and B (× 10). They are highly depleted in Li, Mg, Ca, Sr, and Br and have low concentrations of Si, Ba, and Mn. They are also depleted in Na, and their Na/Cl ratio increases to nearly twice that in seawater. All of these changes take place near the seafloor, in the upper few meters of the serpentinite silt, and most persist to at least 130 mbsf, the maximum depth sampled.

Pore waters from serpentinite silt at Sites 783 and 784 on Torishima Seamount contrast greatly with those from the summit of Conical Seamount (Table 1). The pH elevation from 9 to 10 is not nearly as high as that at the summit of Conical Seamount. There is no freshening relative to seawater; chlorinity and Br are nearly constant with depth. Among the hydrocarbons, only methane is detectable, at very low concentrations (less than 3 mmol/kg). Many species that are elevated relative to seawater at Conical Seamount are depressed in the Torishima serpentinite, including alkalinity, sulfate, K, and B. Conversely, species that are depressed at Conical are elevated at Torishima, including Ca, Sr, Ba, and $S_{32}^4SO_4$. Species that are greatly changed at Conical, but not at Torishima, include Na (and Na/Cl), Rb, and, relative to concentrations in pore waters of the overlying claystone, Li and ammonia, as well as Cl and Br. The only elements that behave similarly at the two localities are Si, Mg, and Mn, all of which are present at very low concentrations. Pore waters from the flanks of Conical Seamount at Sites 778 and 779 are generally intermediate in composition between the summit Site 780 and the Torishima Sites 783 and 784. Site 778 resembles the Torishima sites for all species in Figures 4 through 9 except for lower chlorinity and higher Si, Br, Na/Cl, and Sr. Site 779 is intermediate for all species, except that it has the lowest ammonia and the highest Li, Na, methane, ethane, and propane of any site, and it resembles the Torishima sites in Br.

### Table 1. Comparison of pore water in serpentinite silts from two forearc seamounts.

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<th>Torishima Forearc Seamount</th>
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as well as from seawater, and methane was found in fluid inclusions along with aromatic compounds, long-chain paraffins, naphthenes, and acetate ions (Haggerty, 1989). The presence of aromatic compounds in the chimneys and of ethane and propane along with methane in the pore waters indicates a thermogenic origin for the organic compounds. The presence of acetate ions limits the temperature range in the source region of the organics to less than 150°C. Similar methane-derived carbonate chimneys have been described from the Oregon subduction zone (Kulm et al., 1986; Ritger et al., 1987; Kulm and Suess, 1990). The summit of Conical Seamount is well below the aragonite compensation depth (Berger, 1970), and the bottom waters there are certainly undersaturated with aragonite (Li et al., 1969; Berner and Horjo, 1981).

Video photography taken during drilling of the A hole at summit Site 780 revealed no chimneys. The large gradients in pore-water composition in the surficial sediments, however, are almost certainly maintained by upflow of the freshened water through the central serpentine conduit of the seamount. These gradients cannot originate by reaction in the surficial sediments because the gradients are too steep and they exist for nearly every species measured and because the chemistry and mineralogy of the serpentine sediments are too simple. In the absence of upflow, the gradients would be quickly erased by diffusion to and from the overlying seawater. The steepness of the gradients suggests upflow velocities of a few millimeters per year (Fig. 10), much slower than the focused flow through the chimneys. The large variation in composition at a given depth among the four holes drilled within 150 m of each other indicates a patchy distribution of flow. Temperature logging yielded a value of 13.5°C at 58 mbsf in Hole 780C, vs. only 3.7°C extrapolated for the same depth from discrete measurements in Hole 780D, and 1.52°C for bottom seawater. This suggests that warmer water from the bottom of the deepest Hole 780C at 164 mbsf flowed upward in the hole after it was drilled (Shipboard Scientific Party, 1990a-f).

In addition to the chimneys, upflow is almost certainly also responsible for extensive precipitation of aragonite needles in the serpentine silt at and near the seafloor, leading to carbonate enrichment in these zones at both the summit and flank sites (Fig. 11). Precipitation presumably occurs where the carbonate-rich, but Ca-

Figure 4. Composition of pore waters from cores at Sites 778 (squares), 779 (diamonds), 780 (x's), 783 (right-pointing triangles), and 784 (left-pointing triangles), compared with the composition of surface seawater (+'s on x-axis). For Sites 783 and 784, samples from claystone are shown by open triangles and samples from serpentine by solid triangles.
poor, upwelling solutions mix with Ca-rich seawater. Site 784 on the Torishima Seamount flank shows a similar zone of carbonate enrichment near the top of the serpentine section in the form of dispersed calcite, rather than as discrete needles of aragonite. This enrichment suggests that Torishima Seamount may also have once vented solutions rich in carbonate ion and that the original aragonite has recrystallized.

**ORIGIN OF PORE WATERS AT CONICAL SEAMOUNT**

Two end-member hypotheses for the origin of the low-chlorinity pore waters at Conical Seamount must be considered: (1) the fluids originate mainly as seawater, and chloride is taken up into solid phases during alteration in greater proportion than is H₂O and (2) the fluids do not originate mainly as seawater but contain a large component of freshwater that originates by some other process. Whatever the process that causes the chlorinity decrease at Site 780, it must be occurring at a greater depth than was penetrated by drilling. The steep gradients in chlorinity and other dissolved species in the surficial sediments require upflow of pore water, as noted earlier. The persistence of these surficial changes in pore-water composition to the maximum depth drilled indicates that the upwelling fluid originates at a greater depth and that the upwelling rate overwhelms the reaction rate within the drilled interval.

**Seawater-Rock Interaction and Chloride Uptake**

Chloride uptake into solid phases during serpentinization has been reported, although the conditions under which this phenomenon can occur and the nature of the chlorine-rich phases are unclear. Janecky and Seyfried (1986) performed experiments reacting seawater and Mg-free and Mg- and sulfate-free seawater with peridotites at 200°C and 300°C and a pressure of 500 bars. They found that dissolved chloride behaved conservatively when seawater was the reactant solution but that significant chloride was taken up into partially serpentinized harzburgite when the starting solution lacked Mg. The distinctive feature of the Mg-free-seawater experiments was the high pH in the solutions, which reached 11.5 in the longest experiment. This experiment was performed at 300°C, 500 bars, and with a water/rock mass ratio of 10, and ran for 714 days. The loss of chloride...
from solution was 27 mmol/kg, indicating that the altered harzburgite must have contained about 10,000 ppm, or 1 wt%, chloride that it had picked up from solution during the experiment. Janecky and Seyfried (1986) were unable to determine in what solid phase or phases this chloride resided. They considered several possibilities: (1) a Mg-Fe-chloride mineral like iowaite, which has been reported from serpentinites (Kohls and Rodda, 1967) and which occurs in minor amounts in some samples from the Conical Seamount sites (Fryer and Mottl, this volume; Heling and Schwarz, this volume); (2) an Fe-hydroxide-chloride phase, such as was suggested to occur in serpentinites by Rucklidge (1972) and Rucklidge and Patterson (1977); (3) a Mg-hydroxide-chloride phase, Mg₂(OH)₃Cl, produced during experiments...
Figure 8. Composition of pore waters from cores at Sites 778, 779, 780, 783, and 784. Symbols are the same as in Figure 1.
Figure 9. Composition of pore waters from cores at Sites 778, 779, and 780, in mmol/kg of pore water. Symbols are the same as in Figure 1. Methane concentrations at Sites 783 and 784 are even lower than those at Site 778, and ethane is absent at these three sites.
The large decreases in chlorinity of the pore waters at Site 780, relative to seawater, are balanced only one-quarter to one-third by loss can be accounted for. The other one-quarter is balanced about decreased Na. When the nearly complete removal of Mg and Ca from the altered solids from this experiment contained 1.59 wt% NaO, although for some reason this Na did not appear in the bulk analysis of the altered rock, which yielded only 0.07 wt%. The large decreases in chlorinity of the pore waters at Site 780, relative to seawater, are balanced only one-quarter to one-third by decreased Na. When the nearly complete removal of Mg and Ca from solution is included as well, only three-quarters of the negative charge loss can be accounted for. The other one-quarter is balanced about equally by the large gains in sulfate and alkalinity. Relative to seawater, the increase in sulfate is proportionally about equal to that in K. If the gains in sulfate and K are caused mainly by loss of H$_2$O from seawater to the serpentinized rocks, and the rocks gain on average about 12 wt% H$_2$O, then each gram of seawater would have reacted with and hydrated 2 to 3 g of fresh rock. If the rock also took up chloride under these conditions, it would have to gain about 4300 ppm, or 0.43 wt% chloride, to account for the chloride decrease observed in the interstitial waters at Site 780. This is well within the range of uptake that apparently occurred in the experiments of Janecky and Seyfried (1986), indicating that chloride uptake below the drilled interval could account for the freshening observed in the pore waters at Site 780. The high concentrations of alkalinity, methane, and ammonia could also be explained by this model, as there are adequate concentrations of carbonate and organic carbon and nitrogen in the solids, as measured within the drilled interval (Shipboard Scientific Party, 1990a-f), to account for the concentrations in the pore waters.

However, the very different manner in which charge is balanced at Site 780 vs. in the experiments, noted above, strongly suggests that some other process has dominated at Conical Seamount summit. In addition, the assumption in the above model that sulfate is conservative is highly unlikely in view of the large decrease in S$^{4+}$SO$_4$ at Site 780 (Mottl and Alt, this volume). The chloride-uptake model by itself cannot explain many of the other chemical changes in the Conical summit pore waters relative to seawater, including the large decreases in Br, Li, Na, Ca, Sr, and Ba, which would have to be taken up into the rock along with chloride during alteration, and the large increases in B and Rb, which would have to be leached from the rock. These directions of net transport between solids and solution are very different from what was observed at Torishima Seamount, where there is no evidence of any fluid other than reacted seawater. Neither chlorinity nor Br decreases in pore waters from the serpentine section at Torishima; in fact, they increase slightly with depth. Methane is present only in very low concentrations, and ethanol and propane are absent. Values of δ$_{18}$O increase due to sulfate reduction, probably by bacteria (Mottl and Alt, this volume), Li, Na, and Rb change little, whereas Ca, Sr, Ba, K, B, alkalinity, and sulfate change in the opposite direction to what was found at the Conical Seamount summit. The pore waters at Torishima can be taken as end-member examples of harzburgite-seawater interaction at relatively low temperatures and are very different from the pore waters that are flowing upward through the Conical Seamount summit (Table 1).

### Evidence from Drilling in Other Subduction Zones

Low-chlorinity interstitial waters have been sampled from drill holes in other subduction zones, including the Japan Trench, the Middle America Trench off Guatemala and Mexico, the Barbados accretionary prism, and the Peru continental margin. In the Japan Trench, at Sites 438 and 439 drilled during DSDP Leg 57, the chlorinity of the pore waters decreases with depth to half the value in seawater at 900 to 1000 mbsf. Moore and Gieskes (1980) suggested that this low-chlorinity water originated from a fossil aquifer.

In the Middle America Trench off Guatemala, the chlorinity of the pore waters is lower than that of seawater by up to 54% at 337 mbsf. Moore and Gieskes (1980) suggested that this low-chlorinity water originated from a fossil aquifer.
Figure 11. Concentration of calcium carbonate in the upper 30 m of serpentine silt at Sites 778, 779, and 780. The low concentrations typical of the interval between 20 to 30 mbsf persist to the maximum depths drilled.

as in the overlying sediments, originated by upward migration of gas and fresh water (von Huene and Lee, 1983) from subducted sediment.

On the Northern Barbados Ridge, drilled during Leg 110, chlorinity is lower than that in seawater by up to 28% at Site 674, but by less than 12% at all other sites drilled there. This decrease was tentatively attributed by the Shipboard Scientific Party (1988a) to an ultrafiltration process, in which clay minerals act as semipermeable membranes, selectively passing H₂O but restricting the transport of ionic species. On the basis of additional data, Gieskes et al. (1990) attributed the decrease to dehydration of clays and to deeper prograde diagenetic and metamorphic dehydration reactions. All of these processes were postulated to occur deeper than the maximum depth drilled; the freshwater produced is thought to move upward along faults and other permeable pathways, and especially along the shallowly inclined décollement surface.

On the Peru continental margin, chlorinity increases greatly with depth at five sites drilled during ODP Leg 112 and decreases at the other five. At Site 679, these decreases are up to 38% relative to seawater, but they are less than 18% at the other four sites. Several processes may have contributed to these decreases in chlorinity, including decomposition of hydrocarbon gas hydrates, membrane filtration, dewatering of clays and opal, and the possible presence of fossil freshwater (Suess et al., 1988; Elderfield et al., 1990; Kastner et al., 1990). Several chlorinity maxima and minima at Sites 682, 683, 685, and 688 clearly result from formation and decomposition, respectively, of gas hydrates at corresponding depths in the drill holes. As in the case of the Barbados accretionary prism, however, the larger-scale decrease with depth indicates that the major source of freshwater lies below the maximum depth drilled. This freshwater is thought to have risen upward along permeable channels consisting of unconformities, coarse-grained sedimentary units, and conjugate and intersecting fracture zones (Shipboard Scientific Party, 1988b).

Possible Sources of Low-Chlorinity Water at Conical Seamount

The 57% decrease in chlorinity at 27 mbsf in Hole 780D is larger than any observed in sediment pore waters above subduction zones elsewhere.
It has also been sampled at much shallower depths below the seafloor, as a result of upflow of pore waters through the summit of Conical Seamount. The explanations suggested for freshening of pore waters in other subduction zones must be considered for Conical Seamount.

No direct evidence of hydrocarbon gas hydrates was found in any of the cores during Leg 125, nor is there convincing seismic evidence of a bottom-simulating reflector that might indicate their presence. Given the temperature gradient of about 0.038 °C measured in Hole 780D, however, the phase diagram for methane hydrate (Kvenvolden and McMenamin, 1980) suggests that this phase should be stable from the seafloor to a depth of about 600 mbsf beneath the summit Site 780, below which temperatures exceeding about 24 °C would cause any gas hydrate present to melt. Gas hydrate will form, of course, only if methane concentrations are sufficiently high, in excess of about 58 mmol/kg, according to Claypool and Kaplan (1974). The highest concentration measured at Site 780 was only 2.7 mmol/kg, and at flank Site 779, 36.6 mmol/kg. These concentrations are minima, however, because of probable degassing during core recovery, which can be avoided only by use of a core barrel capable of maintaining in situ pressure. The large maximum in light hydrocarbons measured in Hole 779A at 227 mbsf, and other maxima deeper in the same hole, may result from melting of gas hydrate during core recovery, but there is no correlation between the hydrocarbon maxima and chlorinity decreases. Thus, the evidence is inconclusive. Gas hydrates could be present in a zone between the maximum depth drilled and about 600 mbsf, at which depth they might continuously melt, supplying freshwater and dissolved hydrocarbon gases that may upwell to the summit if diluted somewhat by seawater during the ascent to prevent refreezing. Note, however, that methane concentrations are up to 12 times higher at flank Site 779 than at summit Site 780, yet the decrease in chlorinity is much smaller at Site 779: only 9% lower than seawater, compared with 57% lower at Site 780. Unless methane and water derived from melting of gas hydrates are somehow separated from each other during ascent through the upper few hundred meters of
Conical Seamount, this discrepancy suggests that decomposition of gas hydrates is probably not the dominant source of freshwater on Conical Seamount.

The low abundance of clay minerals in the rocks and sediments recovered from Conical Seamount, along with the absence of a large volume of sedimentary substrate in the Mariana forearc in general, makes membrane filtration through clay minerals an unlikely mechanism for generating freshwater there. Conical Seamount is too far from land for any fossil freshwater aquifers to be present.

Dehydration of hydrous minerals at depth beneath Conical Seamount is therefore the most likely mechanism for generating freshwater. The chemistry of the pore waters sampled at Sites 778, 779, and 780, especially the high pH, indicates that the drilled interval is undergoing serpentinization at the present time. Thus, any dehydration of previously formed hydrous minerals would have to be taking place well below the maximum depth drilled during Leg 125 and would have to supply enough H$_2$O both for serpentinization at shallower depths and to feed the freshened pore waters upwelling through Conical Seamount.

Possible dehydration reactions that could supply fluids in a subduction zone, discussed by Peacock (1990, and references therein), include (1) diagenetic transformation of opal-A, including expulsion of interlayer water at about 30° to 80°C; (2) expulsion of interlayer water from smectite and its transformation to illite at about 150° to 180°C; (3) metamorphic dehydration of sedimentary minerals, mainly clays, beginning at about 250° to 300°C; and (4) metamorphic dehydration of chloride, amphibole, and other hydrous phases in the basaltic oceanic crust, beginning at about 450° to 500°C. These dehydration reactions would be accompanied by breakdown of hydrocarbons to methane in sediments at about 60° to 130°C and by decarbonation of sedimentary carbonates. As noted earlier, the organic compounds in the pore waters of Conical Seamount are thermogenic, and the presence of acetate ions in fluid inclusions in the carbonate chimneys there limits the temperature range in the source region to less than 150°C for at least some of the organic molecules (Haggerty, 1989). As there is little pelagic sediment in the Mariana forearc crust, the most likely source of sedimentary material is that at the top of the subducting slab 30 km below Conical Seamount, where subducted basaltic crust also is available. The thermal models of Peacock (1990) for a subducting slab suggest temperatures in the range of 150° to 600°C at the pressure at this depth of about 10 kbar. Water driven from these subducted oceanic sediments and basement rocks could serpentinize the overlying wedge of mantle, resulting in serpentinite mud volcanoes such as Conical Seamount, as well as the unusual pore fluids upwelling through its summit.

**Evidence for Mixing of Fluids from Several Sources**

If the pore waters upwelling through Conical Seamount summit do in fact originate largely from dehydration of the subducting oceanic crust, then the relative simplicity of their serpentine host matrix, both chemically and mineralogically, should provide us with the clearest picture to date of what such a fluid component looks like. From the composition of the pore waters at Site 780, we can infer that it is rich in H$_2$O, C both as dissolved carbonate species and as light hydrocarbons and other organic molecules, and S as both sulfate and H$_2$S. It is depleted in $^{34}$S relative to seawater. By comparison with pore waters from Torishima Seamount that result from harzburgite-seawater interaction, we can also infer that it is rich in B, Na, K, and Rh.

For other species, the picture is less clear. The pore waters from the summit of Conical Seamount have low concentrations of Mg, Mn, and Si, but so do those from serpentinite at Torishima Seamount. They also have much lower Li, Ca, Sr, and Ba than at Torishima, but this probably results from the much higher concentrations of carbonate alkalinity and sulfate associated with the deep-fluid component at Conical, and from the much higher pH. Ammonia is somewhat higher at Conical than in the Torishima serpentines for reasons that are unclear. The organic carbon content of the serpentinite sediments on both seamounts is generally in the range of 0.1 to 0.3 wt% (Shipboard Scientific Party, 1990a–f). Sulfate decreases greatly in the Torishima pore waters within the serpentinite-rich section, whereas its $^{34}$S/$^{32}$S ratio increases, presumably as a result of bacterial sulfate reduction using this organic matter. Although the alkalinity increase expected from this reaction is prevented by a high concentration of Ca leached from the rock, ammonia increases as expected. The increase in ammonia at Conical Seamount summit may result from the same process of sulfate reduction; some H$_2$S is present there, but any decrease in sulfate is masked by the sulfate supplied with the deep-fluid component.

In summary, the deep-fluid component, as typified by Conical Seamount summit Site 780, is characterized by high carbonate alkalinity, methane, ethane, propane, sulfate, H$_2$S, B, Na, K, and Rh; and low chlorinity, Br, Li, Ca, Sr, Ba, and $^{34}$S/$^{32}$S. The component derived by reaction of seawater with peridotite, as typified by the serpentinite section at Torishima Seamount Sites 783 and 784, is characterized by very low methane; absence of ethane and propane; low carbonate alkalinity, sulfate, B, and K; slightly lower Li and Rh; little-changed chlorinity, Br, Na, and Na/C1; and high Ca, Sr, Ba, and $^{34}$S/$^{32}$S. Both components have high pH and ammonia and low Mg, Si, and Mn (Table 1). To a first approximation, Conical Seamount flank Sites 778 and 779 are mixtures of these two fluid components, Site 779 being richer in the deep-fluid component than Site 778.

In addition to the general trends with depth, however, many of the dissolved species in pore waters at Conical Seamount show more complex profiles that include maxima and minima. This is especially true at flank Site 779, but it also occurs in the lowermost one or two samples in Holes 778A and 780D. Pore waters from closely spaced holes at Sites 779 and 780 also show large differences in composition at similar shallow depths beneath the sea floor. Both types of variation can be caused by mixing of fluids from different sources in a complex plumbing system within Conical Seamount. Besides the two endmember fluids described above, locally derived seawater may have reacted with mafic rocks in the vicinity of the seamount. Such reaction could explain the lower pH and higher Si at Site 778, which has ubiquitous small mafic clasts suspended in the serpentinite (Johnson, this volume) along with the ultramafic clasts that occur at all of the sites. Reaction with these mafic clasts could also explain the relatively high Sr concentrations and Sr/Ca ratios in the pore waters from Site 778 (Fig. 12). Site 779, with its complex profiles, may exhibit the effects of mixing of all three fluids at various depths. At least three separate fluids may be required for explaining the multiple maxima and minima of the various dissolved species, as they generally do not correlate with one another.

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