



## Subsurface structure of a submarine hydrothermal system in ocean crust formed at the East Pacific Rise, ODP/IODP Site 1256

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[1] ODP/IODP Hole 1256D penetrates an in situ section of ocean crust formed at the East Pacific Rise, through lavas and sheeted dikes and ~100 m into plutonic rocks. We use mineralogy, oxygen isotopes, and fluid inclusions to understand hydrothermal processes. The lavas are slightly altered at low temperatures (<150°C) to phyllosilicates and iron oxyhydroxides, with a stepwise increase in grade downward to greenschist minerals in the upper dikes. This resulted from generally upwelling hydrothermal fluids in the dikes mixing with cooler seawater solutions in the lavas, also producing minor metal sulfide mineralization

in the upper dikes. Alteration grade increases downward in the dikes, with increasing recrystallization to amphibole and loss of metals at higher temperatures (>350°C up to ~600°C). Intrusion of gabbro bodies into the lower dikes resulted in contact metamorphism to granoblastic hornfels at 850°C–900°C, representing a thermal boundary layer between the axial melt lens and the overlying hydrothermal system. Downward penetration of hydrothermal fluids led to rehydration of granoblastic dikes and plutonic rocks at ~800°C down to <300°C. Fluid inclusion and oxygen isotope data show that vein quartz formed at ~300°C to >450°C from hydrothermal fluids that were affected by supercritical phase separation. Fluids had variable salinities and were enriched in <sup>18</sup>O (+0.4‰ to +3.5‰) relative to seawater, similar to seafloor vent fluids. Dike margins are brecciated and mineralized, suggesting hydrothermal activity coeval with magmatism. Anhydrite formed mainly in the upper dikes when partly reacted seawater fluids were heated as they penetrated deeper into the system. Low-temperature alteration of the volcanic section continued as cold seawater penetrated along fluid pathways, forming minor iron oxyhydroxides in the rocks. Hydrothermal processes at Site 1256 fit with current models whereby greenschist alteration of dikes at low water/rock ratios is overprinted by fracture-controlled alteration and mineralization by upwelling hydrothermal fluids, a conductive boundary layer above gabbroic intrusions, leaching of metals from dikes and gabbros in the deep “root zone,” and stepped thermal and alteration gradients in the basement. The Site 1256 section, however, is intact and retains recharge effects (anhydrite), allowing an integrated view of processes in the subsurface.

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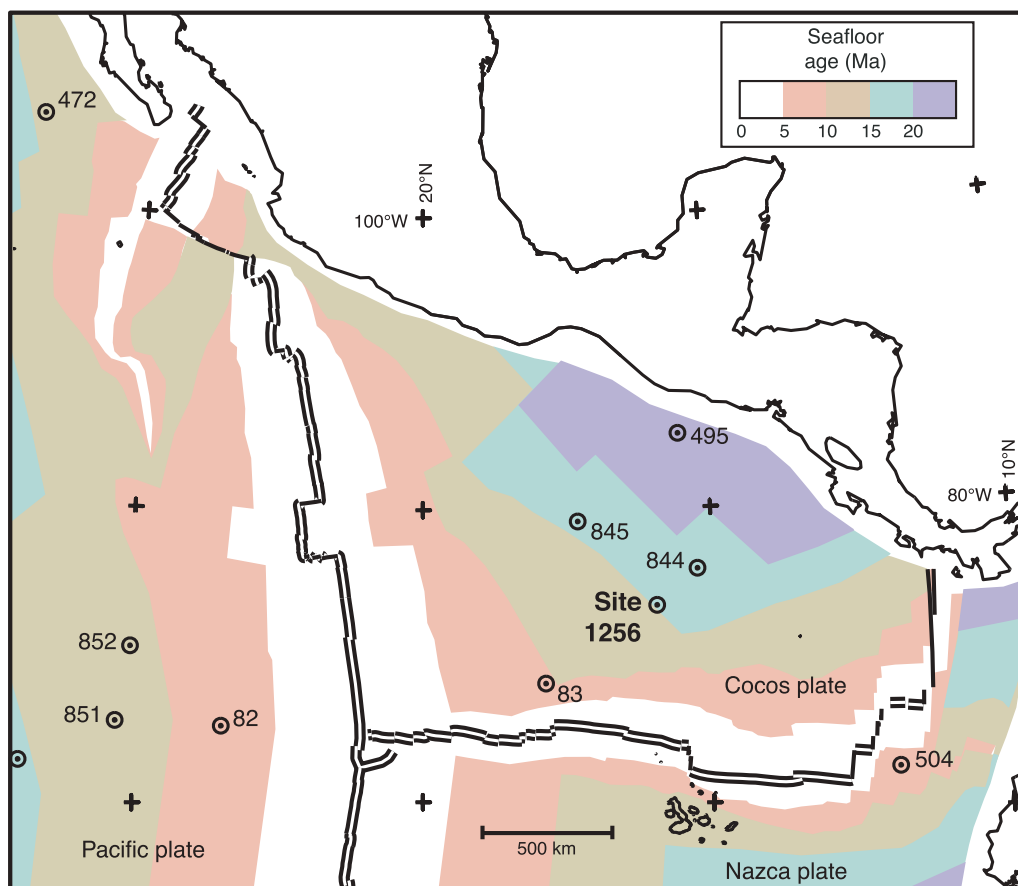
## 1. Introduction

[2] Submarine hydrothermal systems exert important influences on the composition of seawater and the crust, on the physical properties of the lithosphere, and during recycling in subduction zones [Edmond *et al.*, 1979a; Hofmann and White, 1982; Tatsumi, 1989] (see also summary by Alt [1995]). Mid-ocean ridge hydrothermal systems are studied via diverse approaches, all of which are essential to develop an integrated understanding of these systems. However, direct sampling of ocean crust remains critical to understand the subsurface structure and hydrothermal processes in these systems, as well as to test remote geophysical observations. Ophiolites have provided important inspiration for understanding hydrothermal processes at mid-ocean ridges (MOR) [e.g., Spooner and Fyfe, 1973], but evidence suggests that the hydrothermal alteration of ophiolites is anomalous compared to crust in the major ocean basins [e.g., Bickle and

Teagle, 1992; Alt and Teagle, 2000], and hence the study of in situ oceanic crust remains essential.

[3] Hydrothermal alteration of several sections of upper ocean crust formed at intermediate and fast spreading rates has been well studied. These include tectonic windows at Hess Deep and Pito Deep, formed at fast spreading rates on the East Pacific Rise (EPR), and deep drilling of Hole 504B, formed at intermediate spreading rate in the eastern Pacific [Alt *et al.*, 1986, 1996; Teagle *et al.*, 2003; Gillis *et al.*, 2001; Gillis, 2008; Heft *et al.*, 2008].

[4] In this paper we present results of a study of hydrothermal alteration in a ~1.5 km section of upper oceanic crust formed at a fast spreading rate at the EPR, as sampled by drilling at ODP/IODP Site 1256. We document the secondary mineralogy, textural relationships, and distribution of hydrothermal effects in this section, and present fluid inclusion and oxygen isotope analyses of vein minerals. These data are used to understand the overall thermal structure and evolution of hydro-



**Figure 1.** Location of Site 1256 in the eastern Pacific. Isochrons at 5 Myr intervals and selected DSDP and ODP sites are indicated. The wide spacing of 10–20 Myr isochrons to the south reflects the superfast (>200 mm/yr) full spreading rate (modified from *Wilson et al.* [2003]).

thermal alteration at this site. We evaluate our results for Site 1256 in terms of current models for circulation and alteration of the upper oceanic crust at intermediate and fast spreading rates.

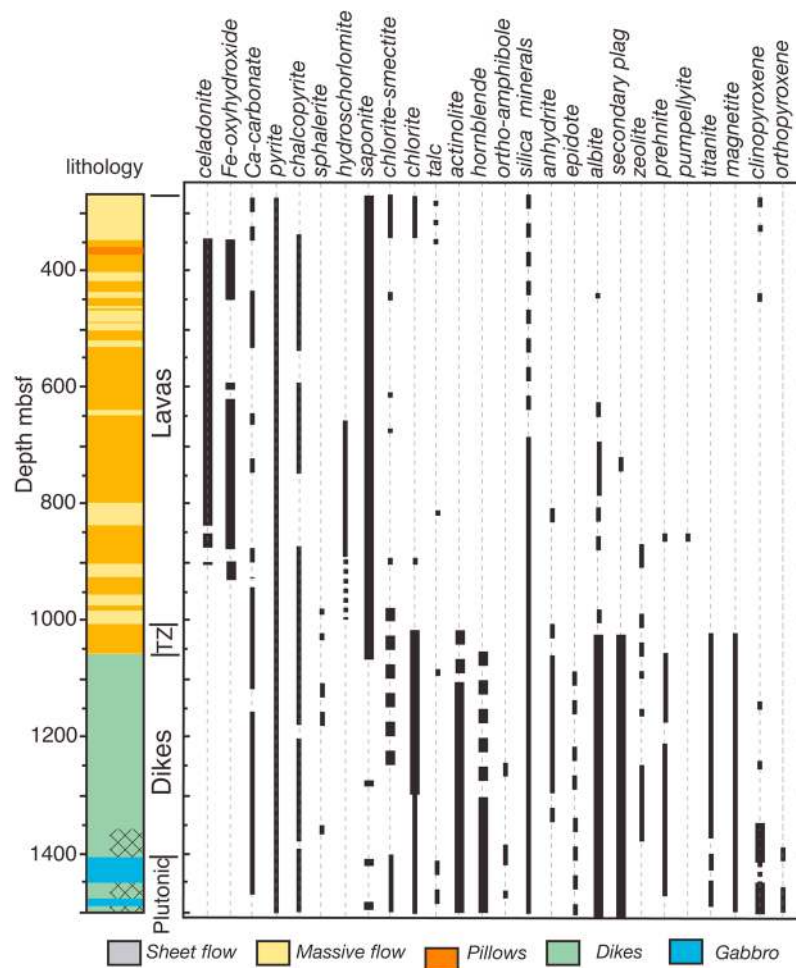
## 2. Geological Setting and Methods

### 2.1. Hole 1256D Background and Lithostratigraphy

[5] Site 1256 is located at 6.736°N, 91.934°W in the eastern Pacific, in crust that formed at a superfast spreading rate at the EPR ~15 Myr ago (Figure 1) [*Teagle and Expedition 309/312 Scientists*, 2006; *Wilson et al.*, 2006]. Drilling took place on ODP Leg 206 [*Wilson et al.*, 2003] and IODP Expeditions 309 and 312 [*Teagle and Expedition 309/312 Scientists*, 2006], and core descriptions are presented in those reports. Hole 1256C penetrates the 250 m sediment section and extends 72 m into basement. Hole 1256D, ~30 m to the south, was cased

though the sediments and ~20 m into basement [*Wilson et al.*, 2003]. Coring started at 26 m sub-basement (msb), 276 m below seafloor (mbsf), and extends to a depth of 1507.1 mbsf (1257.1 msb; Figure 2) [*Teagle and Expedition 309/312 Scientists*, 2006; *Wilson et al.*, 2006].

[6] The uppermost basement comprises a ~100 m thick sequence dominated by a ponded lava flow >75 m thick in Hole 1256D but only 32 m thick in nearby Hole 1256C. The immediately underlying lavas include sheet and massive flows, and minor pillow flows. The upper 284 m of the volcanic section is interpreted to have formed off axis [*Teagle and Expedition 309/312 Scientists*, 2006; *Tominaga et al.*, 2009]. Sheet flows and massive lavas erupted at the ridge axis make up the remaining extrusive section down to 1004 mbsf, where a lithologic transition is marked by sub-vertical intrusive contacts and mineralized breccias. Below 1061 mbsf, subvertical intrusive



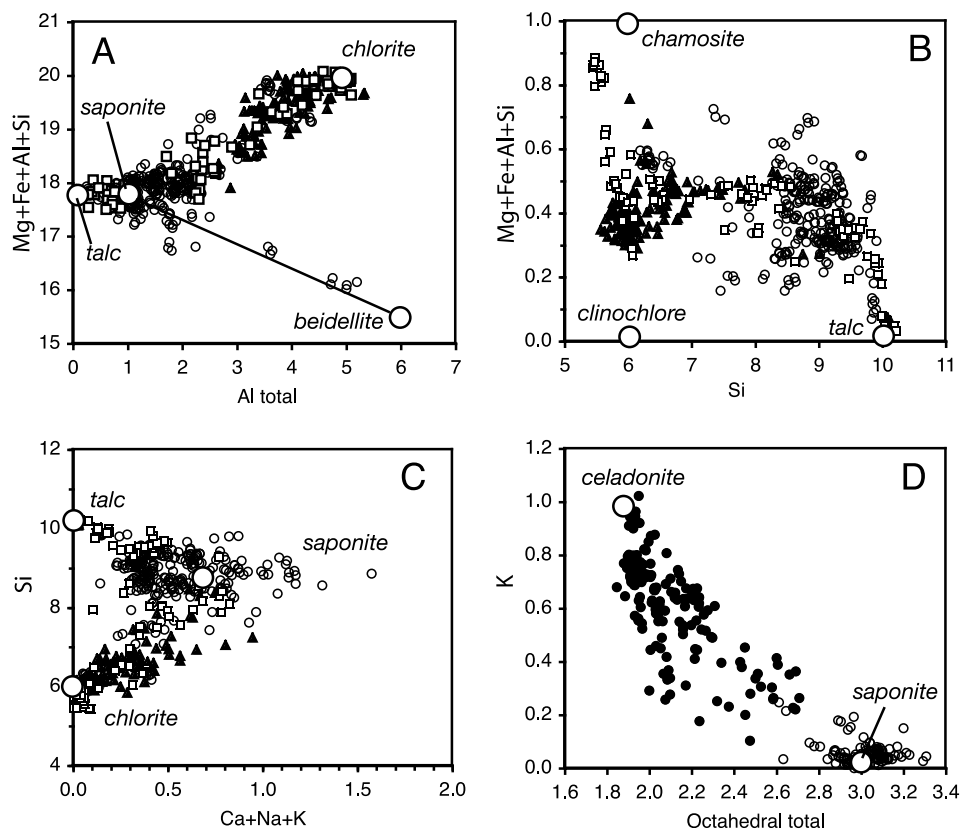
**Figure 2.** Lithostratigraphy and distribution of secondary minerals versus depth in Hole 1256D. Major secondary phases are shown as thick lines, minor and trace phases are shown as finer lines, and crosshatching indicates grano-blastic dikes. Hornblende/actinolite distribution based on textures and electron probe analyses. Shipboard lithostratigraphy from *Teagle and Expedition 309/312 Scientists* [2006]. Mineral distributions based on study of shipboard thin sections [Wilson et al., 2003; *Teagle and Expedition 309/312 Scientists*, 2006] and this study.

contacts indicate a ~350 m thick sheeted dike complex [Teagle and Expedition 309/312 Scientists, 2006; Tominaga et al., 2009]. Some rocks have doleritic textures, and many are cut by subvertical dikes with common brecciated and mineralized chilled margins. The lower ~60 m of the sheeted dike complex is recrystallized to granoblastic textures by contact metamorphism.

[7] The plutonic complex begins at 1407 mbsf. Two gabbro bodies were penetrated, with a 52 m thick upper gabbro separated from a 24 m thick lower gabbro by a 24 m screen of dikes that was recrystallized by contact metamorphism (Figure 2). The upper gabbro comprises gabbros, oxide gabbros, quartz-rich oxide diorites and small trondjhemite

dikelets. The lower gabbro comprises gabbro, oxide gabbro, and lesser orthopyroxene bearing gabbro and trondjhemite [Teagle and Expedition 309/312 Scientists, 2006]. Contacts with the dike screen are intrusive, with partly resorbed, stopped dike clasts within the margins of the lower gabbro. The lowermost rock recovered from the hole is a highly altered actinolite-bearing basaltic dike that post-dates the intrusion of the lower gabbro.

[8] Flows and dikes show a wide range of fractionation, from MgO ~4.5 to 10 wt % (mostly 5.8–9.0) [Teagle and Expedition 309/312 Scientists, 2006; Wilson et al., 2006]. The ranges of most major and many minor element concentrations are similar to those of the northern East Pacific Rise [Teagle and



**Figure 3.** Phyllosilicate compositions. Formulas calculated as chlorites (layer charge = 56) in Figures 3a–3c. (a) Talc, saponite, and beidellite end-members are indicated. Smectite–chlorite occupies trend from saponite to chlorite [after *Schiffman and Fridleifsson, 1991*]. (b) Chlorite classification diagram illustrating variable compositions of chlorites (between chamosite and clinochlore) and trend to higher Si contents of smectite–chlorite, saponite, and talc. (c) Trends from smectite at variable and high interlayer cation contents to chlorite at low contents of Si and interlayer cations and to talc at high Si and low interlayer contents. (d) Trend from celadonite at high K content (0.8–1.0) to saponite at low K and high octahedral contents results from variable mixtures of celadonite and smectite. Formulas calculated as smectites (layer charge = 22). Circles, lavas (solid, celadonite; open, saponite); filled triangles, dikes; open squares, gabbro.

*Expedition 309/312 Scientists, 2006; Wilson et al., 2006*], indicating similar processes at the superfast spreading ridge that formed Site 1256 and the modern EPR. Gabbro compositions span a range similar to the flows and dikes, but are on average more primitive (mean MgO ~8 wt % versus 7.2 wt % for flows and dikes).

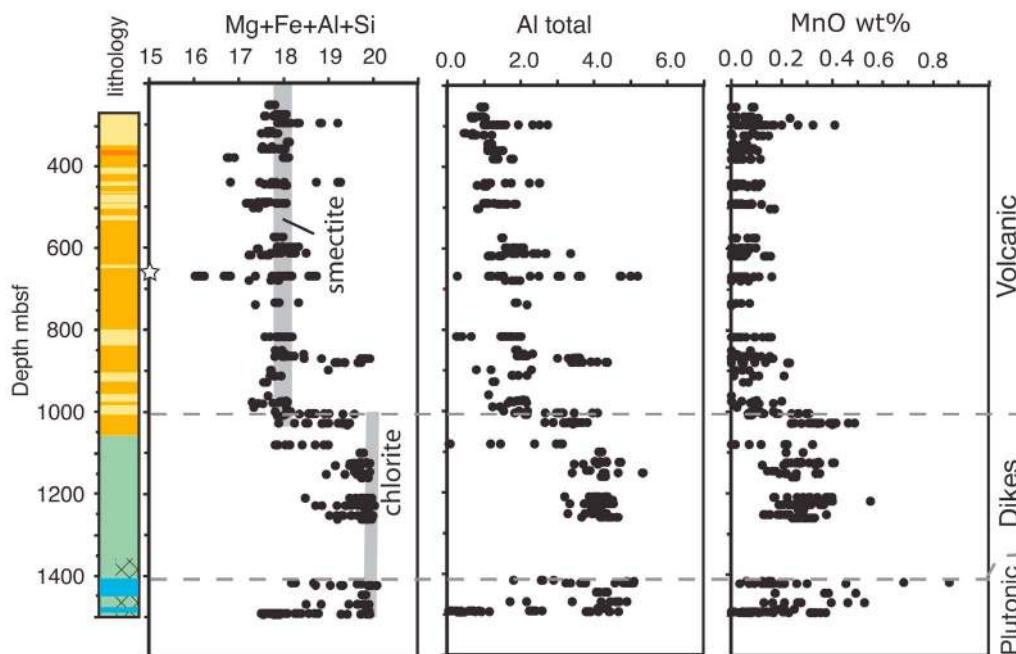
## 2.2. Methods

[9] Alteration petrography and secondary mineralogy were studied by transmitted and reflected light microscopy, with percent secondary phases estimated visually. Compositions of secondary minerals were determined by electron microprobe at The University of Michigan and Ifremer (Brest, France), with methods given by *Alt and Laverne [2006]*. Electron probe data presented in Figures 3–6, 9,

and 10 are given in the auxiliary material.<sup>1</sup> Selected phyllosilicate veins were analyzed by X-ray diffraction at the University of Michigan. Oriented powder mounts were scanned with a Scintag automated diffractometer using Cu K $\alpha$  radiation, and a scan rate of 2° 2 $\theta$ /min, after air drying and glycolation treatments.

[10] Secondary minerals for oxygen isotope analyses were separated from veins by crushing and hand picking, and purity and mineralogy were checked by X-ray diffraction and visual inspection. Oxygen isotope analyses were carried out at the University of Michigan by reacting samples with BrF<sub>5</sub> in externally heated nickel reaction vessels [*Clayton*

<sup>1</sup>Auxiliary materials are available at <ftp://ftp.agu.org/apend/gc/2010gc003144>.



**Figure 4.** Compositions of phyllosilicates versus depth. Volcanic section is dominated by smectite, with local smectite-chlorite (high Mg + Fe + Al + Si and Al contents at ~290, 450, and ~880 mbsf). Beidellite in the intensely altered basalt at 648 mbsf (star) is characterized by low Mg + Fe + Al + Si and high Al. Stepwise change from saponite to chlorite occurs at top of lavas-dikes transition (at 1004 mbsf). Smectite-chlorite in dikes and plutonic section indicated by low Al and Mg + Fe + Al + Si contents relative to chlorite. Talc is present at Al = 0 wt %. Lithology as in Figure 2.

and Mayeda, 1963], and converting O<sub>2</sub> to CO<sub>2</sub> gas by reaction with heated carbon rods. Oxygen isotope ratios were measured using a Finnigan Delta-S mass spectrometer. NBS 28 quartz run with sample unknowns had a mean  $\delta^{18}\text{O}$  value of 9.6‰ ( $\pm 0.2$ ,  $1\sigma$ ) compared to the accepted value of 9.58‰. Anhydrite data are corrected using NBS 127 BaSO<sub>4</sub>, which was run concurrently with unknowns, and sulfate oxygen data were normalized to a value of 9.34‰ for NBS 127 [Gonfiantini *et al.*, 1995].

[11] Microthermometric measurements on fluid inclusions in quartz were made at University of Leeds, using a Linkam THMSG 600 heating-freezing stage. Samples include four quartz veins ( $\pm$ minor chlorite, prehnite, amphibole, laumontite, sulfides) in dikes and gabbros, one quartz-filled vug from the dike section, one plagiogranite and one tonalite vein, and groundmass quartz from an oxide gabbro. Freezing temperature and melting temperature of the solid were measured first, followed by heating measurements, including daughter crystal dissolution ( $T_m$ ) and homogenization temperature ( $T_h$ ). Precision and accuracy were  $\pm 0.2^\circ\text{C}$  below  $0^\circ\text{C}$  and  $\pm 2^\circ\text{C}$  above  $0^\circ\text{C}$ . Salinity was estimated using freezing point depression assuming a pure NaCl solution and is reported as wt % NaCl equivalent [Bodnar, 2003].

Using measured salinities, estimated pressures, and data from Bodnar [2003], homogenization temperatures of two-phase liquid-vapor fluid inclusions were corrected for assumed hydrostatic pressure (350–410 bars, see auxiliary material), yielding trapping temperatures ( $T_t$ ). Pressures were estimated based on depth to the seafloor and depth below seafloor.

### 3. Results

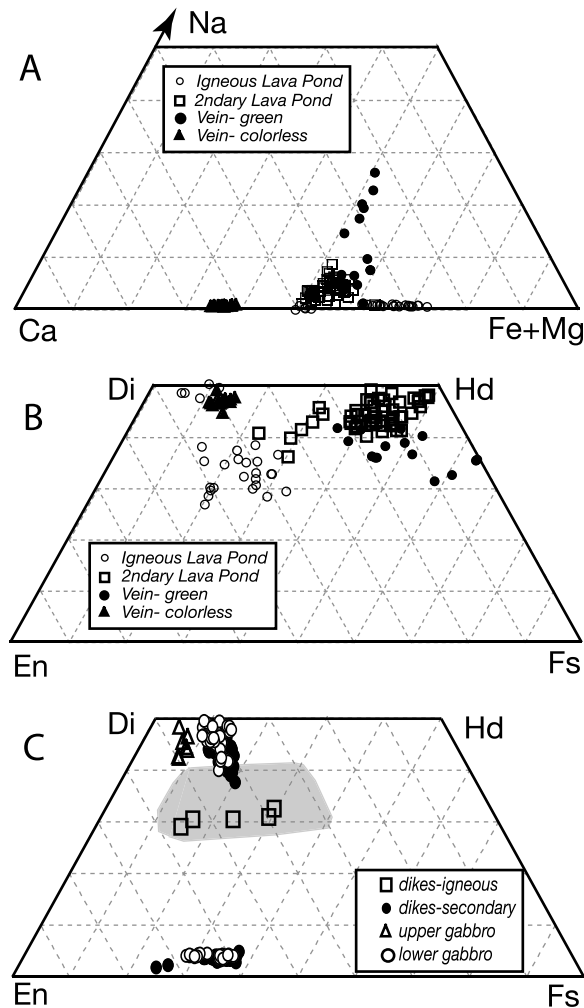
#### 3.1. Hydrothermal Alteration

[12] Documentation of hydrothermal alteration is divided into sections below based on mineralogy and lithology: the volcanic section; transition zone and sheeted dikes; the granoblastic dikes and dike screens; and plutonic rocks.

##### 3.1.1. Volcanic Section

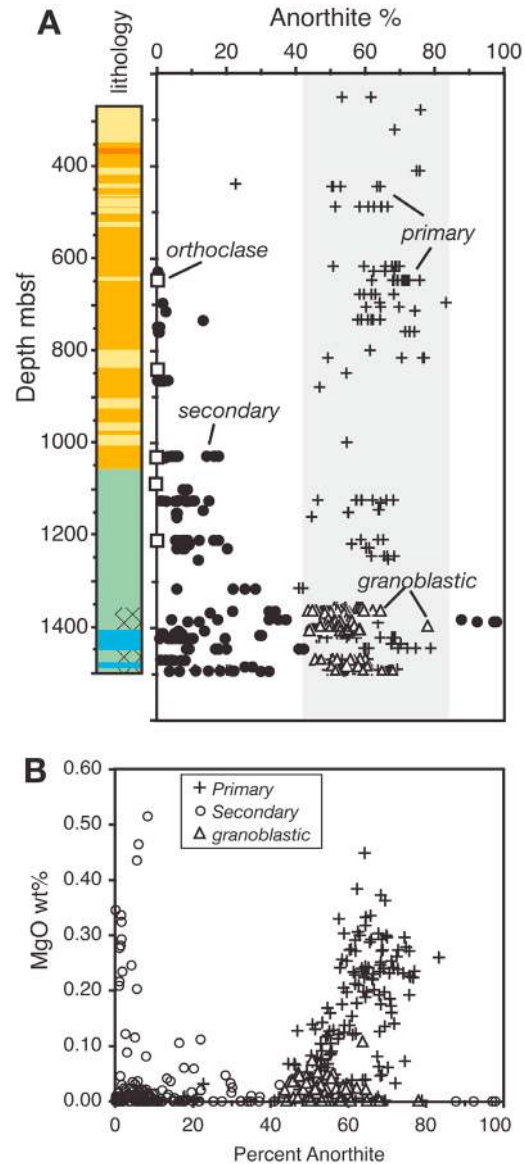
###### 3.1.1.1. Background Alteration

[13] The volcanic section exhibits a pervasive background alteration where the rocks are dark gray and slightly to moderately altered (2%–20%, typically



**Figure 5.** Compositions of pyroxenes. (a and b) Pyroxenes in lavas and sheeted dikes. Igneous clinopyroxenes in the lava pond have compositions similar to the rest of the lavas and dikes but range to high-Ca diopside. Secondary hedenbergite reaction rims in the lava pond have higher Fe, Ca, and Na compared to igneous augites. Green vein clinopyroxenes in lavas (at 445.60 mbsf, sample 26R2, 18–21) and dikes (at 1245.75 mbsf, 168R3, 78–82) are aegerine-augite, with elevated Na and Fe compared to igneous augite. Colorless vein clinopyroxene in a dike (1143.54 mbsf, 146R2, 48–52) is diopside. (c) Compositions of pyroxenes in granoblastic dikes and gabbros. Field for igneous augites in lavas and dikes shown for comparison (gray field [from *Dziony et al.*, 2008]). Secondary pyroxenes in the granoblastic dikes are enstatite and diopsidic clinopyroxene. Also shown are pyroxene compositions in gabbros. Clinopyroxenes and orthopyroxenes in the lower gabbro have textures and compositions similar to those in the granoblastic dikes (see section 3.1.4).

~10%). Saponite replaces olivine and interstitial glass and fills pore space, with common disseminated secondary sulfides. Basaltic glass at flow margins is fresh to variably altered to saponite and trace phillipsite. The most common phyllosilicate in the volcanic section is saponite (Mg-rich, trioctahedral smectite). Saponites have noninterlayer



**Figure 6.** Compositions of feldspars. (a) Percent anorthite versus depth. Locations of orthoclase are indicated. Shading indicates compositions of igneous plagioclase (excluding sodic rims on some crystals). (b) MgO contents of igneous feldspars decrease with decreasing %An. Secondary plagioclase mainly has low MgO contents, except for some low-An plagioclase. Lithology as in Figure 2.

cation (Mg + Fe + Al + Si) contents of ~17.5–18 and elevated interlayer cation contents (Figure 3). Compositions range toward talc, with low Al and interlayer cation contents (Figure 3). Variable amounts of chlorite are present locally, as indicated by elevated contents of Mg + Fe + Al + Si and decreasing Si and interlayer cations (Figures 3 and 4), and by X-ray diffraction. Dark gray mm- to cm-scale alteration patches around pores filled with saponite and silica minerals in the coarser-grained portions of thick flows are more altered, and contain ~50% secondary minerals. The central portion of the thick ponded lava at the top of the basement contains late magmatic/hydrothermal effects, including secondary Na-hedenbergite reaction rims on primary augite (Figure 5) [Laverne *et al.*, 2007].

[14] A significant change occurs below 627 mbsf, where secondary albite, saponite, and chlorite-smectite locally replace the cores of plagioclase phenocrysts and microphenocrysts (Figures 4 and 6). Below 660 mbsf, titanomagnetite is more altered and hydroschorlomite appears (Ti-rich hydrogarnet) [Laverne *et al.*, 2006].

[15] A unique 41 cm interval at 648 mbsf is highly (80%–90%) altered. Olivine, clinopyroxene, and plagioclase are replaced by beidellite, celadonite, K-feldspar and iron oxyhydroxide, imparting blue-green and brick red colors to the rock. These minerals plus quartz and carbonate also fill vugs and pore space. The Al-rich smectite beidellite appears in trends toward decreasing Mg + Fe + Al + Si and increasing total Al in Figure 3.

[16] Pumpellyite is present in a >70 cm interval of basalt at 864 mbsf, where it fills veins and pores with albite, smectite/talc, celadonite, laumontite, minor prehnite, calcite, and titanite, and locally replaces olivine.

### 3.1.1.2. Veins and Vein-Related Alteration

[17] Veins make up 1% by volume of the volcanic section with an average of 23 veins/m of core. Saponite-bearing veins are the most abundant, commonly containing minor pyrite. Celadonite ± iron oxyhydroxide veins are a factor of 10 less abundant. Compositions of celadonite range from celadonite to glauconite [Alt and Laverne, 2006]. Trends of decreasing K with increasing tetrahedral Al and octahedral Mg indicate the presence of variable mixtures with smectite (Figure 3). Also present in veins are silica minerals (opal, chalcedony, and quartz) and small amounts of calcium

carbonate. Early green aegerine-augite in a vein at 445.60 (Figure 5) probably formed during initial seawater penetration into cooling basalt [Laverne, 1987]. Petrographic relationships indicate early formation of celadonite ± iron oxyhydroxides in veins, followed by saponite, and then silica minerals and carbonate. Anhydrite is present locally in veins (around 825 mbsf; Figure 2), and generally postdates associated quartz, pyrite and local calcite.

[18] Alteration halos along veins make up 3.5% of the core (Figure 7a). These are present throughout most of the volcanic section, but are more abundant at 350–450 and 630–820 mbsf [Wilson *et al.*, 2003]. The halos are typically 1 to 10 mm wide, but range up to 30 mm wide. Dark halos are characterized by celadonite + minor later saponite, but iron oxyhydroxide can impart a reddish color to the rock. In rare cases, narrow brown alteration halos contain iron oxyhydroxides as the only secondary mineral.

[19] Below 850 mbsf halos are characteristically 0.1 to 2 mm greenish to gray zones along saponite veins, where the rock is moderately to highly altered to saponite or chlorite-smectite. Below ~950 mbsf, halos are characterized by pyrite and minor chalcopryrite disseminated in the rock along narrow (<1 mm) alteration halos.

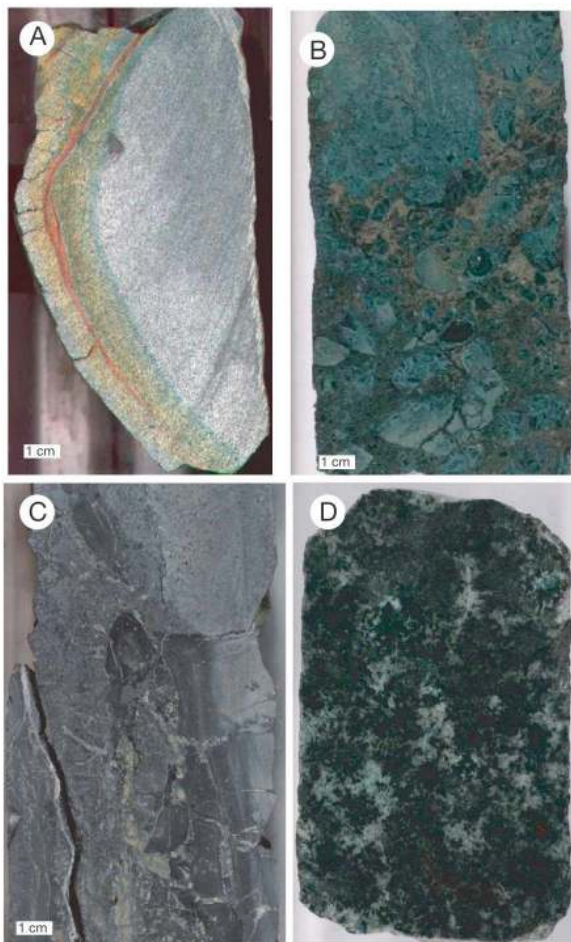
### 3.1.1.3. Breccias

[20] Breccias make up 2% of the core, although wireline logging suggests they are more abundant [Tominaga *et al.*, 2009]. Hyaloclastites make up 90% of breccias and consist of fresh and altered glass clasts in a matrix of saponite and trace quartz. Glass fragments are variably altered to palagonite, saponitic smectite, and trace pyrite. Local basalt breccias are cemented by saponite, local celadonite, chalcedony, red jasper, quartz, pyrite, and minor anhydrite.

### 3.1.2. Transition Zone and Sheeted Dikes

[21] The transition zone and sheeted dikes are characterized by greenschist and subgreenschist minerals in rocks and veins. Hydrothermal alteration is manifest as (1) pervasive but variable background recrystallization; (2) alteration halos along veins; (3) highly altered patches around vugs; and (4) local metal sulfide mineralization in breccias, dike margins, and veins.





**Figure 7.** Characteristic alteration features in lavas and dikes. (a) Mixed dark + brown alteration halo along 2 mm vein of celadonite + iron oxyhydroxide + late saponite (sample 206-1256D-61R-1, 50–59 cm, 678.55 mbsf). Alteration halos are mainly dark gray to green and black but range to brown and reddish. The degree of alteration is similar to the host rock. (b) Mineralized hyaloclastite breccia (at 1028 mbsf). Clasts of aphyric cryptocrystalline basalt and light green altered glass in a matrix of chlorite, albite, and pyrite (brownish in photo), with local sphalerite, quartz, and K-feldspar (309-1256D-122R-1, 68–79 cm). (c) Brecciated and mineralized dike margin. Dike on right intrudes host dike on left, with contact characterized by brecciation, flow structures, and multiple veins. (309-1256D-140R-1, 41–59 cm, 1114.12 mbsf). Host rock is altered to actinolite, chlorite, titanite, and albite. A vein of chlorite + quartz + pyrite + magnetite + trace chalcopyrite is cut by a 1 mm vein of quartz + chalcopyrite + magnetite + actinolite. The host rock is cut by numerous 0.2 mm veins of quartz, sulfides, chlorite, and actinolite. (d) Gabbro exhibiting typical patchy texture. Darker areas of ophitic clinopyroxene enclosing plagioclase are less altered than coarser grained, more felsic material between, imparting patchy appearance to rock.

### 3.1.2.1. Background Alteration

[22] Beginning at the top of the lithologic transition zone (1004 mbsf), rocks are variably (1%–80%; commonly 30%–50%) altered to greenschist and subgreenschist assemblages. Clinopyroxene commonly appears cloudy as the result of partial replacement by fibrous amphibole plus tiny (<1–2  $\mu\text{m}$ ) grains of magnetite (Figures 8a and 8b). Plagioclase phenocrysts and groundmass are variably replaced by albite-oligoclase ( $\text{An}_{0-20}$ ), chlorite, and chlorite-smectite. Small amounts of prehnite, anhydrite, epidote, laumontite and rare calcite occur locally as grains or patches partly replacing plagioclase. Olivine and interstitial areas are variably recrystallized to chlorite-smectite, chlorite, and minor amphibole. Titanomagnetite is partly replaced by titanite, and disseminated pyrite and chalcopyrite are common.

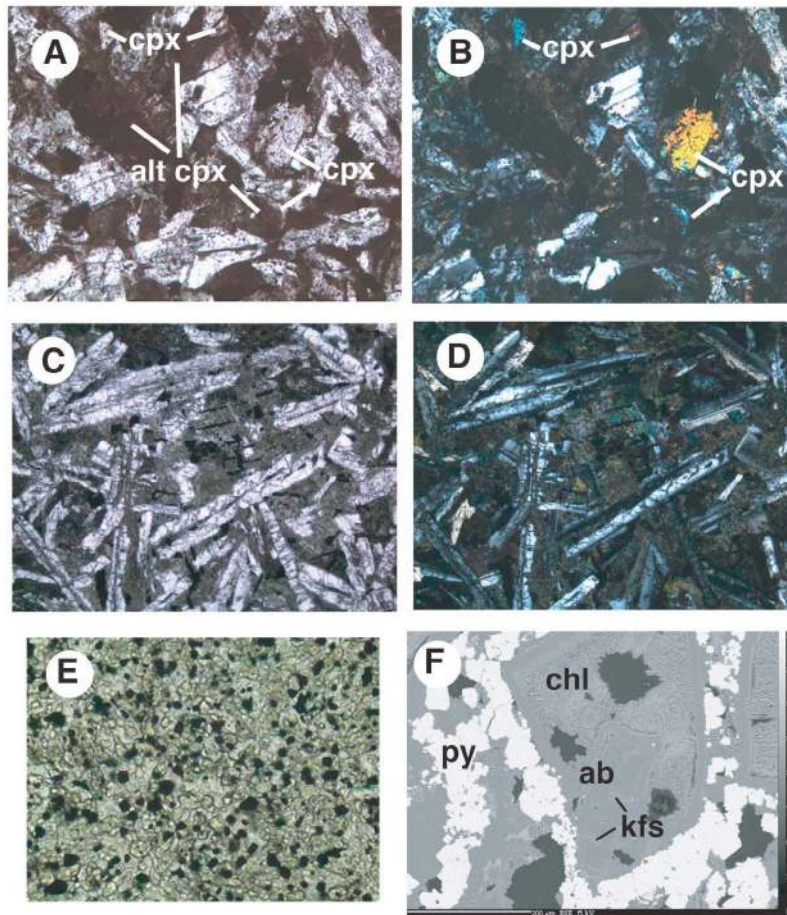
[23] Chlorite is a major alteration phase above ~1300 mbsf, but amphibole becomes dominant below this depth (Figure 2). Below 1300 mbsf clinopyroxene is more intensely recrystallized to amphibole + magnetite, plagioclase is partly altered to albite-oligoclase and minor anhydrite, and titanomagnetite is more highly altered to titanite (Figures 8c and 8d).

[24] Compositions of amphiboles down to 1350 mbsf range from actinolite and minor ferroactinolite to Mg hornblende, and include local Fe-Mg amphiboles (Figures 9 and 10). Data are sparse in the upper dikes (in part because of the fibrous nature of amphibole and patchy replacement of pyroxene), but suggest actinolite in the transition zone and a wider range of composition in the dikes. Local Fe-Mg amphiboles replace interstitial material (e.g., at 1252 mbsf) and pyroxene (e.g., along with actinolite and Mg-hornblende along a chlorite + quartz + titanite vein at 1256 mbsf; Figure 10).

[25] Epidote compositions are variable, with  $\text{Fe}/\text{Fe} + \text{Al} = 0.17\text{--}0.33$  (Figure 10). Secondary plagioclase in the dikes is albite to oligoclase (mostly  $\text{An}_0\text{--}\text{An}_{20}$ , but ranging up to  $\text{An}_{28}$  at 1315 mbsf; Figure 6). Secondary plagioclase tends to have low MgO contents, although some albite has high MgO (Figure 6).

### 3.1.2.2. Veins, Vein-Related Alteration Halos, and Isolated Patches

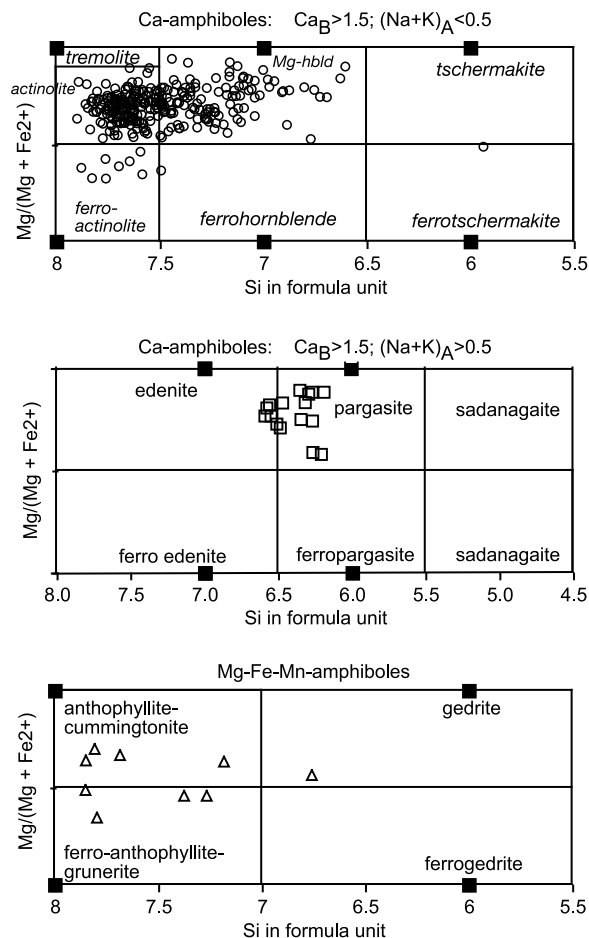
[26] Veins comprise 0.9% of the dikes, at a frequency of ~30 veins/m. Chlorite is the most common vein filling, although quartz, pyrite, chalcopyrite, actinolite, prehnite, laumontite, and calcite are also common. Secondary clinopyroxene was identified



**Figure 8.** Thin section photomicrographs of characteristic alteration effects in sheeted dikes. (a) Background alteration in upper dikes, with clinopyroxene partly corroded and altered to “dusty” clinopyroxene/actinolite+magnetite on the left and with unaltered clinopyroxene on the right (sample 312-1256D-176R-1, 8–12 cm; 1276.18 mbsf) (plane-polarized light; field of view (FOV) = 1.12 mm). (b) Same as Figure 8a but with cross-polarized light. (c) Background alteration in lower dikes, highly altered basaltic dike with clinopyroxene replaced by greenish amphibole ± magnetite (black) (sample 312-1256D-189R-1, 68–69 cm; 1334.58 mbsf) (plane polarized light, FOV = 4.5 mm). (d) Same as Figure 8c but with cross-polarized light. (e) Granoblastic dike, with equigranular assemblage of plagioclase, clinopyroxene, magnetite, and ilmenite (plane-polarized light; FOV = 0.6 mm) (sample 312-1256D-205R-1, 10–14 cm; 1382.20 mbsf). (f) BSE image of mineralized breccia in Figure 7b. Glass clast is replaced by chlorite, albite, K-feldspar (lighter gray), and pyrite, in a matrix of the same minerals. Scale at bottom left is 200  $\mu\text{m}$ .

in two veins (Figure 5): at 1143.54 mbsf colorless diopside occurs along the walls of a vein; at 1245.75 mbsf green aegerine-augite occurs with titanite, hydro-andradite, local epidote, and late anhydrite. Vein relationships indicate a general sequence of mineral formation: (1) chlorite, amphibole, titanite, and pyrite veins can be overprinted by (2) veins of quartz, chlorite, epidote, pyrite, chalcopyrite and rare sphalerite, and (3) late vein minerals include anhydrite, prehnite, laumontite, and calcite [Teagle and Expedition 309/312 Scientists, 2006]. Anhydrite is present locally in veins below 1000 mbsf, but is most abundant from 1100 to 1200 mbsf (Figure 2).

[27] Rock texture and proximity to veins and pore space influence recrystallization of the rocks, with medium grained rocks being more intensely recrystallized than finer grained rocks. Alteration halos, 1–11 mm wide, along veins make up ~4% of the core, and are more intensely recrystallized (10%–100%) than the host rock. Alteration patches comprise 1.4% of the core. Millimeter-scale pores are filled with quartz, chlorite, actinolite needles, pyrite, prehnite and laumontite (and rare sphalerite), and are surrounded by cm-scale, highly altered (60%–100%), alteration halos.



**Figure 9.** Compositions of amphiboles in Hole 1256D. Circles, calcic amphiboles; squares, pargasite-edenite; triangles, Fe-Mg amphiboles.

### 3.1.2.3. Sulfide Mineralization in Breccias, Dike Margins, and Veins

[28] A prominent feature at 1027 mbsf, near the top of the transition zone, is the presence of a ~3 m thick mineralized hyaloclastite breccia (Figures 7b and 8f). Glass shards are replaced by chlorite, albite-oligoclase, K-feldspar, and titanite, and are cemented by abundant quartz and pyrite with lesser chalcopryrite, needles of actinolite, and local concentrations of sphalerite and anhydrite. Minor late calcite is also present in veins.

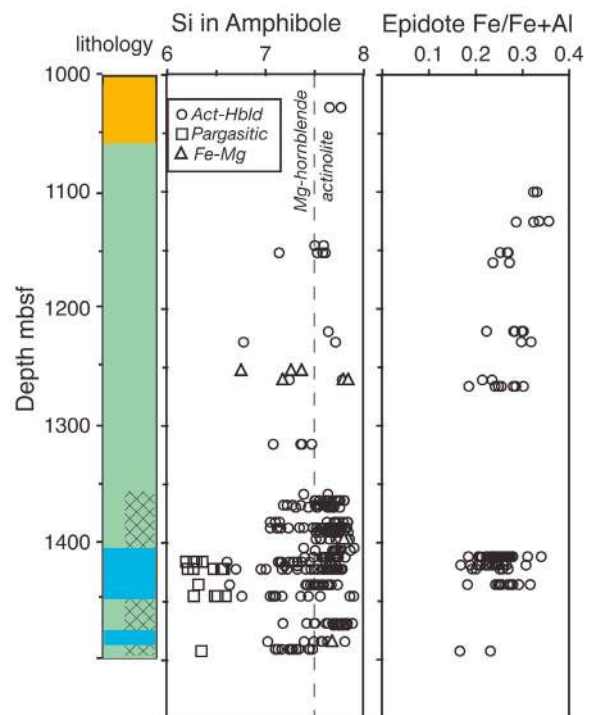
[29] Dike margins are commonly brecciated and mineralized, with the host rocks extensively replaced by chlorite, amphibole, titanite, albite and local K-feldspar (Figure 7c). The brecciated chilled margins are cut by veins of chlorite, amphibole, quartz, magnetite, and pyrite, common chalcopryrite and sphalerite, and local anhydrite. Magnetite is

partly replaced by small amounts of hematite. Similar mineral assemblages are common in veins throughout the sheeted dikes above 1350 mbsf.

### 3.1.3. Granoblastic Dikes and Dike Screens

[30] Below 1348 mbsf, the dikes are variably recrystallized to microcrystalline, granular aggregates of secondary clinopyroxene, plagioclase, magnetite and ilmenite, ±orthopyroxene, ±quartz, and with late amphibole replacing pyroxenes. These rocks contain 0.5–1 mm completely recrystallized patches within variably recrystallized host rocks, and are referred to as the granoblastic dikes (Figure 8e) [Teagle and Expedition 309/312 Scientists, 2006; Koepke et al., 2008]. Rocks from the dike screens between and below the gabbro bodies exhibit alteration identical to that of the granoblastic dikes overlying the gabbros.

[31] Recrystallized plagioclase has compositions of An<sub>44</sub>–An<sub>65</sub>, within the range of igneous plagioclase (Figure 6). MgO contents are low, however, similar to secondary plagioclase (Figure 6). Compared to igneous augites, secondary clinopyroxenes are relatively Ca-rich and range to diopside com-



**Figure 10.** Amphibole and epidote formula compositions versus depth. Amphibole classification from Figure 9. Lithology as in Figure 2.

**Table 1.** Pyroxene Compositions and Calculated Equilibrium Temperatures<sup>a</sup>

Sample	Depth	Area	PX	SiO <sub>2</sub>	Al <sub>2</sub> O <sub>3</sub>	TiO <sub>2</sub>	FeO	MnO	MgO	CaO	Na <sub>2</sub> O	K <sub>2</sub> O	Si	Al	Ti	Fe3+	Fe2+	Mn	Mg	Ca	Na	K	T (°C)
194R1-36	1358.26	4	cpx	51.62	1.17	0.51	11.58	0.30	14.06	20.22	0.28	0.01	1.95	0.05	0.01	0.00	0.37	0.01	0.79	0.82	0.02	0.00	876
194R1-36	1358.26	4	opx	52.53	0.41	0.31	22.74	0.56	21.59	1.69	0.04	0.01	1.97	0.02	0.01	0.00	0.71	0.01	1.21	0.07	0.00	0.00	864
194R1-36	1358.26	5	cpx	51.89	0.77	0.35	11.10	0.32	14.08	20.40	0.25	0.01	1.97	0.03	0.01	0.00	0.35	0.01	0.80	0.83	0.02	0.00	884
194R1-36	1358.26	5	opx	52.40	0.52	0.29	22.86	0.56	21.52	1.65	0.04	0.00	1.97	0.02	0.01	0.00	0.72	0.01	1.21	0.07	0.00	0.00	884
196R1-0	1363.70	1	cpx	51.76	1.29	0.47	10.21	0.32	14.41	20.40	0.27	0.01	1.95	0.06	0.01	0.00	0.32	0.01	0.81	0.83	0.02	0.00	893
196R1-0	1363.70	1	opx	52.86	0.71	0.33	21.20	0.53	22.45	1.78	0.04	0.01	1.97	0.03	0.01	0.00	0.66	0.01	1.25	0.07	0.00	0.00	893
196R1-0	1363.70	2	cpx	51.58	1.39	0.50	10.58	0.29	14.19	20.13	0.27	0.01	1.95	0.06	0.01	0.00	0.34	0.01	0.80	0.82	0.02	0.00	863
196R1-0	1363.70	2	opx	52.65	0.75	0.33	21.33	0.52	22.39	1.76	0.04	0.01	1.97	0.03	0.01	0.00	0.67	0.01	1.25	0.07	0.00	0.00	863
196R1-0	1363.70	3	cpx	52.21	1.11	0.41	10.11	0.31	14.22	20.77	0.26	0.01	1.97	0.05	0.01	0.00	0.32	0.01	0.80	0.84	0.02	0.00	899
196R1-0	1363.70	3	opx	52.76	0.70	0.31	21.39	0.50	22.07	1.68	0.03	0.01	1.98	0.03	0.01	0.00	0.67	0.01	1.23	0.07	0.00	0.00	899
197R1-2	1367.52	1	cpx	51.92	1.32	0.53	11.18	0.34	14.41	19.55	0.30	0.01	1.96	0.06	0.01	0.00	0.35	0.01	0.81	0.79	0.02	0.00	909
197R1-2	1367.52	1	opx	52.81	0.57	0.29	21.58	0.51	22.25	1.72	0.02	0.00	1.974	0.03	0.01	0.00	0.67	0.01	1.24	0.07	0.00	0.00	909
197R1-2	1367.52	2	cpx	51.53	1.36	0.53	11.07	0.33	14.28	19.61	0.30	0.01	1.95	0.06	0.02	0.00	0.35	0.01	0.81	0.80	0.02	0.00	872
197R1-2	1367.52	2	opx	52.47	0.70	0.31	21.58	0.52	21.99	1.67	0.04	0.00	1.972	0.03	0.01	0.00	0.68	0.01	1.23	0.07	0.00	0.00	872
198R1-46	1369.46	A	cpx	52.25	0.53	0.36	11.17	0.45	14.07	20.25	0.21	0.02	1.98	0.02	0.01	0.00	0.35	0.01	0.79	0.82	0.02	0.00	867
198R1-46	1369.46	A	opx	52.61	0.61	0.28	22.51	0.67	20.99	1.70	0.04	0.00	1.983	0.03	0.01	0.00	0.71	0.02	1.18	0.07	0.00	0.00	895
198R1-46	1369.46	D	cpx	51.90	0.93	0.36	10.95	0.39	14.05	20.34	0.22	0.00	1.97	0.04	0.01	0.00	0.35	0.01	0.79	0.83	0.02	0.00	924
198R1-46	1369.46	D	opx	52.10	0.53	0.29	22.77	0.69	20.97	1.77	0.02	0.01	1.97	0.02	0.01	0.00	0.72	0.02	1.19	0.07	0.00	0.00	874
206R1-13	1387.03	4	cpx	50.91	1.45	0.51	10.51	0.39	14.47	20.03	0.29	0.01	1.939	0.06	0.01	0.00	0.33	0.01	0.82	0.82	0.02	0.00	899
206R1-13	1387.03	4	opx	52.47	0.72	0.32	20.86	0.67	22.41	1.82	0.03	0.01	1.97	0.03	0.01	0.00	0.65	0.02	1.25	0.07	0.00	0.00	895
209R1-8	1396.58	B	cpx	51.73	1.25	0.51	11.32	0.35	14.28	19.36	0.30	0.01	1.96	0.06	0.01	0.00	0.36	0.01	0.81	0.79	0.02	0.00	872
209R1-8	1396.58	B	opx	52.92	0.65	0.35	21.28	0.58	22.18	1.87	0.03	0.00	1.97	0.03	0.01	0.00	0.66	0.01	1.23	0.07	0.00	0.00	874
209R1-8	1396.58	D	cpx	51.92	1.04	0.43	10.23	0.33	14.36	20.46	0.29	0.01	1.96	0.05	0.01	0.00	0.32	0.01	0.81	0.83	0.02	0.00	899
209R1-8	1396.58	D	opx	52.54	0.66	0.32	21.31	0.54	21.97	1.82	0.03	0.00	1.97	0.03	0.01	0.00	0.67	0.01	1.23	0.07	0.00	0.00	899
212R1-17	1404.27	F	cpx	51.45	1.04	0.52	12.15	0.39	13.89	19.24	0.25	0.01	1.96	0.05	0.01	0.00	0.39	0.01	0.79	0.79	0.02	0.00	800
212R1-17	1404.27	F	opx	51.97	0.53	0.32	23.29	0.56	20.74	1.75	0.03	0.01	1.97	0.02	0.01	0.00	0.74	0.01	1.17	0.07	0.00	0.00	800
212R1-17	1404.27	G	cpx	51.83	1.01	0.42	10.28	0.27	13.59	21.07	0.28	0.00	1.97	0.05	0.01	0.00	0.33	0.01	0.77	0.86	0.02	0.00	850
212R1-17	1404.27	G	opx	51.99	0.58	0.33	23.28	0.57	20.46	1.90	0.03	0.01	1.98	0.03	0.01	0.00	0.74	0.01	1.16	0.08	0.00	0.00	850
213R1-17	1406.27	G	cpx	51.71	1.23	0.50	10.19	0.39	14.00	20.77	0.36	0.01	1.96	0.06	0.01	0.00	0.32	0.01	0.79	0.84	0.03	0.00	861
213R1-17	1406.27	G	opx	52.50	0.53	0.32	21.08	0.66	21.90	1.80	0.05	0.01	1.98	0.02	0.01	0.00	0.66	0.02	1.23	0.07	0.00	0.00	861
227R1-17	1468.67	3	cpx	52.62	0.81	0.17	10.27	0.30	14.02	20.80	0.24	0.03	1.98	0.04	0.00	0.00	0.32	0.01	0.79	0.84	0.02	0.00	861
227R1-17	1468.67	3	opx	52.45	0.86	0.37	21.76	0.55	21.57	2.01	0.02	0.01	1.97	0.04	0.01	0.00	0.68	0.01	1.21	0.08	0.00	0.00	861
230R1-98	1483.98	1	cpx	52.29	1.16	0.42	10.20	0.28	14.34	20.82	0.25	0.01	1.96	0.05	0.01	0.00	0.32	0.01	0.80	0.84	0.02	0.00	899
230R1-98	1483.98	1	opx	52.47	1.07	0.37	22.10	0.50	21.56	2.04	0.02	0.00	1.96	0.05	0.01	0.00	0.69	0.01	1.20	0.08	0.00	0.00	899
230R1-98	1483.98	6	cpx	51.86	1.45	0.58	11.18	0.32	14.34	19.78	0.32	0.00	1.95	0.06	0.02	0.00	0.35	0.01	0.80	0.80	0.02	0.00	899
230R1-98	1483.98	6	opx	52.51	0.87	0.40	22.34	0.51	21.41	1.87	0.02	0.01	1.97	0.04	0.01	0.00	0.70	0.01	1.20	0.08	0.00	0.00	755
232R1-54	1493.44	5	cpx	53.29	0.37	0.06	7.44	0.24	14.91	22.59	0.23	0.00	1.99	0.02	0.00	0.00	0.23	0.01	0.83	0.91	0.02	0.00	755
232R1-54	1493.44	5	opx	52.90	0.95	0.46	19.52	0.43	23.40	2.01	0.03	0.01	1.96	0.04	0.01	0.00	0.61	0.01	1.29	0.08	0.00	0.00	755

<sup>a</sup>Each composition is an average of multiple analyses within the same area of an individual thin section. All analyses given in auxiliary material. Temperatures calculated using *Brey and Kohler* [1990] (see section 4.1).

positions; orthopyroxenes are hypersthene (Table 1 and Figure 5).

[32] Secondary clinopyroxene and orthopyroxene commonly contain blebs of secondary magnetite like those in amphibole replacing clinopyroxene in the overlying rocks. Clinopyroxene phenocrysts rarely have rims of secondary clinopyroxene with inclusions of magnetite, identical to the occurrence of amphibole + magnetite in the shallower dikes. These textures plus compositional data for phenocrysts and rims [Koepke *et al.*, 2008] indicate derivation of secondary pyroxenes from igneous clinopyroxene that was previously hydrothermally altered to amphibole + magnetite. The patchiness of the granoblastic recrystallization is consistent with prior hydrothermal alteration influencing the intensity of granoblastic recrystallization, which varies even within a single thin section [Teagle and Expedition 309/312 Scientists, 2006].

[33] Two types of vein are present in the granoblastic dikes. The first consists of ~1 mm wide felsic veins that contain subequal amounts of subhedral quartz and plagioclase, with minor coarse grained and fibrous interstitial amphiboles, plus trace FeTi oxides and needles of apatite. These veins are remarkably similar to tonalite veins that result from partial melting of hydrated dike rocks [Gillis and Coogan, 2002; Koepke *et al.*, 2008; France *et al.*, 2009]. In one case (at 1363.70 mbsf), actinolite occurs with oligoclase (An<sub>21-30</sub>) in the vein, and actinolite replaces clinopyroxene and orthopyroxene in the granoblastic wallrock. In another case (at 1367.52 mbsf), actinolite and minor Mg-hornblende are present with andesine (An<sub>48</sub>) in the vein, and Mg-hornblende replaces wallrock pyroxenes.

[34] The second type of vein consists of ~1 mm wide coarse Mg-hornblende + actinolite veins having sharp to diffuse contacts with the host rock, and containing bands and patches of granular material. Some of these veins contain a few percent quartz, apatite, and/or sulfides (pyrite and chalcopyrite) with the amphibole. One Mg-hornblende + actinolite vein (at 1382.19 mbsf) contains up to 400  $\mu\text{m}$  grains of andesine (An<sub>35</sub>), partly replaced by anorthite (An<sub>88</sub>) in small patches. A vein at 1386.97 mbsf contains irregular patches, up to ~1 mm, of granular Mg-hornblende + anorthite (An<sub>92-98</sub>) and local quartz, with anorthite partly replaced by albite. Included granoblastic material in the veins is variably replaced by actinolite and oligoclase. Pyroxenes in the wallrock surrounding these veins are variably

replaced by Mg-hornblende and actinolite, but Fe-Mg amphibole also replaces granular orthopyroxene (e.g., at 1397 mbsf; Figure 10). Orthopyroxenes in the dike screens are also locally partly replaced by smectite, talc, and chlorite-smectite.

[35] Mineral parageneses indicate a series of retrograde reactions, from contact metamorphism and formation of felsic veins, to Mg-hornblende and local anorthite in veins and host rocks, to actinolite and oligoclase in veins and rocks, to late albite replacing earlier more calcic plagioclase, and finally smectite and chlorite-smectite replacing orthopyroxene.

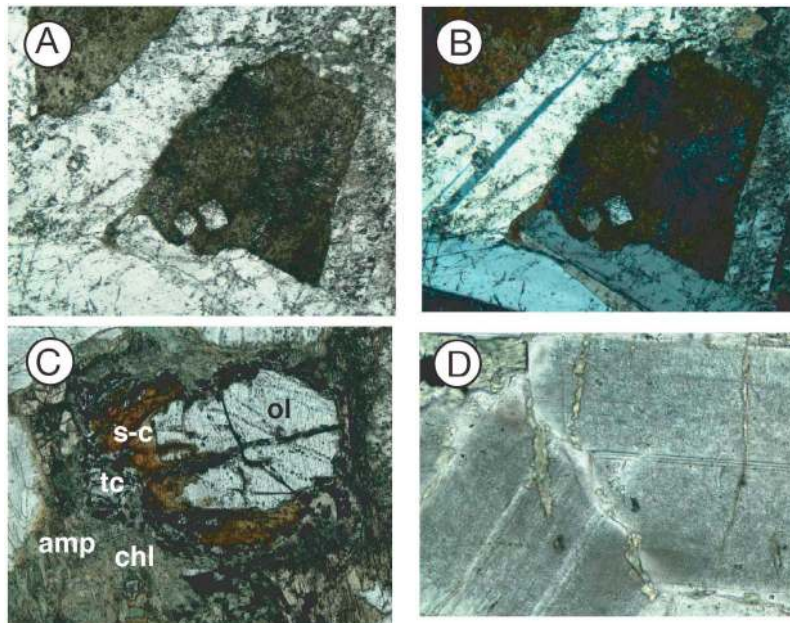
[36] The lowermost dike recovered in the hole exhibits alteration similar to that in the upper dike complex, with partial alteration to actinolite, albite, chlorite, and titanite.

### 3.1.4. Plutonic Rocks

[37] The plutonic rocks are variably altered, with the intensity of replacement related to texture of the rock and proximity to intrusive contacts and veins. Regions of gabbro with large ophitic clinopyroxenes enclosing plagioclase are less altered than irregular coarser grained zones between, giving the rocks a patchy appearance in hand specimen (Figure 7d).

[38] Clinopyroxene in the plutonic rocks is extensively to totally replaced by amphibole + dusty to granular magnetite (~1–40  $\mu\text{m}$ ) (Figure 11). Amphibole compositions typically range from Mg hornblende to actinolite and local ferroactinolite, but clinopyroxene can also contain small patches of brown pargasite, most likely of late magmatic origin [Banerjee and Gillis, 2001; Coogan *et al.*, 2001]. Fe-Mg amphibole is present locally, for example replacing orthopyroxene at 1490 mbsf and with actinolite replacing pyroxene at 1484 mbsf (Figure 10). Plagioclase is generally only slightly altered to albite-oligoclase (An<sub>1-30</sub>), and to small amounts of amphibole, chlorite, epidote, prehnite and laumontite (Figure 11). Olivine, where present, is variably altered to smectite-chlorite, chlorite, talc, calcite, magnetite, pargasite, Mg hornblende, and rare actinolite (Figure 11). These minerals are commonly zoned, from an inner core of fresh olivine with veins of magnetite, outward to smectite-chlorite or chlorite  $\pm$  calcite, to colorless Mg-hornblende + magnetite, to pale green pargasite.

[39] Oxide minerals consist of titanomagnetite and ilmenite, either as subgrains within individual grains or as oxidation/exsolution lamellae of ilmenite



**Figure 11.** Thin section photomicrographs of Gabbro alteration. (a) Clinopyroxene altered to hornblende. Plagioclase is partly altered to secondary plagioclase ( $An_{9-42}$ ; sample 312-1256D-223R-1, 42–45 cm; 1449.72 mbsf) (plane polarized light; field of view (FOV) = 4.5 mm). (b) Same as Figure 11a but with crossed polars. (c) Olivine partly replaced by smectite-chlorite (s-c) + magnetite + minor talc (tc), with an outer zone of chlorite (chl) and amphibole (amp) (sample 312-1256D-223R-1, 42–43 cm; 1449.72 mbsf) (plane-polarized light; field of view (FOV) = 2.25 mm). (d) Plagioclase riddled with tiny inclusions and cut by veinlets of amphibole and albite (sample 312-1256D-232R-2, 98–100 cm; 1494.96 mbsf) (plane-polarized light; FOV = 0.6 mm).

within magnetite, and are variably altered to titanite. Igneous apatite is present in gabbros and felsic veins, but apatite also occurs in the upper gabbro as up to 1 mm irregular porous crystals in altered plagioclase and with hornblende replacing pyroxene. The mode of occurrence of this apatite suggests a possible hydrothermal origin, but the high F contents and  $F/(F + Cl)$  (0.74–2.74 wt % and  $\sim 0.5$ , respectively) favor a magmatic origin.

[40] Hydrothermal effects are generally similar in the upper and lower gabbro bodies, but the lower gabbro contains orthopyroxene, which is variably replaced by Mg hornblende, talc, and chlorite-smectite. Granoblastic assemblages are present in stoned dike fragments (up to 10 cm) at the margins of the lower gabbro body. Granoblastic textures elsewhere within the lower gabbro reflect assimilation of stoned rocks [France *et al.*, 2009].

[41] The plutonic rocks contain an average of 10 veins/m of recovered core. The earliest generation consists of linear zones of amphibole replacing pyroxenes, with 1–2 mm wide amphibole-rich alteration halos. These are cut by 0.5–1 mm wide veins of amphibole filling open fractures, in some

cases also containing granular plagioclase ( $An_{17-40}$ ). Vein amphibole compositions range from Mg hornblende to pargasite and edenite. These early veins are cut by veins of actinolite, chlorite, quartz, epidote, and prehnite, bordered by 2–5 mm alteration halos containing actinolite, chlorite, albite, titanite, and local FeMg amphibole.

### 3.2. Fluid Inclusions

[42] Most fluid inclusions in quartz in the dikes and gabbros have salinities of 0.5–12.2 equivalent wt % NaCl, and homogenization temperatures of 213°C–446°C. Assuming 2.5–3.0 km water depth and up to 1 km seafloor sample depths gives hydrostatic pressures of  $\sim 350$ –400 bars. Applying these pressure corrections to the homogenization temperatures gives trapping temperatures of 269°C–469°C (Table 2). Average values of trapping temperatures for individual samples are 355°C–425°C, and average salinities are 2.2–9.2 wt % NaCl (Figure 12 and Table 2). Samples from the sheeted dikes tend to have lower average salinities than those from the granoblastic dikes and upper gabbro (<6 versus >7 wt % NaCl, respectively; Figure 12). Otherwise, there are no consistent differences

**Table 2.** Hole 1256D Fluid Inclusion Data Summary<sup>a</sup>

Core Section, cm Interval	Depth (mbsf)	Rock Type	Occ	n	Salinity Range	Salinity Mean	1s	Pressure (bars)	Tt Range	Tt Mean	1s
173R1, 107–110	1261.67	dike	Qz vein	19	1.3–10.1	5.7	2.5	350	269–422	386	34
176R1, 41–44	1276.51	dike	Qz vug	23	2.0–7.3	5.6	1.1	350	250–431	355	57
176R1, 107–109	1277.17	dike	Qz vein	15	0.5–6.8	2.2	1.9	350	352–456	425	25
194R1, 45–47	1358.35	Gb	tonalite vein	16 17	2.8–11.7 47.2 to >50	7.6	3.1	390	312–436 398 to >450	369 >420	34
212R1, 29–31	1404.39	felsic	ground vein mass	11 6	5.8–11.2 >50	9.2	1.8	400	275–469 >450	394	60
214R1, 94–95	1411.84	gabbro	ground mass	4 8	6.34–9.0 >50	7.6	1.1	400	346–383 >400	375	19
219R1, 61–62	1430.61	gabbro	Qz vein	4	3.7–12.2	7.0	3.9	410	332–427	372	43
222R1, 144–149	1446.04	gabbro	Qz vein	10 2	5.8–11.2 >50	7.0	1.4	410	275–469 >450	386	42

<sup>a</sup>All inclusions contained within quartz. Salinity in wt % NaCl equivalent; Tt is trapping temperature (corrected for assumed hydrostatic pressure, see section 3.2). N, number of analyses; 1s, 1 sigma. Inclusions fall into two groups, lower salinity and hypersaline. Mean values given for lower-salinity group, and minimum values given for hypersaline group (see section 3.2 for details). Data given in auxiliary material. Gb, granoblastic texture; Occ, occurrence.

between veins and vugs in the dikes, or between quartz veins and groundmass in the gabbros. Data for the tonalite vein are similar to those for quartz veins, whereas the plagiogranite vein has the highest average salinity (9.2 wt %; Table 2). Eight percent of all fluid inclusions were tentatively identified as secondary, but there is no consistent difference in homogenization temperature or salinity between primary and possible secondary inclusions.

[43] Some inclusions from felsic veins in the granoblastic dikes and from vein and groundmass quartz in the upper gabbro contain daughter halite crystals and did not homogenize at temperatures of 450°C, approximately the maximum usable temperature with the heating-freezing stage. Assuming a simple NaCl-H<sub>2</sub>O system, this implies a salinity of greater than about 50% for these inclusions, which are all identified as primary inclusions (Table 2).

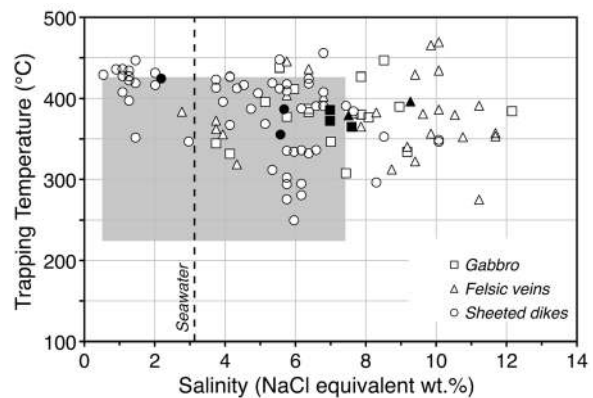
### 3.3. Oxygen Isotopes

[44] Oxygen isotope analyses of vein minerals are given in Table 3. In the volcanic section, the  $\delta^{18}\text{O}$  values of secondary minerals generally decrease with depth. Smectite and chlorite-smectite in the lavas have  $\delta^{18}\text{O}$  values of 6.8‰–17.8‰, and two celadonites have values of 13.2‰ and 17.2‰. Quartz and chalcedony in the volcanic section have  $\delta^{18}\text{O} = 17.1\text{‰}$ –30.2‰, whereas quartz from the transition zone, dikes, and plutonic section has lower  $\delta^{18}\text{O}$  values of 5.5‰–9.4‰. Quartz from the tonalite vein in the granoblastic dikes has  $\delta^{18}\text{O} = 7.0\text{‰}$ , within the range of vein quartz. Two epidote separates from alteration patches in a dike and a gabbro have  $\delta^{18}\text{O}$  values of 1.4‰ and 0.5‰, respectively. Fourteen anhydrites have  $\delta^{18}\text{O}$  values

in the range 9.0‰–17.7‰, whereas two others have more extreme values of 2.2‰ and 29.0‰.

## 4. Discussion

[45] The hydrothermal system and alteration processes at Site 1256 are discussed below in terms of depth, lithology, and time. We start with high-temperature reactions at the bottom of the section, documenting changes in the system through time,



**Figure 12.** Trapping temperatures and salinities for fluid inclusions in quartz (sample average values in solid symbols). Most inclusions have trapping temperatures and salinities similar to seafloor vent fluids (shaded box [Von Damm, 1995]) and are explained by supercritical phase separation processes (see section 4.4). Hypersaline inclusions (~50 wt % NaCl, see Table 2) did not homogenize at the maximum temperature measured (450°C) and plot off scale at top right. Dashed line is seawater salinity shown for comparison. Solid symbols are sample averages.

**Table 3.** Hole 1256D Oxygen Isotope Data, Equilibrium Temperatures, and Fluids<sup>a</sup>

Core Section	Centimeter Interval	Depth (mbsf)	Mineral	Occurrence	$\delta^{18}\text{O}$ (‰)	T <sup>b</sup> (°C)	Fluid Inclusion T (°C)	Fluid $\delta^{18}\text{O}$	T <sup>c</sup> (°C)
5R1	17–32	252.57	saponite	vein	16.2	60			
6R4	53–56	261.6	chalcedony	vein	30.2	49			
8R1	47–48	313.77	saponite	vein	16.7	56			
8R6	0–3	319.11	quartz	vein	24.0	81			
12R7	101–104	350.12	quartz	vein	28.1	58			
14R1	23–28	359.73	saponite	vein	17.8	48			
18R2	136–138	380.62	saponite	vein	17.3	52			
22R2	67–70	407.9	quartz	vein	27.7	60			
26R3	90–93	441.82	quartz	vein	25.0	75			
27R2	105–108	447.89	celadonite	vein	17.2	70			
35R2	108–111	491.33	quartz	vein	26.4	67			
35R2	108–111	491.33	saponite	vein	14.3	75			
43R2	13–16	535.53	quartz	vein	22.6	90			
47R2	141–144	573.91	saponite	vein	13.9	79			
53R2	13–21	611.4	quartz	vein	21.3	99			
55R2	118–122	628.58	chalcedony	vein	23.9	81			
57R2	70–74	647.55	quartz	vein	20.1	108			
57R2	70–74	647.55	chalcedony	vein	22.1	93			
57R2	75–76	647.6	saponite	vein	15.4	66			
60R1	35–38	668.95	saponite	vein	8.9	135			
68R1	112–113	720.62	quartz	vein	21.2	99			
70R2	8–12	730.24	celadonite	vein	13.2	100			
72R1	140–149	739.3	quartz	vein	24.3	79			
81R-1	58–60	789.4	anhydrite	haloclastite	14.5	136			
81R2	88	789.68	quartz	vein	17.1	133			
85R2	118–123	814	saponite	vein	11.9	98			
85R6	26	818.51	quartz	vein	25.6	72			
99R1	82	908.22	quartz	vein	22.1	93			
100R2	10–14	912.14	saponite	vein	9.6	125			
108R1	119	959.99	quartz	vein	22.7	89			
114R-1	113–116	989.93	anhydrite	vein w. zeolite	17.7	105			
117R1	12–14	1003.32	saponite	vein	5.8	185			
118R-1	11–13	1008.1	anhydrite	alt. patch	14.1	140			
122R-2	71–73	1029.51	anhydrite	breccia, glass, ab,py	14.9	131			
122R-2	73–78	1029.54	anhydrite	breccia, glass, ab,py	14.5	136			
123R-1	31–34	1032.41	anhydrite	vein	13.1				
128R-1	47–51	1056.57	anhydrite	hyalo br: sap, zeol, sulf	10.9	180			
130R-1	85–89	1066.55	anhydrite	anh + sap + qz vein	11.3	174			
135R-1	104–108	1090.74	anhydrite	vug w. qz + py	16.3	117			
137R1	4	1099.34	quartz	vein	7.8				356
140R-1	29–33	1113.99	anhydrite	dike margin br: chl + cp	2.2	408			
145R-2	83–89	1139.83	anhydrite	chl + py vein (late anh)	9.0	211			
151R-1	111–114	1166.41	anhydrite	chl + py vein	29.0	36			
151R1	111	1166.41	quartz	vein	9.0				318
156R-2	86–91	1191.69	anhydrite	chl + qz + py vn (late anh)	11.0	179			
157R-1	142–146	1195.62	anhydrite	vein	9.7	199			
170R-1	66–70	1252.46	anhydrite	chl + qz vein (late anh)	9.8	197			
173R1	88–94	1261.48	quartz	vein	7.1		386 <sup>d</sup>	1.6	382
174R1	91–98	1266.31	quartz	patch	6.8	354 <sup>c</sup>		0.4	
174R1	91–98	1266.31	epidote	patch	1.4				
191R1	6–11	1343.56	quartz	vein	5.5				454
194R1	9–14	1357.99	quartz	tonalite vein	7.0		369 <sup>f</sup>	1.0	386
202R1	5–15	1372.85	quartz	vein	7.8				358
219R1	61–65	1430.61	quartz	vein	9.4		372 <sup>g</sup>	3.5	308
222R1	67–71	1445.27	quartz	vein	6.9		386 <sup>h</sup>	1.4	390
225R1	8–9	1458.98	quartz	vein	5.9				434
227R1	92–96	1469.42	quartz	patch	9.0	226 <sup>e</sup>			
227R1	92–96	1469.42	epidote	patch	0.5			–2.8	



and work upward to lower-temperature processes in the lavas.

#### 4.1. Early Hydrothermal Alteration and Contact Metamorphism in the Granoblastic Dikes and Plutonic Section

[46] Petrographic and lithologic relationships show that the granoblastic dikes and the dike screens were initially hydrothermally altered, then underwent a stage of high-temperature contact metamorphism. The variability in high-temperature recrystallization, plus the presence of magnetite grains within secondary pyroxenes, identical to the occurrence of magnetite with amphibole replacing pyroxene in the upper dikes, are textural evidence for an early stage of hydrothermal alteration. Losses of Cu and Zn from the granoblastic dike are also consistent with an early period of hydrothermal alteration of these rocks (see section 4.5).

[47] The mineral assemblages of the granoblastic dikes and dike screen provide a record of the contact metamorphism. Applying the two-pyroxene thermometer of *Brey and Kohler* [1990] to pyroxene data for 8 samples of granoblastic dikes gives temperatures of 800°C–924°C, with no clear depth trend (Table 1). Temperature estimates from different areas within individual thin sections generally vary by less than 50°C, but in one sample at 1404 mbsf calculated temperatures vary by 100°C (Table 1). *Koepke et al.* [2008] obtained higher temperatures of 935°C–1045°C for the granoblastic dikes using the QUILF two-pyroxene thermometer of *Andersen et al.* [1993]. Comparison of temperatures of various published experiments with those calculated using two-pyroxene thermometers, however, shows that the *Brey and Kohler* [1990] thermometer gives smaller average deviations from the experimental temperatures than does the QUILF thermometer, 70°C versus 113°C, respectively [*Putirka*, 2008; *Blundy and Cashman*, 2008]. Recalculation of the pyroxene data from *Koepke et al.* [2008] using the *Brey and Kohler* thermometer for comparison gives

temperatures of ~850°C–950°C, similar to results in Table 1.

[48] Data for the dike screen between the two gabbro bodies and for the lower gabbro unit indicate that these rocks experienced peak metamorphic temperatures similar to those in the granoblastic dikes. Two-pyroxene thermometry yields a temperature estimate of 861°C for one sample from the upper dike screen, and 755°C–900°C for two samples of the lower gabbro (Table 1).

[49] Temperatures of contact metamorphism are consistent with those for hydrous partial melting [*Gillis and Coogan*, 2002; *Koepke et al.*, 2008], and felsic veins in the granoblastic dikes are suggested to result from partial melting of previously hydrated sheeted dikes [*France et al.*, 2009]. The tonalite veins in the granoblastic dikes are probably the result of this process. The presence of local andesine and apatite in the amphibole-rich veins in the granoblastic dikes suggests that these veins may also have a magmatic component.

[50] The two gabbro sills penetrated by the drill hole have insufficient thermal mass to produce a 60 to 90 m contact metamorphic halo [*Koepke et al.*, 2008; *Coggan et al.*, 2008], suggesting a larger intrusion below. Alternatively, there may be significant lateral relief on the sheeted dike – gabbro boundary at Site 1256. The most strongly recrystallized diabases are not at the boundary with the underlying gabbro sills, suggesting that other gabbroic bodies may be present at slightly shallower levels nearby [*Teagle and Expedition 309/312 Scientists*, 2006].

#### 4.2. Post-Contact Metamorphism Hydrothermal Alteration of the Granoblastic Dikes and Plutonic Section

[51] The earliest hydrous alteration phases in the granoblastic dikes and plutonic section are amphiboles. We did not use plagioclase-amphibole thermometry for our samples because adjacent

##### Notes to Table 3:

<sup>a</sup>Abbreviations are as follows: anh, anhydrite; sap, saponite; qz, quartz; py, pyrite; br, breccia. See section 4.4 for details and discussion.

<sup>b</sup>Temperatures in the volcanic section (<1004 mbsf) assume equilibrium with seawater (0‰) for saponite [*Buatier et al.*, 1995], celadonite [*Savin and Lee*, 1988] and quartz (<200°C [*Zheng*, 1993]).

<sup>c</sup>Temperatures for quartz in the transition zone, dikes, and gabbros (below 1004 mbsf, T > 200°C) based on fluid inclusions and assuming hydrothermal fluid d18O = 1.5‰ using quartz-water fractionation from *Sharp and Kirschner* [1994]. Temperatures for anhydrite assume equilibrium between sulfate and seawater [*Chiba et al.*, 1981].

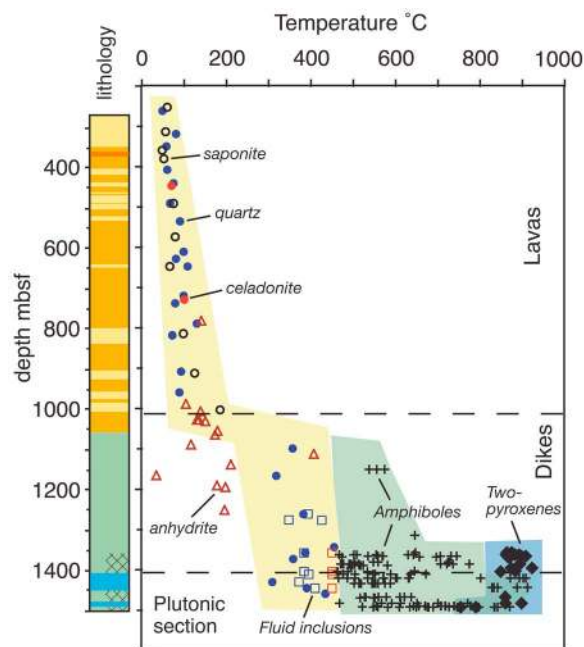
<sup>d</sup>From 173R1, 107–110.

<sup>e</sup>Using quartz-epidote fractionation from [*Matthews and Schliestedt*, 1984], and quartz-water fractionation from *Sharp and Kirschner* [1994].

<sup>f</sup>From 194R1, 45–47.

<sup>g</sup>From 219R1, 61–62.

<sup>h</sup>From 222R1, 144–149.



**Figure 13.** Temperatures estimated from various mineral thermometers in Hole 1256D. Note step in thermal gradient from low temperatures in the lavas to hydrothermal alteration in the sheeted dikes. Amphibole and two-pyroxene thermometry (solid diamonds) record stepwise increase to near-magmatic temperatures in the granoblastic dikes and plutonic section. Anhydrites (open triangles) record either nonequilibrium temperatures of precipitation or precipitation during collapse and cooling of the system. Temperatures estimated from oxygen isotopes (saponite, open circles; celadonite, solid red circles; quartz, solid blue circles; anhydrite, open triangles), fluid inclusions in quartz (open squares; hypersaline inclusions plotted in red at minimum temperature of 450°C), Ti contents of amphiboles (crosses), and two-pyroxene thermometry (solid diamonds). Lithology as in Figure 2.

plagioclase and amphibole were not always analyzed, and because textural evidence for equilibrium between these phases is commonly not clear. Compositions of plagioclase in the granoblastic dikes may reflect the contact metamorphism stage, with amphibole forming later during hydration of the granoblastic assemblages. *Koepke et al.* [2008] used the plagioclase-amphibole thermometer of *Holland and Blundy* [1994] for what they termed later “wet” granoblastic metamorphism of these rocks, and obtained temperatures of 708°C–854°C.

[52] *Ernst and Liu* [1998] show that  $\text{TiO}_2$  and  $\text{Al}_2\text{O}_3$  contents of calcic amphibole formed from mid-ocean ridge basalt protoliths increase with temper-

ature, providing a semiquantitative thermometer. Application of this semiquantitative amphibole thermometer to Hole 1256D analyses gives temperature estimates of 464°C–913°C (Figure 13). Equilibrium was at best attained locally in these samples during amphibole formation, and uncertainties in temperatures at low  $\text{TiO}_2$  contents (<0.5 wt %) from *Ernst and Liu’s* [1998] correlations are high, but uncertainties of  $\pm 100^\circ\text{C}$  for these estimates are probably reasonable. The highest temperatures, 829°C–913°C, are for pargasite-edenite in the upper gabbro unit, where these amphiboles occur in olivine reaction coronas, replacing clinopyroxene, and in a vein with Mg-hornblende and actinolite. Some of these high-temperature amphiboles are most likely late magmatic in origin. Temperatures estimated for other amphiboles in the gabbro bodies vary from 840°C down to 480°C.

[53] The highest temperatures estimated for amphibole in the granoblastic dikes (700°C–776°C) are for coarse amphibole veins and for wallrock associated with these veins (at 1387.03, 1382.19, and 1367.52 mbsf). Other amphibole temperatures for these rocks range from 650°C to 460°C. Using the same semiquantitative thermometer, *Koepke et al.* [2008] obtained temperatures of 650°C–747°C for amphibole formation in the granoblastic dikes. Fluid inclusion and oxygen isotope data, discussed below, along with the mineralogy indicate that hydrothermal alteration continued throughout the cooling of the granoblastic dikes and plutonic section. Actinolitic amphiboles may have continued to form at temperatures down to  $\sim 300^\circ\text{C}$  [*Bird et al.*, 1984].

### 4.3. Early Alteration of the Sheeted Dikes and Transition Zone

[54] Petrographic relationships indicate that clinopyroxene, amphibole (Mg-hornblende to actinolite), chlorite, albite-oligoclase, titanite,  $\pm$ pyrite formed relatively early in veins and bulk rocks throughout the transition zone and sheeted dikes. Application of the semiquantitative amphibole Ti thermometer [*Ernst and Liu*, 1998] to amphiboles in dikes gives temperature estimates as high as 540°C–650°C (Figure 13). Local secondary clinopyroxenes in veins formed at comparable temperatures of  $\sim 400^\circ\text{C}$ –800°C, based on the one-atmosphere solvus of *Lindsley* [1983] (Figure 5). Based on comparison with active geothermal areas, however, actinolite, chlorite and the other phases probably continued to form at lower temperatures, down to  $\sim 300^\circ\text{C}$  [*Bird et al.*, 1984].

#### 4.4. Formation of Quartz and Associated Minerals in the Sheeted Dikes and Plutonic Section

[55] Veins of quartz with other phases (chlorite, epidote, pyrite, chalcopyrite, magnetite, and local sphalerite) are common and overprint early hydrothermal alteration. Fluid inclusion and oxygen isotope data for quartz and epidote provide constraints on the quartz-forming hydrothermal fluids.

[56] Fluid inclusions in quartz veins and vugs are mainly relatively low salinity, having individual sample averages of 2.2–7.0 wt % NaCl (Table 2). These inclusions have salinities and trapping temperatures similar to those of hydrothermal vent fluids at mid-ocean ridges (Figure 12) [Butterfield *et al.*, 1994; Von Damm, 1995], but range to higher salinities. The variable salinities of the vent fluids result from processes related to supercritical phase separation. The critical point of seawater is 407°C and 298 bars [Bischoff and Rosenbauer, 1985]. At higher pressure and temperature, decompression or heating can lead to supercritical phase separation and formation of a lower-salinity vapor-like phase plus a high-salinity brine phase [Bischoff and Rosenbauer, 1985]. Separation and recombination of these phases in variable proportions in the subsurface results in the observed salinity variations of vent fluids [Von Damm, 1995; Edmonds and Edmond, 1995].

[57] The depth to the seafloor at the EPR where Site 1256 formed is ~2.5 km, and the lower dikes and plutonic section lie ~1 km subbasement, which would have been at pressures of ~360 bars at the spreading axis. This is above the critical point pressure for seawater, and the trapping temperatures of the inclusions in quartz veins and vugs range up to 469°C, spanning across the supercritical two-phase boundary. At 350–360 bars and temperatures of 425°C–440°C, supercritical phase separation leads to formation of brines having salinities of 7–10 wt % NaCl equivalent plus a vapor phase having salinity of ~1 wt % NaCl, as observed in other oceanic sheeted dikes and plutonic rocks [Schöps and Herzig, 1990; Kelley *et al.*, 1993; Saccocia and Gillis, 1995; Heft *et al.*, 2008]. If some cooling of fluids is taken into account, these temperatures and salinities are comparable to those for the inclusions in quartz veins and vugs from Hole 1256D.

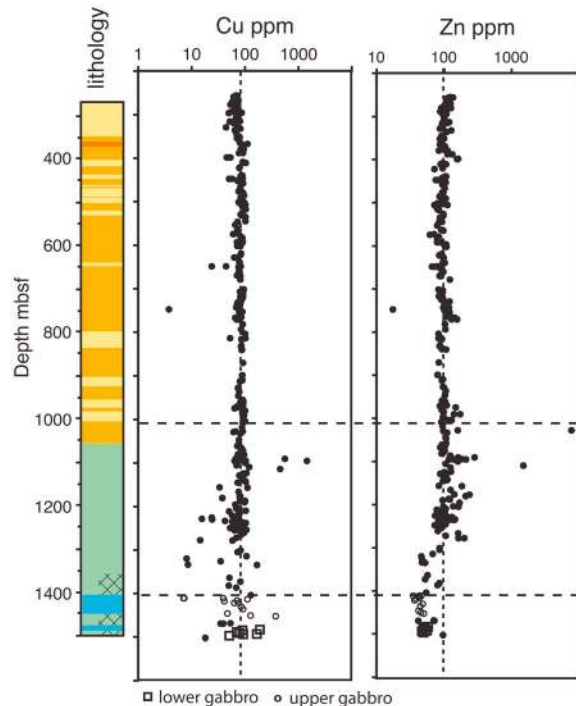
[58] Quartz in the felsic veins and gabbro groundmass contain lower-salinity inclusions similar to those in the quartz veins, but also contain hyper-

saline inclusions that range to ~50 wt % NaCl (Table 2). Given their occurrence in magmatic veins and groundmass, these hypersaline inclusions probably record magmatic fluids. One hydrothermal quartz vein in gabbro (at 1446.04 mbsf) also contains a component of high-salinity inclusions (Table 2). Such brines can form through supercritical phase separation of seawater at 350–400 bars and temperatures of 450°C–500°C [Kelley *et al.*, 1993; Saccocia and Gillis, 1995], but could also reflect input of magmatic fluids into circulating hydrothermal fluids within the plutonic section.

[59] The quartz-forming fluids can be further constrained with oxygen isotope analyses. Using average trapping temperatures of fluid inclusions in quartz veins and  $\delta^{18}\text{O}$  values of quartz adjacent to the fluid inclusion samples, application of the quartz-water oxygen isotope fractionation of Sharp and Kirschner [1994] shows that fluids in equilibrium with quartz veins had  $\delta^{18}\text{O}$  values of 1.4‰–3.5‰ (Table 3). Assuming that quartz from a tonalite vein (at 1357.99 mbsf) was in equilibrium with an aqueous fluid and using temperature of the lower-salinity inclusions in an adjacent quartz sample, the fluid would have had a  $\delta^{18}\text{O}$  value of 1.0‰ (Tables 2 and 3).

[60] Temperatures and fluid compositions can also be estimated using quartz-epidote mineral pairs. Applying the quartz-epidote temperature fractionation of Matthews and Schliestedt [1984] and using a pistacite content of 0.29, one quartz-epidote pair from an alteration patch in the dikes (at 1256 mbsf) gives a temperature of 354°C (Table 3). The fluid in equilibrium with this quartz would have had  $\delta^{18}\text{O} = 0.4\text{‰}$  [Sharp and Kirschner, 1994]. A second quartz-epidote pair from a quartz-epidote band in a gabbro at 1469 mbsf yields an equilibrium temperature of 228°C and a fluid  $\delta^{18}\text{O}$  of  $-2.8\text{‰}$  (Table 3). This temperature estimate is low for epidote formation, and the negative fluid  $\delta^{18}\text{O}$  value is unreasonable in rocks dominated by hydrothermal alteration by seawater at temperatures  $>250\text{°C}$ , where fluid-rock interactions lead to  $^{18}\text{O}$  enrichment of seawater fluids. Quartz in this sample has highly variable grain size suggesting multiple generations of quartz, and that this mineral pair does not represent equilibrium.

[61] The quartz-forming fluids in the dikes and plutonic section had  $\delta^{18}\text{O}$  values of 0.4‰–3.5‰, similar to seafloor hydrothermal vent fluids ( $\sim 0.5\text{‰}$ – $2\text{‰}$ ) [Shanks, 2001]. Assuming an intermediate  $\delta^{18}\text{O}$  value of 1.5‰ for hydrothermal fluids, other quartz samples in the sheeted dikes and plutonic section



**Figure 14.** Bulk rock metal contents versus depth. Note metal enrichments in upper dikes and metal depletions in lower dikes and plutonic section. Additional metal enrichments of upper dikes are indicated by local Fe, Cu, and Zn sulfides in veins. Dashed vertical lines for reference. There is little variation in metal contents with fractionation in lavas and dikes, so plots of changes in metal contents look the same. See section 4.5 for discussion (D. A. H. Teagle, unpublished data, 2010). Lithology as in Figure 2.

formed in equilibrium with such fluids at temperatures of 320°C–450°C (Figure 13 and Table 3).

#### 4.5. Mineralization and Metal Mobility in the Sheeted Dikes and Plutonic Section

[62] Local sulfide mineralization and metal enrichments occur in the upper dikes, as indicated by elevated Zn and Cu contents at ~1025–1110 mbsf (Figure 14). Veins of quartz + pyrite ± chalcopyrite ± sphalerite are also common down to ~1200 mbsf, documenting minor local metal enrichments not shown by the bulk rock analyses in Figure 14. Data for fluid inclusions and oxygen isotopes presented above show that hydrothermal fluids had temperatures, salinities, and oxygen isotope compositions similar to those of seafloor vent fluids. The association of metal sulfides with quartz in veins indicates that these subsurface fluids were also enriched in metals and sulfide like the vent fluids. The common occurrence of sulfide mineraliza-

tion and anhydrite in brecciated dike margins suggests that this hydrothermal activity was coeval with dike intrusion at the spreading axis [Teagle and Expedition 309/312 Scientists, 2006].

[63] The mineralized breccia near the top of the lavas-dikes transition coincides with the downward change from low-temperature seawater alteration to higher-temperature hydrothermal alteration. The intense sulfide mineralization, stepwise increase in alteration grade and temperatures, plus a peak in whole-rock Sr isotope ratios in this zone [Harris et al., 2008], indicate that alteration and mineralization of the breccia occurred in a mixing zone between generally upwelling hydrothermal fluids in the dikes and cooler seawater solutions in the overlying volcanic section.

[64] The dikes exhibit variable depletions in Cu below ~1150 mbsf and have low Zn contents below 1300 mbsf (Figure 14). These metal depletions reflect leaching of metals during hydrothermal alteration at high temperatures (>350°C), where the solubility of metals in hydrothermal fluids increases significantly [Seewald and Seyfried, 1990]. Metal depletions roughly coincide with a change in silicate alteration at ~1300 mbsf: below this depth chlorite diminishes and amphibole increases in abundance, and the dikes are more intensely recrystallized. The granoblastic dikes (below 1360 mbsf) and the dike screen between the two gabbro units have low Cu and Zn contents, but because these rocks exhibit only slight hydration to amphibole, the metal losses must have occurred during the initial stage of hydrothermal alteration, prior to contact metamorphism.

[65] The variable Cu and Zn contents of the plutonic rocks in part reflect igneous processes, but also result from hydrothermal leaching of metals. Moderately altered gabbros having high Cu contents (>150 ppm) contain mainly recrystallized igneous sulfides, reflecting igneous accumulation of CuFe and Fe sulfides. An oxide diorite at 1411.16 mbsf contains only 7 ppm Cu, in part reflecting the leaching of metals from the highly altered rock.

[66] The metal losses in the lower dikes and plutonic section are consistent with observations of other oceanic and ophiolitic sections [Alt et al., 1996; Alt and Anderson, 1991; Harper et al., 1988; Nehlig et al., 1994; Gillis, 1995; Gillis et al., 2001; Heft et al., 2008], and with this portion of the crust being the deep “root zone” of submarine hydrothermal systems. Inclusion-rich anorthite rims around magmatic plagioclase that occur in the hydrothermal root zone in the lower dikes of Hole 504B and in Oman [Vanko and Laverne, 1998] have not been

observed in Hole 1256D, perhaps because equivalent rocks have been recrystallized to granoblastic assemblages.

#### 4.6. Anhydrite Formation in Lavas and Dikes

[67] The lower lavas and upper dikes at Site 1256 are the sites of anhydrite precipitation (Figure 2). Using the sulfate-water oxygen isotope fractionation of *Chiba et al.* [1981] and assuming equilibrium with seawater, anhydrite formed at temperatures of 36°C–408°C, with most samples falling in the range 105°C–210°C (Figure 13 and Table 3). Anhydrites shallower than 1100 mbsf in the upper dikes are significantly enriched in <sup>18</sup>O relative to seawater sulfate (10.9‰–16.3‰ versus 9.3‰, respectively) [*Lloyd*, 1968], and have equilibrium temperatures of 105°C–180°C. Formation of anhydrite at these temperatures is consistent with heating of sulfate-bearing seawater fluids that may also have been enriched in Ca through reaction with basalt [*Shanks et al.*, 1981]. The kinetics of sulfur and oxygen exchange in sulfate are slow, however, at these temperatures and at the pH expected during basalt seawater reactions (~6 at 150°C) [*Seyfried and Bischoff*, 1979; *Chiba et al.*, 1981]. The measured sulfate  $\delta^{18}\text{O}$  values thus may represent only partial reaction toward higher equilibrium  $\delta^{18}\text{O}$  values at temperatures less than those calculated assuming equilibrium.

[68] Several anhydrites below 1100 mbsf in the upper dikes are generally only slightly enriched in <sup>18</sup>O compared to seawater sulfate, yielding equilibrium temperatures of 179°C–211°C (Table 3 and Figure 13). These slight <sup>18</sup>O enrichments could result from higher reaction temperatures than the shallower anhydrites, but because of the slow reaction rates at these conditions, the data may reflect kinetic effects. Seawater sulfate that reacted with basalt at temperatures similar to the shallower anhydrites, but to a lesser extent, would show the slight <sup>18</sup>O enrichments like the deeper anhydrites. The high- $\delta^{18}\text{O}$  anhydrite in a chlorite + pyrite + anhydrite vein at 1166 m is in equilibrium with seawater at 36°C: this sample must have formed from seawater sulfate that was highly reacted (near equilibrium) at low temperatures and which was then heated and mixed with hydrothermal fluids to form anhydrite. In contrast, anhydrite with chalcopyrite in a brecciated dike margin at 1114 mbsf has very low  $\delta^{18}\text{O}$  (2.2‰) reflecting high-temperature reactions (408°C if in equilibrium with seawater) [*Chiba et al.*, 1981].

[69] The anhydrites in Hole 1256D thus formed over a range of temperatures and with varying reaction histories, but are generally consistent with heating of seawater fluids and mixing with hydrothermal fluids as seawater penetrated downward into hotter dikes as also observed in Hole 504B [*Teagle et al.*, 1998]. Prior to anhydrite formation, the seawater-derived fluids reacted with basalt to varying degrees at temperatures <200°C. Anhydrite could reflect rapid precipitation at high temperatures during heating and mixing, freezing in the oxygen isotope exchange effects that occurred at lower temperatures. Alternatively, the temperatures could reflect equilibrium during anhydrite formation as seawater penetrated to greater depths as the system waned and cooled (Figure 13).

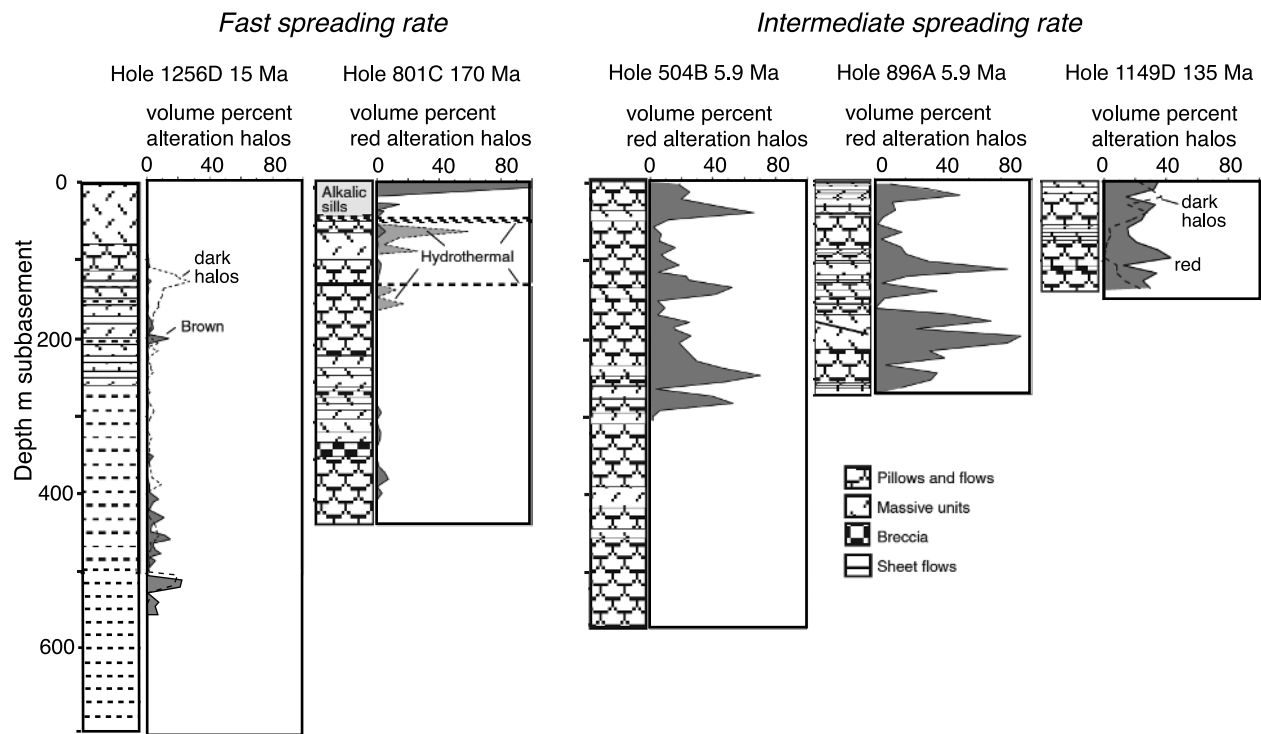
#### 4.7. Late Alteration Effects in Dikes and Plutonic Section

[70] Generally late alteration phases in veins and rocks include anhydrite, prehnite, laumontite, and calcite. With the exception of anhydrite, formation of these phases reflects overall decreasing temperatures and more evolved hydrothermal fluids (e.g., Mg-depleted, elevated Ca and pH) [*Seyfried et al.*, 1999]. Prehnite is stable relative to epidote at higher Ca activity and pH and at lower temperatures [*Rose and Bird*, 1987], and laumontite forms in geothermal systems at temperatures below about 240°C [*Neuhoff and Bird*, 2001]. Minor chlorite-smectite and smectite occur locally in the dikes and plutonic section, and reflect lower alteration temperatures (<200°C) and/or incomplete reaction of olivine and orthopyroxene to chlorite [*Alt et al.*, 1986].

#### 4.8. Low-Temperature Alteration of the Volcanic Section

[71] The mineralogy and hydrothermal processes in the volcanic section are generally typical of low-temperature alteration of upper oceanic basement (summary by *Alt* [2004]). Temperatures can be estimated from oxygen isotope data for secondary vein minerals. Assuming equilibrium with seawater (0‰), saponite, celadonite, quartz, and chalcedony formed at temperatures of 50°C–125°C, generally increasing downward, and ranging up to higher temperatures (185°C) near the base of the lavas (Table 3 and Figure 13).

[72] Petrographic relationships show that the earliest alteration was the formation of dark celadonite-bearing alteration halos along fractures. These



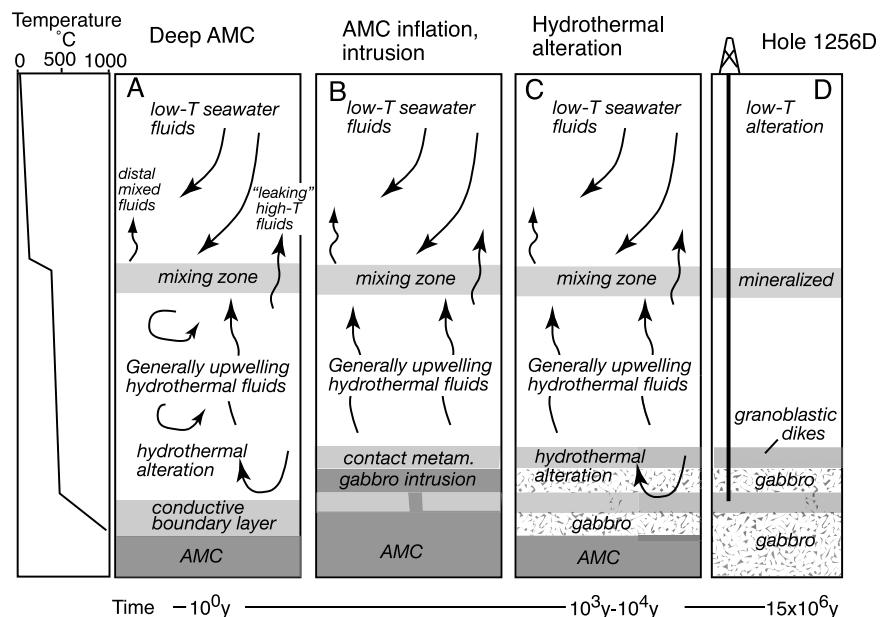
**Figure 15.** Distribution of alteration halos in upper oceanic basement. Alteration halos in crust formed at fast spreading rates are less abundant than those in basement formed at intermediate spreading rates, reflecting more rapid sealing of basement by sediment at fast spreading rates. See section 4.8 for discussion. [After *Alt et al.*, 1996; *Alt*, 2004.]

alteration halos form in very young basalts ( $<10^5$  yrs old) and result in additions of iron, silica, and alkalis to the rock [*Alt*, 2004]. Celadonite in these halos forms from upwelling distal hydrothermal fluids. Mixing of upwelling black-smoker-type hydrothermal fluids in the subsurface causes deposition of metal sulfides at depth, and the resulting cool ( $\sim 10^\circ\text{C}$ – $100^\circ\text{C}$ ) mixed fluids are enriched in iron, silica, and alkalis compared to seawater and to ridge flank basement fluids [*Edmond et al.*, 1979a, 1979b; *Wheat and Mottl*, 2000]. Such mixed fluids are manifest at the ridge crest as the diffuse venting that surrounds black smokers and that occurs sporadically between hydrothermal vent sites. The heterogeneity of diffuse flow in the subsurface contributes to variability in the thickness and distribution of the dark celadonitic alteration halos.

[73] Petrographic observations show that, subsequently, iron oxyhydroxides formed in unaltered rocks or superimposed on celadonitic alteration halos along fractures. It is inferred that this results from flow of cold seawater through the rocks [e.g., *Alt*, 2004]. Saponite, rich in ferrous iron, and pyrite may have formed simultaneously with iron oxyhydroxides, but in the adjacent host rock at lower seawater/rock ratios and under more reduc-

ing conditions. Circulation throughout the volcanic section evolved to lower seawater/rock ratios and reducing conditions as indicated by the ubiquitous late saponite  $\pm$  pyrite rocks and veins. Small amounts of late carbonates and zeolites formed from more highly evolved (Mg-depleted, elevated pH) low-temperature seawater-derived fluids [*Alt*, 2004].

[74] Compared to crust formed at intermediate spreading rates, the abundance of red iron-oxyhydroxide-rich alteration halos is much lower at Site 1256 and in the one other deep drillhole in fast spread crust (Hole 801C, Figure 15). This likely reflects sealing of basement to seawater access by sediment blanketing the smoother basement topography at fast spreading rates [*Alt and Teagle*, 2003]. Volcanic eruptive style may also play a role, with more abundant pillow basalts at intermediate spreading rates perhaps providing greater permeability and enabling more seawater circulation than do sheet and massive flows that dominate at fast spreading rates. The massive ponded lava flow capping the section at Site 1256 may have restricted flow in the immediately underlying rocks, but upper basement flow is generally horizontal over the scale of tens of kilometers, controlled by permeable horizons at



**Figure 16.** Summary of alteration processes at Site 1256. (a) Conductive boundary layer separates deep axial magma chamber (AMC) from high-temperature hydrothermal alteration and leaching of metals in the lower dikes. Generally upwelling hydrothermal fluids in the dikes mix in the transition zone with cooler seawater circulating in the overlying lavas. Distal mixed fluids percolate upward to alter basalt lavas locally and vent as diffuse flow on the seafloor. Some high-T fluids leak upward to locally alter lavas. Idealized temperature profile at left shows steps in thermal gradient across hydrothermal mixing zone and conductive boundary layer. (b) Inflation of AMC, intrusion of gabbro units into lower dikes, and contact metamorphism producing granoblastic dikes. (c) Crystallization and cooling of gabbro intrusions and upper AMC, with downward penetration of hydrothermal fluids and hydration of granoblastic dikes and plutonic section. Gabbro crystallized from upper AMC now acts as conductive boundary layer. (d) Present alteration stratigraphy at Site 1256. Time line based on  $200 \text{ mm yr}^{-1}$  full spreading rate and 1–2 km width of AMC.

shallow (100 s of m) depth [Fisher, 2005], so it is unlikely that the massive flow affected alteration of rocks hundreds of meters below.

[75] Oxygen isotope data indicate that temperatures generally increased downward in the volcanic section, with a stepwise increase to hydrothermal temperatures across the top portion of the lavas-dikes transition (Figure 13). The presence of local chlorite and chlorite-smectite in the lower lavas suggests locally higher temperatures ( $>200^\circ\text{C}$ ), probably related to flow of higher-temperature hydrothermal fluids “leaking” upward across the lithologic transition from the underlying dikes. The narrow chlorite-smectite-bearing green-gray halos and pyrite-bearing halos at the base of the lavas may also be related to “upward leaking” higher-temperature fluids.

[76] The intense alteration of the 41 cm interval at 648 mbsf (90% altered to celadonite, beidellite, K-feldspar, iron oxyhydroxide, quartz, and calcite) reflects focused fluid flow. The enrichments of K, Fe and Si in these rocks [Teagle and Expedition 309/312 Scientists, 2006] are inconsistent with seawater but are consistent with flow of cooler,

mixed hydrothermal fluids derived from the mineralized mixing zone below. The mineralogy of these intensely altered rocks is identical to that of brecciated pillow lavas beneath a low-temperature ( $\sim 60^\circ\text{C}$ ) seafloor iron-silica deposit in ODP Hole 801C, where the link between this intense alteration and distal hydrothermal fluids is clear [Alt and Teagle, 2003].

#### 4.9. Summary: Structure and Evolution of the Hydrothermal System at Site 1256

[77] In this section we combine the observations and interpretations for the different basement sections into an integrated view of the hydrothermal system at Site 1256, documenting interactions between hydrothermal circulation, basement lithology, and magmatism (Figure 16).

[78] The earliest reactions in the cored section involved hydrothermal alteration of the dike section above a melt lens at the EPR. Hydrothermal fluid circulation in the dikes is separated from the melt lens by a conductive boundary layer, where heat is transferred by conduction from magma to the

overlying convecting hydrothermal system [Gillis, 2008]. Hydrothermal alteration of the lower dikes at Site 1256 occurred at relatively high temperatures (>350°C), mobilizing metals from these rocks. Intrusion of the two gabbro bodies into the lower sheeted dikes and contact metamorphism of these rocks at temperatures of ~800°C–900°C may be related to inflation of the axial melt lens. This produced granoblastic textures in the lower dikes, the dike screen, and in the lower gabbro unit, with the previous hydrothermal alteration controlling the high-temperature recrystallization effects. Hydrous partial melting of altered dikes also occurred [Koepke *et al.*, 2008; France *et al.*, 2009]. This contact metamorphism represents the upward migration of the conductive boundary layer into hydrothermally altered rocks [Teagle and Expedition 309/312 Scientists, 2006; Koepke *et al.*, 2008]. Hydrothermal circulation and alteration later penetrated downward into the contact metamorphosed lower dikes and plutonic section as these rocks cooled, with initial hydrothermal fluid penetration into the granoblastic dikes at temperatures of 700°C–800°C. The inferred highest temperatures of fluid reactions in the gabbros as recorded by amphibole formation were 830°C–910°C, reflecting late magmatic fluids or the earliest penetration of seawater fluids [Coogan *et al.*, 2001; Koepke *et al.*, 2008].

[79] Contact hornfels metamorphism of basal dikes, hydrous partial melting, highly variable gabbroic rocks, and complex intrusive relationships have been documented at this lithostratigraphic level elsewhere in ocean crust and ophiolites [Gillis and Roberts, 1999; Gillis, 2008; Nicolas *et al.*, 2008; France *et al.*, 2009]. This “root zone” of the sheeted dikes is ~100 m thick, with contact metamorphic aureoles ~40 m thick that migrate vertically as much as 200–400 m as magmatism waxes and wanes. Temperatures of contact metamorphism (756°C–1024°C) and of initial hydration (701°C–997°C) in these rocks [Gillis, 2008] are similar to those estimated for Site 1256.

[80] Initial hydrothermal alteration of the shallower dikes involved formation of amphibole + magnetite, albite-oligoclase, chlorite, and titanite. Hydrothermal fluids in this section rapidly evolved to upwelling fluids similar to seafloor vent fluids, leading to mineralization at the lavas-dikes boundary where hydrothermal fluids mixed with cooler seawater fluids circulating in the overlying lavas. Quartz + sulfide ± anhydrite veins also formed locally in the upper dikes, with anhydrite recording prior evolution of seawater fluids at temperatures generally less than 200°C. Up to ~500 m of lavas

accumulated while Site 1256 was at the spreading axis [Tominaga *et al.*, 2009]. Distal mixed hydrothermal fluids percolating upward from the mixing zone in the lavas-dikes transition caused formation of the dark celadonite-bearing alteration halos along flow pathways in the volcanic section and may have fed diffuse venting at the seafloor. Local focused flow of these distal hydrothermal fluids and mixing with seawater led to localized intense alteration of lavas to celadonite + K-feldspar + beidellite + iron oxyhydroxide assemblages, which are now deep in the volcanic section (at 648 mbsf), but which were probably altered at or near the seafloor at the spreading axis.

[81] Hydrothermal activity in the dike complex was likely coeval with dike intrusion, as indicated by hydrothermal alteration and mineralization of brecciated dike margins, but continued as individual dikes and their surrounding host rocks cooled to greenschist facies conditions. Fluids evolved from early Mg-bearing hydrothermal fluids forming chlorite and amphibole in veins and host rocks at temperatures as high as ~600°C to “black-smoker-like” hydrothermal fluids at temperatures of 320°C–450°C. These mineralizing fluids had variable salinities resulting from processes related to supercritical phase separation in the lower dikes and plutonic section. Anhydrite formed in fractures in the lower lavas and upper dikes as partly reacted seawater penetrated downward and was heated and mixed with hot hydrothermal fluids.

[82] Prehnite and laumontite formed in the dikes and plutonic section from more evolved hydrothermal fluids as the hydrothermal system cooled. With further cooling, minor late chlorite-smectite and smectite formed locally in the dikes and plutonic section through hydration of relict olivine and orthopyroxene. The current temperature at the base of the cored section is ~100°C, and these phases could be forming at the present time.

[83] Fluids in the volcanic section evolved to more seawater dominated, with formation of iron oxyhydroxides in the rocks via flow along the same pathways as previous alteration. There is much less oxidation in Site 1256 lavas compared to crust formed at intermediate spreading rates, and where present it is spatially related to zones of concentrated subvertical fracturing. This lack of oxidative low-temperature alteration mostly likely reflects the efficient sealing of the relatively smooth basement by sediment. Only minor amounts of late carbonates and zeolites are present, perhaps related to sealing of basement and restriction of fluid flow.



It is not known precisely when the seawater oxidation occurred, but there is no present evidence for continuing fluid circulation or advective heat transport at Site 1256 [Wilson *et al.*, 2003].

#### 4.10. Comparison With Current Models

[84] In a general model for the subsurface of submarine hydrothermal systems, Alt [1995] documented the effects of early alteration during seawater recharge followed by diffuse hydrothermal upflow and mineralization during subsurface mixing with seawater at the lavas-dikes boundary in ocean crust and ophiolites. It was proposed that this process is responsible for the observed stepped thermal gradient. Recent work based on new observations of altered crust have suggested that such stepped thermal gradients can only be maintained if all hydrothermal alteration occurs during fluid upflow [Barker *et al.*, 2008; Coogan, 2008; Gillis, 2008; Heft *et al.*, 2008].

[85] Heft *et al.* [2008] show that effects of upflow in the dikes are variable. In some areas fluid flow is of sufficiently short duration that only the initial alteration stages are observed, like the “background” alteration at Site 1256. Elsewhere, upflow develops and continues long enough for fluids to achieve black-smoker-like compositions and deposit veins of quartz  $\pm$  sulfide, as observed at Site 1256. Variable focusing of upflow results in differing intensities of alteration and mineralization in fault zones and during mixing at the lavas-dikes boundary [Honnorez *et al.*, 1985; Saccocia and Gillis, 1995; Gillis, 1995; Gillis *et al.*, 2001; Heft *et al.*, 2008; Barker *et al.*, 2010].

[86] Results from Site 1256 agree in general with these models. In particular, the upper dike section is characterized by the effects of hydrothermal upflow and mixing with seawater at the lava-dikes boundary. There is evidence for trace amounts of lower-temperature (<200°C) alteration in the plutonic section, but there is no evidence for significant low-temperature fluid flow in these rocks or the dike section.

[87] There is little anhydrite at depth in the Site 1256 dikes and none in the plutonic section, in contrast to what would be expected from significant circulation of little-reacted seawater at these depths. The main zone of anhydrite formation is within the upper dikes (at 1100–1200 mbsf), where seawater fluids were heated and mixed with hydrothermal fluids. This is consistent with observations of anhydrite in the one other drillhole to penetrate this lava-dike boundary [Alt *et al.*, 1986; Teagle *et al.*, 1998],

and as predicted for the zone where seawater is heated as temperatures increase downward [Mottl, 1983].

[88] Recent 3-D modeling of hydrothermal circulation at spreading axes shows cylindrical focused upflow zones, surrounded by dispersed downward recharge of seawater [Coumou *et al.*, 2008]. The upflow zones in the models are unstable, and migrate along-axis, consistent with interpretations of microearthquake data [Tolstoy *et al.*, 2008]. These fluid circulation models are based on a uniform basement permeability, and result in the 100°C isotherms outside the upflow zones being quite deep in the basement [Coumou *et al.*, 2008]. In contrast, circulation models that take into account the stepwise decrease in permeability downward from lavas to dikes result in shallower isotherms, with a step-like temperature increase corresponding to the lavas-dikes transition, as observed in oceanic basement [Rosenberg *et al.*, 1993]. In these models greater volumes of fluid circulate through the lavas, maintaining low temperatures there, whereas the dikes remain hot. Evidence from anhydrite in Hole 1256D suggests that anhydrite precipitates as seawater, partly reacted in the volcanic section, penetrates downward from the cooler lavas into the hot dikes. This could happen around the periphery of upflow zones, or at the site of a former upflow zone as these migrate along axis or as a crustal section moves away from focused upflow as the result of seafloor spreading.

## 5. Concluding Remarks

[89] Alteration of the upper crustal section at Site 1256 documents the thermal and chemical structure of a submarine hydrothermal system, and the evolution of fluid-rock interactions with time. A critical part of this is the interplay between fluid flow, lithology, and magmatism, from the mobile top of an axial magma chamber up to the seafloor. Results are generally consistent with current models for fast spread crust but there is little evidence for significant low-temperature fluid flow or significant cooling deep in the dikes. Planned deeper penetration at this site will allow evaluation of models for fluid flow, and chemical and heat transport from the lower oceanic crust.

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