29. PETROLOGY OF ULTRAMAFIC ROCKS FROM SITES 556, 558, AND 560 IN THE NORTH ATLANTIC

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ABSTRACT

Strongly serpentinized peridotites occur at shallow crustal levels at DSDP Sites 556 (35-Ma-old crust), 558 (35-Ma-old crust), and 560 (12-Ma-old crust) in the North Atlantic. Primary peridotite silicate minerals are preserved only at Sites 556 and 558, but all three sites have primary spinel preserved. The serpentinites at all three sites were probably spinel harzburgite tectonites, with little or no clinopyroxene.

For orthopyroxene from Site 556, Mg/(Mg + Fe) ratios average 0.904, and Al₂O₃ contents are about 2.0–3.3 wt.%. Compositions of pyroxene-derived bastites from Sites 558 and 560 suggest that the original pyroxenes were probably low-alumina orthopyroxene. Spinel from the three sites have low Fe⁢³⁺ and TiO₂. Spinel from Sites 556 and 560 have high Cr/(Cr + Al) (0.45–0.53). Those from Site 558 have Cr/(Cr + Al) values that are among the highest measured for any plagioclase-free oceanic peridotites (0.55–0.62).

Whole-rock chemistry, although affected by serpentinization, suggests that the peridotites had relatively low CaO, TiO₂, Al₂O₃, and Na₂O. Mineral and whole-rock chemistry and modal abundances suggest that Leg 82 peridotites are residua from partial melting, and that the extent of partial melting may have been greater in this region than farther south in the North Atlantic, or that the mantle was originally more depleted here.

INTRODUCTION

Serpentinized peridotites were recovered at DSDP Sites 556, 558, and 560 (Fig. 1). Site 556 was drilled in 35-Ma-old crust on a flow line away from the Azores Triple Junction. Peridotites were recovered under 461 m of sediments and 96 m of basalts. A cemented gabbronorite breccia directly overlies the serpentinized peridotites, which are interlayered with cemented serpentinite breccias, gabbronorites, and cemented gabbronorite breccias from 561–575 m subsurface. Gabbronorite and cemented gabbronorites are present from 575 m to the hole bottom at 639 m, with two narrow serpentinite horizons at about 620 m.

Site 556 was drilled in 35-Ma-old crust on a flow line from the FAMOUS area. Serpentinitized peridotites were recovered from 533 m depth to the hole bottom at 563 m. There are several serpentinite horizons with subtle color and textural differences. They are separated from each other by uncemented serpentinite breccias.

DSDP Site 560 was drilled in 12-Ma-old crust, midway between the Oceanographer and Hayes fracture zones. Serpentinitized peridotites were encountered directly under 373 m of sediments. Two serpentinite horizons, at 373 m and at about 400 m depth are separated by gabbronorites and gabbronorite breccias.

PETROGRAPHY

About ten standard 20 × 30-mm thin sections from Sites 556 and 558 and five from Site 560 were examined. Serpentinites from all three sites had peridotite protoliths. Only those from Site 556 have primary silicates preserved, and even these samples contain no olivine. Primary spinel is preserved in serpentinites from all three sites, however.

Peridotites from the three sites exhibit the pseudomorphous texture that is typical of many oceanic peridotites (Pritchard, 1979). The rock is composed predominantly of mesh-textured serpentine and magnetite. In plane polarized light, the color or tint of the mesh-textured serpentine may vary from clear to light green or brown, and the sharpness of the magnetite framework is also variable. The mesh is the alteration product of olivine, which is generally the least stable phase during serpentization. Within the mesh matrix are large (up to 10 mm) bastites or serpentized pyroxene porphyroclast relics, which preserve the shape and fine details of the internal structure of the pyroxene, including kink-band ing and exsolution lamellae.

It is difficult to discern the difference between orthopyroxene bastite and clinopyroxene bastite, or to reconstruct the original modes of the samples. Samples from Site 556 have islands of orthopyroxene preserved within the bastites. Small blebs of clinopyroxene, exsolved from the orthopyroxene in these samples, suggest that clinopyroxene was more resistant than orthopyroxene to serpentization. The lack of clinopyroxene porphyroclasts is therefore a primary feature of the peridotite from Site 556. Sites 558 and 560 required a less direct approach for modal reconstruction. Petrographic and microprobe analysis of partly serpentinitized peridotites from other locations (the Owen Fracture Zone), in which clinopyroxene and orthopyroxene porphyroclasts and their enclosing bastites are preserved, suggests that clinopyroxene bastite often has a different appearance than orthopyroxene bastite. For example, relict exsolution lamellae.
often "die out" toward the edges of clinopyroxene bastites. Clinopyroxene porphyroclasts generally have higher Cr contents than coexisting orthopyroxene porphyroclasts, and Cr concentrations and concentration differences are reflected in the bastites as well (Bonatti et al., in preparation^3^). With these observations in mind, we examined the bastites from Sites 558 and 560, and approximately 20 bastites from two sections of each site were analyzed for Cr content. The observations suggest that little or no porphyroclastic clinopyroxene was present in peridotites from Sites 558 and 560. Thus, even though the silicates have been completely serpentinized, it is possible to infer original peridotite modes and textures (Table 1). The reconstructions suggest that the samples were originally spinel harzburgite tektontites.

Spinel is preserved in all of the serpentinized peridotites. It is generally brick-red, with a varied habit, shape and association and is often contained within or near orthopyroxene or orthopyroxene bastite. In this association, spinel forms as long, curved grains (up to 8 mm) with xenomorphic texture around orthopyroxene; as symplectite-textured clusters of small grains (0.1-0.5 mm); and as small grains near orthopyroxene. Similar associations of spinel with orthopyroxene in peridotites have been described by other authors (Hamlyn and Bonatti, 1980; Smith, 1977) and ascribed to the reaction: olivine + garnet → spinel + pyroxene. Spinel also occurs within the olivine-derived mesh-textured serpentine, as rounded or rectangular porphyroclasts (0.3-3 mm) or as tiny (0.1-0.3 mm) disrupted grains scattered throughout a larger (~1 cm) region.

Opaque spinel is present in three associations: (1) with clinopyroxene and plagioclase (An^3^), a probable gabbroic inclusion; (2) as an alteration product along the edges and cracks in red spinel; and (3) as complete grains, possibly also as a reaction product.

Veins of calciferous and noncalciferous amphiboles are variably present in the peridotites. Where they cut the rock, the nearby pyroxenes have been transformed into predominant amphibole and lesser serpentine. This may represent an earlier, high-temperature hydration of the original peridotite, and the amphibole may be more resistant to serpentinization than the primary, anhydrous silicates.

### MINERAL CHEMISTRY

Mineral compositions were obtained with an automated, three-channel ARL-EMX microprobe at Lamont-Doherty Geological Observatory, using a 15-kV accelerating voltage, and a 0.15 µA beam current. Data were reduced online using a Bence-Albee procedure and reference to a suite of natural and synthetic mineral standards. For spinel analyses, a narrow (3-5 µm) beam was

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^3^ Bonatti, E., Morandi, N., Brueckner, H., and Michael, P. J., Progressive serpentinization of mantle-derived ultramafic rocks from the Owen Fracture Zone.
used. Fe\(^{2+}\) and Fe\(^{3+}\) were calculated assuming stochiometry. A broad (70 µm) microbeam was used to obtain a more representative analysis of the heterogeneous orthopyroxenes and bastites. The mineral compositions presented in Tables 2-4 are based on averages of more than four spot analyses in each of two separate grains.

**Orthopyroxene.** Site 556 peridotites have orthopyroxenes with high and variable Mg/(Mg + Fe) ratios, averaging 0.904 (Table 2). These ratios are slightly higher than those of pyroxenes from the Vema Fracture Zone (Hamlyn, unpublished data, 1982) and DSDP Site 395 (Sinton, 1979), but similar to those from the Kane and Atlantis fracture zones. The ratios are lower than orthopyroxenes from the Oceanographer Fracture Zone (Fig. 2).

Orthopyroxene alumina contents are quite low, averaging only 2.8% (Table 2). Site 556 orthopyroxenes contain less alumina than orthopyroxenes from most other oceanic peridotites, a feature shared by some other North Atlantic peridotites (Fig. 3). Site 556 orthopyroxenes approach the compositions of pyroxenes from some alpine peridotites, which are probably more depleted than oceanic peridotites (Dick, 1975). Chrome concentrations are quite high, about 1%, and Cr/(Cr + Al) is about 0.16-0.19; higher than other oceanic peridotites because of the low alumina. Orthopyroxene has about 3-5 mole% wollastonite component, which is similar to orthopyroxene of other oceanic peridotites. Measurement of this quantity is difficult because of the large-scale heterogeneity caused by exsolution.

Bastites of serpentinites from Sites 558 and 560 have very low alumina concentrations, about 1.2-2%. Preliminary work (Bonatti et al., in preparation) suggests that bastite’s alumina content reflects to some degree the alumina content of the original pyroxene from which it derives, and that Al has limited mobility during the serpentinization process. Thus, orthopyroxenes from Sites 558 and 560 may have also had low alumina.

**Clinopyroxene.** In serpentinized peridotites from Site 556, clinopyroxene occurs mostly as tiny (0.1-0.5 mm) blebs included within or spatially associated with much larger orthopyroxenes or orthopyroxene bastites. These are most likely exsolution products from the orthopyroxene. In one sample (556-13-1, 6-8 cm), a large primary clinopyroxene grain is associated with plagioclase -An\(_{55}\), and high-Ti spinel. This assemblage might be an inclusion from the surrounding gabbros, since the interval is adjacent to a gabbro breccia.

Exsolved clinopyroxene has high Mg/(Mg + Fe) ratios (0.907-0.921) and low alumina contents (2.30-4.48 wt.%) (Table 3). TiO\(_2\) and Na\(_2\)O are positively correlated and quite variable. Chrome contents (about 1%) are lower than those observed from nonexsolved clinopyroxenes from other oceanic peridotites (Sinton, 1979).

**Spinel.** Spinel has variable and high Cr concentrations and very low Ti and Fe\(^{3+}\) (calculated assuming stochiometry; Table 4). Spinel from Site 558 are among the most chromium-rich spinels measured for any oceanic peridotites (Cr/[Cr + Al] -0.58), and spinels from Site 556 and 560 are only slightly less chromium rich (Fig. 4).

Opaque spinel, which is associated with plagioclase and clinopyroxene, has high TiO\(_2\) (up to 3 wt.%) and Fe\(^{3+}\) and is more similar to spinel, which is in equilibrium at low pressure in basalts (Sigurdsson and Schilling, 1976). Opaque spinel that invades cracks and margins of brick red spinel approaches pure magnetite in composition. The third association of opaque spinel, as discrete grains, is slightly more iron-rich than coexisting brick-red spinel in a similar association, and underscores the fact that spinel compositions vary within individual grains and between grains within thin section. The significance of the variation is not clear, but has been ascribed (Hamlyn and Bonatti, 1980) to subsolidus reequilibration by the reaction: orthopyroxene + clinopyroxene + Al-spinel - olivine + plagioclase + Cr-spinel. Recently, Ozawa (1983b) noted that intergrain and intra-grain variations in spinel chemistry arose from incomplete reequilibration of spinel with surrounding pyroxenes and olivine. Spinel compositional variation can also arise from the serpentinization reaction (Hoffman and Walker, 1978). Thus, the spinel compositional variation that is observed between different drilling sites or fracture zones may reflect the variation of the original peridotite protolith, or possibly later reactions.

**WHOLE-ROCK MAJOR ELEMENT CHEMISTRY**

Major element compositions were determined by standard X-ray fluorescence (XRF) methods, using a Siemens SRS-1 spectrometer and employing the sample preparation procedures of Harvey and others (1973). The results are shown in Table 5. Evaluation of these data must take into account that the samples are highly serpentinized, and that serpentinization, in addition to increasing the water content of the rock, may involve some elemental exchange.

The relatively high CaO and LOI in Samples 556-14-1, 61-65 cm and 558-41-3, 111-116 cm are due to
carbonate veins in the sample. Thus, in these two samples loss on ignition (LOI) must include significant quantities of carbonate-derived CO₂. If the data are recalculated on a LOI-free basis, the Al₂O₃ content of all samples, which should not be greatly affected by serpentinization, appears to be low compared to other oceanic serpentinites, paralleling the low Al content of orthopyroxenes, clinopyroxenes, and bastites. CaO, K₂O, and Na₂O are also rather low, but these elements, particularly CaO, are probably affected by serpentinization.

**DISCUSSION**

The following points can be drawn from the study of the ultramafic rocks recovered during DSDP Leg 82.

1. The rocks appear to be spinel harzburgites and to have undergone intense serpentinization. Their relic textures and the chemistry of relic pyroxene and spinel suggest they are tectonites of upper mantle derivation.

2. Pyroxenes have low Al content, which might be related to greater extent of melting compared to other analyzed oceanic ultramafics (Mysen and Kirshiro, 1977; Jaques and Green, 1980). Relatively high Mg/(Mg + Fe) and low Na contents both of pyroxenes and of whole rock suggest that we are dealing with residual material, depleted of a basaltic fraction.

3. Spinel is in general rich in Cr, but their composition is variable in these rocks, suggesting that their equilibrium with surrounding pyroxenes and olivine was incomplete. Although spinel composition may be related to the serpentinization process, its correlation with modal abundance of pyroxene suggests that it is primarily a function of the peridotite protolith composition.

4. Ultramafic rocks from DSDP Sites 556, 558, and 560, recovered in crust 35 Ma old (556 and 558) and 12 Ma old (560), are rather similar to those from Site 395, recovered in crust about 7 Ma old, though the latter con-
Table 3. Selected microprobe analyses of clinopyroxene from Hole 556 peridotites.

<table>
<thead>
<tr>
<th>Core-Section (interval in cm)</th>
<th>13-1</th>
<th>13-1</th>
<th>14-1</th>
<th>14-1</th>
<th>14-2</th>
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<tr>
<td>Major elements (wt. %)</td>
<td></td>
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<tr>
<td>SiO₂</td>
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<td>51.87</td>
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<td>TiO₂</td>
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<td>0.78</td>
<td>0.73</td>
<td>0.14</td>
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<td>Al₂O₃</td>
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<td>2.30</td>
<td>2.96</td>
<td>3.72</td>
<td>3.05</td>
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<td>CaO</td>
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<td>1.11</td>
<td>1.06</td>
<td>1.46</td>
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<tr>
<td>FeO</td>
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<td>2.22</td>
<td>2.76</td>
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<td>MnO</td>
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<td>0.07</td>
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<td>MgO</td>
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<td>17.66</td>
<td>16.97</td>
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<td>GaO</td>
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<td>22.29</td>
<td>21.59</td>
<td>22.57</td>
<td>22.92</td>
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<tr>
<td>Na₂O</td>
<td>0.04</td>
<td>0.31</td>
<td>0.42</td>
<td>0.38</td>
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<td>Total</td>
<td>100.99</td>
<td>100.25</td>
<td>100.69</td>
<td>99.98</td>
<td>100.01</td>
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Note: All except 13-1, 6-8 cm are exsolution blebs from much larger orthopyroxene. All iron (Fe*) is assumed to be Fe²⁺.

Cations on the basis of 6 oxygens

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<thead>
<tr>
<th></th>
<th>Si</th>
<th>Ti</th>
<th>Al</th>
<th>Cr</th>
<th>Fe²⁺</th>
<th>Fe³⁺</th>
<th>Mn</th>
<th>Ni</th>
<th>Mg</th>
<th>Ca</th>
<th>Na</th>
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<td></td>
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<td>3.997</td>
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<tr>
<td>Fs</td>
<td>48.59</td>
<td>50.13</td>
<td>50.85</td>
<td>48.92</td>
<td>49.02</td>
<td>4.20</td>
<td>5.05</td>
<td>4.46</td>
<td>4.32</td>
<td>4.19</td>
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<tr>
<td>Wo</td>
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<td>44.69</td>
<td>46.77</td>
<td>46.79</td>
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<td>5.05</td>
<td>4.46</td>
<td>4.32</td>
<td>4.19</td>
<td>100.99</td>
<td>100.25</td>
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Table 4. Selected microprobe analyses of nonopaque spinels from peridotites of Sites 556, 558, and 560.

<table>
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<tr>
<th>Hole</th>
<th>Core-Section (interval in cm)</th>
<th>13-1</th>
<th>13-1</th>
<th>14-1</th>
<th>14-1</th>
<th>14-2</th>
<th>42-1</th>
<th>42-1</th>
<th>43-1</th>
<th>43-47</th>
<th>44-1</th>
<th>44-7</th>
<th>4-1</th>
<th>5-1</th>
<th>81-84</th>
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<tbody>
<tr>
<td>Major elements (wt. %)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.30</td>
<td>0.00</td>
<td>0.12</td>
<td>0.00</td>
<td>0.11</td>
<td>0.02</td>
<td>0.02</td>
<td>0.00</td>
<td>0.02</td>
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<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
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<tr>
<td>Cr₂O₃</td>
<td>41.64</td>
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<td>45.51</td>
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<td>FeO</td>
<td>3.52</td>
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<tr>
<td>Total</td>
<td>99.59</td>
<td>99.35</td>
<td>99.61</td>
<td>99.29</td>
<td>100.15</td>
<td>100.84</td>
<td>100.29</td>
<td>100.26</td>
<td>100.35</td>
<td>100.45</td>
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Cations on the basis of 4 oxygens

<table>
<thead>
<tr>
<th></th>
<th>Cr</th>
<th>Cr + Al</th>
<th>Mg</th>
<th>Mg + Fe²⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.525</td>
<td>0.456</td>
<td>0.475</td>
<td>0.444</td>
</tr>
<tr>
<td>Cr + Al</td>
<td>0.630</td>
<td>0.613</td>
<td>0.573</td>
<td>0.618</td>
</tr>
<tr>
<td>Mg</td>
<td>0.551</td>
<td>0.551</td>
<td>0.551</td>
<td>0.551</td>
</tr>
</tbody>
</table>

Note: Fe²⁺ and Fe³⁺ calculated assuming stoichiometry.

5. The recoveries of ultramafic rocks at relatively shallow levels in the crust away from transform zones confirm earlier suggestions that mantle-derived ultramafic bodies are an important component of slow spreading, Atlantic-type crust. The complex stratigraphy at Holes 556, 558, and 560 suggests a major role of faulting in the shallow emplacement of ultramafics in the crust, as proposed by Bonatti (1976) and Francis (1981). The observation that peridotite tectonites are sandwiched between gabbro layers suggests that only the topmost layer of peridotites is being sampled.

6. Comparison of ultramafics from non-fracture-zone Sites 556, 558, and 560 with ultramafics recovered from nearby fracture zones (i.e., Oceanographer Fracture Zone at 35°N, see Fig. 1) suggests that fracture zones contain some ultramafic rocks that are slightly less refractory than ultramafic rocks recovered in nearby non-fracture-zone crust. This observation is supported by studies at DSDP Site 395 and the Kane Fracture Zone at about 24° (Michael and Bonatti, in preparation). It is possible that the upper mantle associated with fracture zones may have undergone less extensive melting than nearby upper mantle material (Michael and Bonatti, in preparation). Similarly, comparison between Leg 82 and Oceanographer Fracture Zone ultramafics with ultramafics from farther south in the Atlantic (see, for instance, Fig. 4),
suggests systematic regional differences in upper mantle mineral chemistry and modes. The differences between the peridotite protoliths may be related either to greater degree of partial melting in the northernmost region or

to preexisting heterogeneities in the mantle, as further discussed elsewhere (Michael and Bonatti, 1983; Dick and Bryan, 1983; Michael and Bonatti, in preparation).

NOTE ADDED IN PROOF

Subsequent sampling of peridotites from Hole 560 revealed the presence of primary orthopyroxene porphyroclasts averaging 2.8% Al$_2$O$_3$ and 91.1 Mg/(Mg + Fe) x 100.

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REFERENCES


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