Sm-Nd and Rb-Sr Systematics in Volcanics and Ultramafic Xenoliths From Malaita, Solomon Islands, and the Nature of the Ontong Java Plateau

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A Rb-Sr and Sm-Nd study was carried out on five ultrabasic inclusions, three alnoite host rocks, and one sample of basalt from Malaita in an effort to determine the isotopic stratigraphy of the mantle in this area. The results of this study show the presence of relatively undepleted mantle segments underlying relatively older depleted upper mantle. The alnoite samples, three megacrysts and one lherzolite inclusion, and the basalt are relatively uniform isotopically and yielded eNd(T) ≈ +3.5 and eNd(T) ≈ −6 to +17 (T = 100 m.y. for the basalt and lherzolite and 34 m.y. for the rest of the samples). It is suggested that a mantle segment of that isotopic composition exists at a depth of more than 100 km. This mantle segment is relatively undepleted. A lherzolite inclusion with eNd(T) = 6.6 is included in the alnoite and indicates the location of a light rare earth element depleted layer overlying a less depleted mantle at a depth of approximately 100 km. The mantle source for the alnoite and basalt cannot be old, depleted oceanic mantle or subducted old continental crust but may come from a relatively young, slightly depleted mantle segment. The isotopic results are compatible with a mixture of about 99% depleted oceanic mantle homogenized with 1% continental crust possibly by metasomatic processes. Such a mixture is only slightly fractionated and has $f^{\text{Sm/Nd}} \approx 0.13$ and $f^{\text{Rb/Sr}} \approx 0.0$. Assuming that the basalt is characteristic of the Ontong Java Plateau, a volume greater than $10^4$ km$^3$ of mantle source with $e_{\text{Nd}} = 3.5$, $e_{\text{Sr}} = 0$ was involved. This corresponds to a block of continental crust of $\sim 10^5$ km$^3$ which had to reach a depth greater than 100 km to produce the proposed nearly homogeneous isotopic mixture.

Oceanic plateaus are anomalous regions embedded in the earth’s seafloor, located mainly in the western Pacific and the Indian Ocean [see Nur and Ben Avraham, 1982]. They are distinguished by their shallow water depth, thick crust, and low upper crustal seismic velocities. Numerous explanations have been advanced for the nature and origin of the oceanic plateaus. As summarized by Nur and Ben Avraham [1982], they include extinct arcs, abandoned spreading ridges, detached continental fragments, anomalously thick oceanic piles, hot spots, and uplifted oceanic crust.

The origin of the OJP is somewhat controversial. Its thick crust, up to 40 km, of Lower Cretaceous age and the details of its seismic profile led Carlson et al. [1980] and Nur and Ben Avraham [1982] to propose that the OJP represents a continental block. Nur and Ben Avraham suggested that the plateau is a remnant of Pacifica (a hypothetical eastern extension of Gondwanaland). Kroenke [1974] suggested that the excess thickness of the basaltic rocks represented flood basalts which mark the beginning of continental formation. Nixon and Coleman [1978] and Nixon [1980] support the view that the OJP is a "protocontinent" and predict the existence of true kimberlites under the thicker parts of the plateau. According to Kroenke [1974] and Coleman and Kroenke [1981], the plateau was formed along the crest of a slow spreading ocean ridge. The slow spreading, according to this view, is responsible for the thick crust of the plateau. In this model the magmas which formed the plateau would presumably be derived from depleted oceanic mantle [Tatsumoto et al., 1965; Gast, 1968], similar to other mid-ocean ridge basalts (MORB) yielding a characteristic, isotopic signature of old, depleted mantle [DePaolo and Wasserburg, 1976a; Richard et al., 1976]. From their study of seismic data, Hussong et al. [1979] proposed that the OJP consists of normal oceanic crust simply expanded by a factor of 5 in thickness. This would also show the above isotopic characteristics.

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Hughes and Turner [1977] described the basaltic rocks of Malaita. The more extensive basalts (more than 2 km are exposed in central Malaita) belong to the old, tholeiitic phase assumed to be part of the Lower Cretaceous OJP volcanism. The younger (Upper Cretaceous to Eocene) alkalic phase is much less voluminous and is restricted only to the southern part of the island. On the basis of chemical compositions, Hughes and Turner pointed out that the old tholeiitic phase (consisting of flows of nonpillowed and pillowed lavas) shows similarities to both oceanic island tholeiites and ocean ridge tholeiites. On the basis of field evidence, they favor the idea that the Malaita basalts resemble oceanic island volcanism because they consider it unlikely that the basalts were derived from extensional rift features where dikes would be expected to be much more common. From surveying these hypotheses it appears that there is no consensus concerning the nature and origin of the OJP. It is, however, remarkable that no xenoliths of continental crust were found in the basalts nor were any found among the xenoliths of the alnoite which intruded the OJP during the Tertiary.

3. The Alnoite

Field evidence [Nixon and Coleman, 1978] suggests that the alnoite is a pipelike intrusion emplaced in steeply folded Upper Cretaceous and lower Tertiary mudstones and limestones (Figure 2). These sedimentary rocks are found as xenoliths in the alnoite [Nixon and Boyd, 1979]. This age relationship is in accord with the U-Pb age of 34 m.y. [Davis, 1978] obtained by dating a zircon xenocryst from the alnoite. The Cretaceous sediments which are crosscut by the alnoite overlie basalt, pillow lavas, and tuffs (the Alite Volcanics) of tholeiitic composition [Hughes and Turner, 1977].

The alnoite, which is described in detail by Allen and Deans [1965] and Nixon et al. [1980], consists mainly of fragmented rocks grading from tuffs to soft breccias, but a fresh hard aphanitic variety is also present. Primary minerals in the fresh alnoite are olivine (Fo 85-90), melilite up to 30%, clinopyroxene, phlogopite (5-55%), perovskite, and spinel. Accessory minerals are nepheline, melilite, and apatite. The fragmented rocks are rich in calcite (more than 50%) and contain coarse, rounded phlogopite (>1 cm). Coarse phlogopite is often found in the cores of autoliths, which are subcircular structures composed of alnoitic magma wrapped around a rock or mineral inclusion. Nixon et al. [1980] suggested that the alnoite is a primary melt of a pyrolite-type mantle formed by approximately 4% partial melting at depths greater than 120 km, under high CO2 pressure. The only previous isotopic data on this rock type from the same locality were a 143Nd/144Nd ratio determination by Basu and Tatsumoto [1980]. These workers compared this result with kimberlites and concluded that they came from a different source.

4. The Xenoliths

As reported by Nixon and Boyd [1979], the proportion of country rock sedimentary fragments, ultramafic xenoliths, and discrete monomineralic nodules in the alnoite varies but is generally greater than 10%. Lherzolites predominate among
Fig. 2. Stratigraphic section of the upper crust in northern Malaita (Solomon Islands) using information from Rickwood [1957] and other sources referred to in the text.

The mantle ultramafics. They consist of olivine (50–70%), orthopyroxene (10–35%), and clinopyroxene (5–40%). Some of the specimens might have garnet (up to 18%) and/o r spinel or chromite (up to 7%). The discrete monomineralic nodules are large, rounded, single crystals, megacrysts of garnet, cpx, opx, and ilmenite. Nixon and Boyd used the xenoliths and megacrysts to construct a paleogeotherm. Compared with similarly derived geotherms from garnet-bearing nodules in continental Kimberlites, the Malaita geotherm has a higher gradient, reflecting a higher heat flow. A high-temperature inflection is interpreted as the occurrence of the lithosphere-asthenosphere transition at a depth of 110 km. Figure 3 (taken from Nixon and Boyd [1979]) gives their proposed depth profile of the lithosphere and top of asthenosphere under Malaita. Not all of these depths are well defined by geobarometers. The discrete nodules are believed to come from the top of the asthenosphere at a depth of 110–120 km. The lherzolites are considered to be representative of the overlying lithosphere from 110 to 60 km depth. They include both deeper seated depleted and undepleted samples as determined by large ion lithophile (LIL) element abundances. The alnoitic magma must have cut across all of these lithic units during its ascent.

5. Samples

Ten samples from the Malaita alnoite and an older basalt were used in this study. They include four alnoite samples, five xenoliths and megacrysts found in the alnoite, and one sample of the country rock basalt believed to be part of the Lower Cretaceous volcanic phase of the OJP. A short description of the samples is given below. More information can be found in the references cited before. No depth assignment can be made on rocks in which garnet is absent.

MALAITA OCEANIC MODEL

<table>
<thead>
<tr>
<th>Depth, km</th>
<th>Succession</th>
<th>Spinel/Garnet</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–30</td>
<td>Basaltic Crust</td>
<td></td>
</tr>
<tr>
<td>30</td>
<td>Harzburgite (very depleted)</td>
<td>Cr spinel</td>
</tr>
<tr>
<td>40</td>
<td>Lherzolite</td>
<td>Al spinel</td>
</tr>
<tr>
<td>60</td>
<td>Depleted lherzolite (i.e., Removal of basalt and loss of Al relative to Cr)</td>
<td>Garnet</td>
</tr>
<tr>
<td>80</td>
<td>(ilmenite-rich zone) Zone rich in large crystals of garnet and pyroxene</td>
<td>Cr Al spinel &amp; Cr garnet</td>
</tr>
<tr>
<td>100</td>
<td>Source of Alnoites and Basalts</td>
<td>LVZ</td>
</tr>
<tr>
<td>&gt;140</td>
<td>(ilmenite-rich zone) Zone rich in large crystals of garnet and pyroxene</td>
<td>Garnet</td>
</tr>
</tbody>
</table>

Source of Alnoites and Basalts

Fig. 3. Stratigraphic cross section of the lower crust mantle underlying Malaita from a study of xenoliths by Nixon and Boyd [1979] (slightly modified).
TABLE 1. Rb-Sr and Sm-Nd Analytical Data

<table>
<thead>
<tr>
<th>Sample</th>
<th>Rb, ppm</th>
<th>Sr, ppm</th>
<th>Sm, ppm</th>
<th>Nd, ppm</th>
<th>( f^{186}\text{Rb}/f^{86}\text{Sr} )</th>
<th>( f^{147}\text{Sm}/f^{144}\text{Nd} )</th>
<th>( \Delta f^{147}\text{Nd}/f^{144}\text{Nd} )</th>
<th>( e_{\text{Rb}}(T)^{\ast} )</th>
<th>( e_{\text{Nd}}(T)^{\ast} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>PHN-3935</td>
<td>79.1</td>
<td>1380</td>
<td>19.4</td>
<td>100</td>
<td>+1.00</td>
<td>-0.40</td>
<td>0.70461 ± 3</td>
<td>0.511992 ± 25</td>
<td>1.0</td>
</tr>
<tr>
<td>PHN-3855</td>
<td>61.4</td>
<td>3055</td>
<td>17.2</td>
<td>89.7</td>
<td>-0.30</td>
<td>-0.40</td>
<td>0.70467 ± 6</td>
<td>0.511956 ± 28</td>
<td>2.6</td>
</tr>
<tr>
<td>PHN-3554</td>
<td>32†</td>
<td>827</td>
<td>13†</td>
<td>64.4</td>
<td>+0.35</td>
<td>-0.34</td>
<td>0.70455 ± 7</td>
<td>0.512002 ± 28</td>
<td>0.5</td>
</tr>
<tr>
<td>PHN-3544</td>
<td>74†</td>
<td>1188‡</td>
<td>24.4‡</td>
<td>143.7‡</td>
<td>+1.18</td>
<td>-0.47</td>
<td>-0.38</td>
<td>-0.15</td>
<td>-0.6</td>
</tr>
<tr>
<td>PHN-3995</td>
<td>0.5†</td>
<td>17.1</td>
<td>0.2†</td>
<td>0.093</td>
<td>+0.02</td>
<td>-0.33</td>
<td>0.70510 ± 5</td>
<td>0.52167 ± 27</td>
<td>8.5</td>
</tr>
<tr>
<td>PHN-3852</td>
<td>0.03†</td>
<td>48.8†</td>
<td>2.1†</td>
<td>12.0†</td>
<td>-0.98</td>
<td>-0.46</td>
<td>0.70440 ± 7</td>
<td>0.511946 ± 15</td>
<td>-0.9</td>
</tr>
<tr>
<td>PHN-3861</td>
<td>13†</td>
<td>31†</td>
<td>1.1†</td>
<td>3.7†</td>
<td>+0.43</td>
<td>-0.08</td>
<td>0.70571 ± 2</td>
<td>0.512002 ± 15</td>
<td>16.9</td>
</tr>
<tr>
<td>PHN-3866A</td>
<td>0.72†</td>
<td>10.2</td>
<td>1.1</td>
<td>1.48</td>
<td>+1.47</td>
<td>+1.47</td>
<td>0.70453 ± 4</td>
<td>0.512086 ± 15</td>
<td>-0.4</td>
</tr>
<tr>
<td>PHN-3916</td>
<td>1.9†</td>
<td>107†</td>
<td>1.9†</td>
<td>6.96</td>
<td>-0.38</td>
<td>-0.15</td>
<td>0.70406 ± 4</td>
<td>0.512055 ± 14</td>
<td>-6.0</td>
</tr>
</tbody>
</table>

* \( T = 100 \text{m.y.} \) for the basalt (PHN-3916) and \( T = 34 \text{m.y.} \) for all the rest of the samples.
† Concentration determined by aliquot only (see text).
‡ Data from Nixon et al. [1980].
§ Data from Basu and Tatsumoto [1980], using \( 150\text{Nd}/144\text{Nd} = 0.23643 \) for normalization of isotopic data.

PHN-3935. Tough fresh alnoite. Partly serpentinized olivine phenocrysts. Minor calcite is associated with deformed phlogopite.


PHN-3554. Soft alnoite agglomerate with spherical autoliths. About 50% calcite.

PHN-3544. Fresh tough alnoite (description and chemical analysis from Nixon et al. [1980]).

PHN-3995. Tough unserpentinized spinel lherzolite (olivine 60–70%, cpx ~ 30%) with a deformatonal texture: defined by layers of recrystallized cpx granules (~1 mm) and much coarser, aligned spinels and opx (up to 7 mm). Spinel granules (0.1 mm) are conspicuously developed at grain boundaries.

PHN-3852. Cpx (chromium diopside) separated from a spinel lherzolite: olivine (60%), opx (20%), cpx (5%), serpentine and calcite rims surrounding fresh grains.

PHN-3861. Coarse garnet-opx-cpx xenolith with cumulate texture. Garnet is highly kelyphitic.

PHN-3866A. Discrete garnet (megacryst). Fractured but fresh.

PHN-3968. Discrete cpx (megacryst) (originally 1-kg nodule). A few spherical inclusions containing phlogopite and serpentine.


6. Analytical Method

The analytical method used for the determination of Rb, Sr, Nd, and Sm concentration as well as Sr and Nd isotopic composition is described by Papanastassiou et al. [1977] and DePaolo [1978]. Element concentrations in some of the samples (Table 1) were determined only by an aliquot run in which the four elements are spiked and measured in a single mass spectrometric run prior to chemical separation. Uncertainties associated with these concentration values are typically better than 5%. Samples which were totally spiked and separated have concentrations determined to better than ~0.1%.

Sample PHN-3995, because of its low rare earth element (REE) concentration, was processed in a somewhat different way: 1 g of the sample powder was dissolved, aliquot concentrations were determined, and accurate spiking was performed the usual way. Then the sample solution was split into two fractions: 0.1 g was processed for Rb-Sr determinations. To the remaining 0.9 g, a few milliliters of pure ammonia solution was added until iron precipitation as Fe(OH)₃ was completed. The REE were adsorbed on this precipitate [Goldberg et al., 1963] and more than 95% of the REE, as determined for Eu and Nd, were recovered. This removed most of the other major elements. The precipitate was centrifuged, separated from the supernate and dissolved in HCl. This solution, containing the REE fraction, was then processed through the separation columns following the usual procedure.

Rather pure cpx was hand-picked from a lherzolite (PHN-3852) and then cleaned from small, attached olivine and possible alteration products by soaking in HF solution. Olivine disintegrated, and the cpx remained unattacked.

7. Data Representation

Rb-Sr and Sm-Nd data are summarized in Table 1. The enrichment factors \( f^{186}\text{Rb}/f^{86}\text{Sr} \) and \( f^{147}\text{Sm}/f^{144}\text{Nd} \) provide a measure of the deviation of Rb/Sr and Sm/Nd of a rock from a model primitive undifferentiated bulk earth source composition. An estimate for the REE composition of undifferentiated bulk earth (CHUR) was given by DePaolo and Wasserburg [1976]. The revised values for CHUR, given by Jacobsen and Wasserburg [1980] and Wasserburg et al. [1981] will be used here. The unfractionated reservoir values (UR) for the bulk earth Rb and Sr are taken from DePaolo and Wasserburg [1976b] and
O’Nions et al. [1977].

\[ f^{\text{Sm/Nd}} = \frac{[\text{Sm/Nd}][/\text{Sm/Nd}]_{\text{CHUR}} - 1}{(143^{\text{Nd}}/144^{\text{Nd}})_{\text{CHUR}} = 0.1967} \]

\[ f^{\text{Rb/Sr}} = \frac{[\text{Rb/Sr}[/\text{Rb/Sr}]_{\text{CHUR}} - 1}{(87^{\text{Rb}}/86^{\text{Sr}})_{\text{CHUR}}} = 0.0827 \]

Here “m” represents the measured value. Isotopic ratios are expressed in \( \varepsilon \) notation [DePaolo and Wasserburg, 1976], where \( \varepsilon (T) \) is the measured deviation in parts of 10\(^6\) from the isotopic ratio from present-day CHUR or UR values.

\[ \varepsilon^{\text{Nd}}(T) = \left[ \frac{(143^{\text{Nd}}/144^{\text{Nd}})_{\text{CHUR}} - 1}{(143^{\text{Nd}}/144^{\text{Nd}})_{\text{CHUR}}} \right] \times 10^4 \]

\[ \varepsilon^{\text{Sr}}(T) = \left[ \frac{(87^{\text{Sr}}/86^{\text{Sr}})_{\text{CHUR}} - 1}{(87^{\text{Sr}}/86^{\text{Sr}})_{\text{CHUR}}} \right] \times 10^4 \]

The evolution of the isotopic deviation with age can be calculated for any age (T) of interest. A useful approximation [DePaolo and Wasserburg, 1976] was used to calculate those values:

\[ \varepsilon_{\text{Nd}}(T) = \varepsilon_{\text{Nd}}(0) - Q_{\text{Nd}} f^{\text{Sm/Nd}} T \]

\[ Q_{\text{Nd}} = 24.74 \text{ aeon}^{-1} \]

\[ \varepsilon_{\text{Sr}}(T) = \varepsilon_{\text{Sr}}(0) - Q_{\text{Sr}} f^{\text{Rb/Sr}} T \]

\[ Q_{\text{Sr}} = 16.70 \text{ aeon}^{-1} \]

The values of \( \varepsilon (T) \) for the alnoite and xenolith samples was calculated with \( T = 34 \text{ m.y.} \), the U-Pb age of the alnoite intrusion. For the xenoliths and megacrysts this is a minimum age but is useful for investigating the alnoite-xenolith genetic relations. The \( \varepsilon (T) \) for the basalt sample was calculated with \( T = 100 \text{ m.y.} \) This is an estimate for the age of the intrusive volcanic episode, responsible for the formation of the Ontong Java Plateau. The stratigraphic age of those volcanics is at least as old as Early Cretaceous [Hughes and Turner, 1977]. As the \( f^{\text{Sm/Nd}} \) value of the basalt is close to zero, \( \varepsilon_{\text{Nd}}(T) \) is not sensitive to a possible error in the chosen age.

We will show that the resulting \( \varepsilon(T) \) values calculated with the above mentioned ages are very similar for all the samples (excluding only the lherzolite). Using another set of ages will cause a scatter of \( \varepsilon(T) \) values. We will conclude that the homogeneity of \( \varepsilon_{\text{Nd}}(T) \) is accidental and indicates genetic relations between the samples.

8. Results

8.1. The Alnoite and Basalt

The alnoite samples are all LIL and REE enriched by a large factor in comparison to any bulk earth values or even to average continental crustal rocks: up to 80 ppm Rb, 3050 ppm Sr, 25 ppm Sm, and 145 ppm Nd (Table 1). Some of these samples have La/Lu of ~45 [Nixon et al., 1980]. Samples PHN-3935, PHN-3855, and PHN-3544 are very similar in their concentration levels except for the Sr in PHN-3855 which is much higher, possibly due to a high mellite content. PHN-3554 does not show such high enrichments. This sample is a soft agglomerate with 50% calcite, whereas the other three samples are aphanitic alnoite, poor or devoid of calcite. We believe that as calcite is low in REE, it is a diluant for PHN-3554. The \( f^{\text{Sm/Nd}} \) values for all the samples are very similar around -0.4 reflect a high light rare earth element (LREE) enrichment compared with a chondritic Sm/Nd ratio. A more complete REE spectrum had previously been measured by Nixon et al. [1980], who showed the high LREE enrichment. The high LIL and REE content and strong LREE enrichment are very typical of alkali basalts, kimberlites, and carbonatites [Haskin et al., 1966; Kay and Gast, 1973]. This is an additional indication that the alnoite belongs genetically to this group of rocks.

The \( f^{\text{Rb/Sr}} \) values range from -0.3 to +1.18. Leaving out the only negative value (-0.3) corresponding to the mellite-rich sample (PHN-3855), the mean \( f^{\text{Rb/Sr}} \) of the alnoite is +0.85. A melt with such Rb/Sr enrichment can be derived from a mantle source with bulk earth composition \( f^{\text{Rb/Sr}} = 0 \) by ~3% of partial melting (calculated using bulk partition coefficients of \( D_{\text{Rb}} = 0.0025, D_{\text{Sr}} = 0.033 \)) (taken from literature list as given by Jacobsen and Wasserburg, [1979a]).

The Sr and Nd isotopic compositions are very close to each other for all four alnoite samples. The \( \varepsilon_{\text{Nd}} \) (34 m.y.) is close to zero, suggesting an unfractonated source close to the bulk earth value. The \( \varepsilon_{\text{Nd}} \) (34 m.y.) is 2.5-3.7, indicating some time-integrated LREE depletion but far smaller than for typical depleted oceanic mantle. These isotopic values are similar to the \( \varepsilon_{\text{Nd}} \) (100 m.y.) = 3.8 of the basalt sample, which is believed to represent the bulk of the OJP. Despite the age difference, the sources for the alnoite and basalt can be treated as isotopically identical sources because of the short time interval. The isotopic composition of the mantle source will not change by more than one epsilon unit for Nd and 10 epsilon units for Sr within 0.1 aeon, providing \( f^{\text{Sm/Nd}} < 0.4 \) and \( f^{\text{Rb/Sr}} < 0.5 \), which is the case for all the major tectonic environments, excluding the upper continental crust [DePaolo and Wasserburg, 1979a].

Figure 4 shows the isotopic data for the Malaita alnoite and basalt in comparison with oceanic basalts values. The distinction of the Malaitan values from MORB and island arc values is obvious. It is therefore concluded that the source of the alnoite and basalt is not an old time-integrated depleted mantle such as MORB and island arc basalt sources. The high LIL and REE contents of the alnoite relative to bulk earth MORB and island arc values are therefore concluded to be isotopically identical sources because of the incompatibility elements [Boettcher et al., 1979]. The isotopic compositions, nevertheless, clearly indicate that even if the origin of the alnoite involved multistage chemical enrichment processes, the sources involved were not of an old, depleted nature.

The Malaita sample falls to the right of the mantle array close to the field of ocean islands (Figure 4). Ocean islands display a wide range of \( \varepsilon_{\text{Nd}}-\varepsilon_{\text{Sr}} \) values, and deviations from the mantle array are not rare. For instance, the Azores, Samoa, and Society islands' basalt values [Hawkesworth et al., 1979; White and Hofmann, 1982] are shifted to the right of the mantle array, thus including the Malaitan values within their range. The Malaitan isotopic characteristics are thus not unique and can be found in oceanic environments. DePaolo and Wasserburg [1979b] and Wasserburg and DePaolo [1979] presented the possibility that some oceanic islands represented plumes from an undepleted lower mantle which were partially contaminated or mixed with overlying oceanic mantle during
ascent. White and Hofmann [1982] propose a reinjection of crustal material into the mantle as an explanation of the observed isotopic values for ocean islands. The $e_{sr}$ and $e_{nd}$ values of the Malaitan samples differ conspicuously from negative $e_{sr}$ and high $e_{nd}$ values typical of old continental crust [McCulloch and Wasserburg, 1978]. There is no hint in the available data from Malaita of the isotopic signature of old continental crust. If subducted old continental crust is a component, it must be sufficiently mixed with other components so that no partial melts, with an old continental signature, are produced. This indicates that no direct melting of typical old continental material is involved either as parental material or as an incompletely homogenized contaminant.

8.2. The Xenoliths and Megacrysts

Five samples of mantle nodules found in the alnoite were analyzed (Table 1). All the samples, excluding PHN-3995, show undepleted contents of Rb, Sr, Sm, and Nd, compared with chondritic levels [Haskin et al., 1966; Mason, 1971]. The lherzolite, PHN-3995, is highly depleted in REE, having Sm and Nd 10 and 6 times lower than chondritic values, respectively. Note that of the “undepleted” samples, only PHN-3861 represents a total rock; the other samples are individual mineral phases so that no strong quantitative comparison regarding concentration levels can be made.

The $e_{sr}$ (34 m.y.) values of the five samples lie within a range of -6.0 to +1.70 and scatter around zero, similar to the alnoite and basalt samples, and none of the samples has low $e_{sr}$ typical of depleted oceanic mantle or high $e_{sr}$ typical of continental crust. The range of $e_{sr}$ (34 m.y.) for the garnet (PHN-3986A), the cpx (PHN-3968), and the gt-pyroxene (PHN-3861) is 3.1-4.2, essentially the same as the alnoite value. The $e_{nd}$ (34 m.y.) of the cpx from lherzolite PHN-3852 is +2.3, but at 100 m.y. the value $e_{nd}$ (100 m.y.) = 3.1 and is similar to the basalt value.

The isotopic similarity between the alnoite, the megacrysts, and the gt-pyroxene might be a result of either isotopic equilibration of these inclusions with the alnoite magma or represent the original mantle layer through which the alnoite passed. The megacrysts, which are homogeneous and unzoned, could not maintain their isotopic similarity at a time substantially prior to 34 m.y. because they have very different $f^{Sm/Nd}_{34}$ values. The garnet has $f^{Sm/Nd}_{34} = 1.47$, and the cpx has $f^{Sm/Nd}_{34} = -0.15$ so that they would differ by one epsilon unit if their age was 60 m.y. instead of 34 m.y. The observed $D_{cpx}^{34} = 1.6$ and $D_{nd}^{34}$ = 4.7 are consistent with values obtained experimentally [Harrison, 1981] and observed in natural lherzolites [Shimizu, 1975]. The megacrysts thus seem to be in chemical equilibrium with each other, as demonstrated by the REE distribution coefficients. The megacrysts, however, cannot be in chemical equilibrium with the very REE rich alnoite magma if we consider the distribution coefficients obtained experimentally by Harrison [1981]. The presence of a $H_2O-CO_2$ vapor phase which concentrate REE [Wendlandt and Harrison, 1979; Mysen, 1979] would facilitate equilibration. Substantial amount of calcite and phlogopite in the alnoite demonstrates the existence of volatile components and suggests the presence of a separate volatile phase in the alnoite magma. Intuitively, we would expect that these volatile components would also facilitate the equilibration of the megacrysts with the alnoite which does not appear to be the case.

In considering the possible origin of kimberlites from “ordinary” mantle, Kramers et al. [1981] have argued that the partition coefficients between kimberlitic liquids and clinopyroxenes may be very low. If this is correct, then our inference using the data of Harrison [1981] is not valid, and the megacrysts could be in chemical equilibrium with the melt. The wide range of possible distribution coefficients presently available or considered plausible do not permit us to draw a firm conclusion on this matter. However, $f^{Sm/Nd}_{34}$ values of the megacrysts range from 0 to -0.35, and therefore the isotopic composition of the megacrysts would not be the same for an age much different from that of the kimberlite intrusion. Regardless of any particular model, we infer from the isotopic similarity that the megacrysts were isotopically equilibrated at or near the time of emplacement either with the volatile-rich alnoite or were produced from partial melts of an overlying mantle layer with the same isotopic characteristics (e.g., only slightly depleted).

The lherzolite (PHN-3995) has $e_{sr}$ (34 m.y.) = +6.5. This value is compatible with the low REE content of this rock (similar to more depleted mantle) and indicates a time-integrated LREE depletion which is substantially greater than for the alnoite. There is, however, a contradiction between these inferred depletions and the low $f^{Sm/Nd}_{34}$ value of -0.33. This $f^{Sm/Nd}_{34}$ value, indicating LREE enrichment, points to a further complication in the history of the lherzolite sample and might point to a contamination (mechanical or chemical alteration) by a component with a LREE-enriched pattern such as the alnoite. If these results are due to the partial contamination with the alnoite, then the $e_{nd}$ and $e_{sr}$ value of the original lherzolite must have been substantially greater. Because of the very low Nd concentration, the $e_{nd}$ values would be most susceptible to such an effect and bring the original uncontaminated value even further toward a MORB-type source value. In summary, this sample shows that lherzolite or segment with the characteristics of a rather old, depleted mantle source was traversed by the alnoite magma. This depleted layer has affinities with old, depleted oceanic mantle and may represent a residue left from forming MORB-type magmas.

In contrast, cpx from the lherzolite PHN-3852 $e_{sr}$ (34 m.y.) (+2.3) which is slightly lower than the values found for the alnoites. If this lherzolite was partially equilibrated with the alnoite, then its original $e_{sr}$ values was even lower. It therefore points to a relatively undepleted zone, either not much different from the alnoite source or to an undepleted or enriched source.

9. DISCUSSION

The Rb-Sr and Sm-Nd data indicate that the mantle underlying the OJP is heterogeneous and layered with a large volume of relatively undepleted material (the basalt and al-
The isotopic composition of the undepleted Malaitan mantle \( \varepsilon_{\text{Nd}} \sim 0; \varepsilon_{\text{Nd}} \sim 3.5 \) is an unusual combination, incompatible with the known major mantle or crustal reservoirs \[ \text{[DePaolo and Wasserburg, 1979a]} \] (excluding oceanic islands, whose mantle source is tectonically not well defined). The \( \varepsilon_{\text{Nd}} \) value is distinct from the bulk earth values and is compatible with a LREE-depleted reservoir with \( f_{\text{Sm/Nd}} = 0.01 \) and \( Q = 0.14 \), where the angle brackets represent the average values. Assuming it represents a depleted mantle with \( f_{\text{Sm/Nd}} = 0 \) for the depletion event, which is much younger than that of the MORB source \( (1.8 \text{ a}, \text{ according to Jacobsen and Wasserburg [1979a]}) \) or to an older, much less depleted source with smaller \( f_{\text{Sm/Nd}} \). The \( \varepsilon_{\text{Nd}} \) value indicates an essentially unaltered reservoir remarkably close to the value for UR. The UR value has been obtained indirectly as the intersection of \( \varepsilon_{\text{Nd}} = 0 \) with the mantle array \[ \text{[DePaolo and Wasserburg, 1976b; O'Nions et al., 1977]} \] and is not so precisely fixed. It is therefore legitimate to consider another value for the present bulk earth \( ^{87}\text{Sr}/^{86}\text{Sr} \) composition. Assuming that the Malaita Sr isotopic composition is a result of a depletion with parameters determined by the Sm-Nd systematics, then the calculated \( ^{87}\text{Sr}/^{86}\text{Sr} \) UR in this case is \( \sim 0.7053 \). However, if we accept the bulk earth composition value as 0.7045, then there are two possible explanations for the observed isotopic composition.

1. It is an ancient mantle reservoir with a unique composition that has not been previously identified.
2. It is a mixture of known reservoirs.

The possibility that the Malaita mantle is an old reservoir implies that it has chemically evolved from primordial composition \( (\varepsilon_{\text{Nd}} = \varepsilon_{\text{Nd}} = 0) \) to yield \( \varepsilon_{\text{Nd}} \sim 3.5 \) and \( \varepsilon_{\text{Nd}} \sim 0 \). The differentiation trend had to be such that it would cause a LREE depletion without fractionating Rb/Sr. As Rb is highly fractionated in partial melting or in volatile phases, such a mechanism is very implausible.

The possibility that the source is produced as a result of mixing was examined. The end-members that were considered are depleted mantle and continental crust reservoirs. The concentration and isotopic parameters for those reservoirs have been estimated previously but are not precisely known. Jacobsen and Wasserburg \[ \text{[1979a]} \] suggested a set of Rb-Sr/Sm-Nd parameters for the present crust and depleted mantle, a set which is consistent with a primitive composition (CHUR and UR) prior to the separation of the crust from the mantle. This self-consistent (but not necessarily correct) set of parameters \( (\text{Table 2}) \) serve as a basis for the mixing calculations. The distribution of the elements among upper and lower crust was constructed \( (\text{Table 2}) \) assuming that (1) Sm-Nd levels are the same for both reservoirs \[ \text{[see Jacobsen and Wasserburg, 1979a; Table 6]} \] and, (2) \( \varepsilon_{\text{upper crust}} = 163 \) (average continental runoff, Veizer and Compston \[ \text{[1974]} \]), and (3) \( \varepsilon_{\text{lower crust}} = 2 \times \text{Rb}_{\text{lower crust}} \).

The isotopic composition of the mixture of the different crustal reservoirs and depleted mantle was calculated using the mixing equation \[ \text{[Steiger and Wasserburg, 1966]} \]

\[
\varepsilon_{\text{mix}} = x_1 \varepsilon_1 + (1 - x_1) \varepsilon_2
\]

where \( x_1 \) is the concentration of Nd or Sr and \( x_1 \) is the weight fraction of component 1 in the binary mixture \[ \text{[DePaolo and Wasserburg, 1979a]} \].

The resultant \( \varepsilon_{\text{Nd}} \) mixing curves are shown in Figure 5. The Malaitan isotopic composition \( (\varepsilon_{\text{Nd}} = 3.5; \varepsilon_{\text{Nd}} = 0) \) falls on the mixing curve of upper crust with depleted mantle at 1% upper crust. It should be noted that (1) as Sm and Nd content is assumed to be the same for both crustal reservoirs, the difference between the mixing curves depends on the Rb-Sr fractionation alone, (2) at a level of \( 1 \% \) upper crust, \( \varepsilon_{\text{Sm-Nd}} \) and \( \varepsilon_{\text{mix}} \) will not change much if \( \varepsilon_{\text{upper crust}} = \varepsilon_{\text{lower crust}} \) used in the model are somewhat different; for \( x_1 = 0.99 \), \( \varepsilon_{\text{mix}} \) will change by only 1.5 \( \varepsilon \) per 10 \( \varepsilon \) change of \( \varepsilon_{\text{upper crust}} \) and \( \varepsilon_{\text{mix}} \) will change by 0.24 \( \varepsilon \) per 1 \( \varepsilon \) change of \( \varepsilon_{\text{lower crust}} \), and (3) the shape of the mixing curves is a function of \( K \) \( (\text{K} = (\text{Sr/Nd})_{\text{mix}}/(\text{Sr/Nd})_{\text{UR}}) \). For \( K < 1 \) the curve is concave downward, and if that was the case for the lower crust, then the Malaitan result \( (\text{Figure 5}) \) would have lain close to the lower crust mixing curve and similarly for the bulk crust. However, the precise \( K \) values are not well known.

The conclusion from the mixing calculations seems to be that it is possible to obtain a reservoir with the Malaitan isotopic characteristics by mixing depleted mantle and crustal reservoir at a ratio of 100:1. By using a self-consistent set of Rb-Sr/Sm-Nd parameters for the crust and mantle and assuming \( \varepsilon_{\text{upper crust}} = \varepsilon_{\text{average continental runoff}} = 163 \) and \( \varepsilon_{\text{lower crust}} = 2 \times \text{Rb}_{\text{lower crust}} \), it appears that the upper crust is the better reservoir for the mixture. The resultant concentrations of this mixture are listed in Table 2. The \( f_{\text{Sm/Nd}} \) and \( f_{\text{Rb/Sr}} \) of the mixture are 0.07 and 0.0, respectively. Such reservoirs will change in \( \varepsilon_{\text{Nd}} \) by less than 0.2 \( \varepsilon \) over 0.1 aequ, a time span...
covering the formation of the OJP and the alnoite. By the
assumptions, $\varepsilon_{Rb}$ will not change with time.

As the volume of mantle involved in the source of the OJP is $5 \times 10^8-10^9$ km$^3$, this requires $5 \times 10^9-10^8$ km$^3$ of crustal contribution. That corresponds to a block of $10^8-10^9$ km$^2$ of continental crust of ~40 km thickness. As the alnoite and basalt were derived over a time span of ~65 m.y. and from a large volume of parental material, it appears that the source must be relatively homogeneous in order to explain the isotopic data. This imposes a difficulty with mixing the proposed components. A mechanical blend of mantle and crust is not an attractive process as we would expect a wide range of isotopic composition in the partial melting of such mixtures. There is no isotopic evidence of the result of partial melting of a crustal end-member which might result from a purely coarse scale mechanical “mixture.” A large-scale metasomatism of the mantle with fluids originated from the crustal component which could mobilize and reequilibrate the trace elements preferentially over large distances seems to be a way to obtain a rather homogeneous isotopic reservoir from such a “mixture.” The crustal component must, nevertheless, have reached a depth of at least 100 km.

The source for continental blocks in the West Pacific is somewhat problematic, as has already been pointed out by Carlson et al. [1980], Nur and Ben Avraham [1977, 1982] call for Pacifica to supply pieces of continental crust, which now appear as oceanic plateaus in the West Pacific. The results of our study are clearly incompatible with the OJP itself being a piece of old continent. However, as indicated above, such a small block of old continental crust might have been involved in producing the mantle source for the OJP if a homogenization process with large volumes of depleted mantle could be found. Another alternative is that young continental block (a ready-made blend of old crustal and young arc material) with bulk LIL element characteristics close to the average earth was subducted under the oceanic lithosphere. This could then provide an appropriate isotopic and chemical source.

DePaolo [1980, 1981] has shown that some plutonic rocks in the Sierra Nevada and Peninsular ranges of California are the product of partial melting of old continental crust mixed with younger magmas derived from the depleted mantle. These composite magmas show a wide range in $\varepsilon_{Nd}$ and $\varepsilon_{Rb}$, but several of them have isotopic compositions similar to the alnoite and basalt from Malaita. The characteristics of the two mixing mechanisms outlined above are different only in requiring mixing at near crustal levels rather than at substantial depths within the mantle. The possibility of producing a “well-mixed” smaller crustal block prior to subduction appears simpler than homogenizing crustal material with large volumes within the mantle. In any event a narrowly defined isotopic composition as a result of mixing is not readily explained.

In conclusion, we infer that the alnoites and basalts of the OJP are derived by partial melting of a lithic reservoir that is relatively undepleted ($\varepsilon_{Nd} \approx 0.1$). This reservoir may be composed of a mixture of ~99% of depleted mantle ($\varepsilon_{Nd} \approx 0.4$) with ~1% of enriched continental blocks ($\varepsilon_{Nd} \approx -0.6$). This continental block or blocks (~10$^6$ km$^3$) would then have been entrained by downward moving, depleted upper mantle to a depth of at least 100 km. Partial melting of these blocks with very extensive and large-scale metasomatic exchange with the depleted oceanic mantle component produce the observed isotopic signature. Alternatively, the reservoir might consist of deeply subducted young continental blocks with the appropriate isotopic signature. In this case the volume of the block would be relatively small. The presence of continental blocks in the ocean sector during Mesozoic time may be related to Gondwanaland fragmentation. It is possible that this mechanism of relatively large-scale homogenization of trace elements by volatile metasomatism could also provide the common characteristics of continental flood basalts ($\varepsilon_{Nd} \approx 0$) which were used by DePaolo and Wasserburg [1979] to infer the existence of an undifferentiated layer in the lower mantle. This model has recently been contested by White and Hofmann [1982], who have argued for subducted continental frag-

### Table 2. Rb-Sr and Sm-Nd Data Used for a Mixing Model for the Alnoite Mantle Source

<table>
<thead>
<tr>
<th>Reservoir</th>
<th>Sm, ppm</th>
<th>Nd, ppm</th>
<th>Rb, ppm</th>
<th>Sr, ppm</th>
<th>$\varepsilon_{Sr}$</th>
<th>$\varepsilon_{Nd}$</th>
<th>$f_{Rb/Sr}$</th>
<th>$f_{Sm/Nd}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depleted mantle</td>
<td>0.32</td>
<td>0.82</td>
<td>0.04</td>
<td>15.8</td>
<td>-27</td>
<td>10</td>
<td>-0.9</td>
<td>0.225</td>
</tr>
<tr>
<td>Bulk crust</td>
<td>5.0</td>
<td>26</td>
<td>33</td>
<td>370</td>
<td>65.5</td>
<td>-17.8</td>
<td>2.18</td>
<td>-0.4</td>
</tr>
<tr>
<td>Upper crust</td>
<td>5.0</td>
<td>26</td>
<td>50*</td>
<td>270†</td>
<td>163†</td>
<td>-17.8</td>
<td>5.4†</td>
<td>-0.4</td>
</tr>
<tr>
<td>Lower crust</td>
<td>5.0</td>
<td>26</td>
<td>24*</td>
<td>430†</td>
<td>28†</td>
<td>-17.8</td>
<td>0.9†</td>
<td>-0.4</td>
</tr>
<tr>
<td>A mixture of 1% upper crust with 99% depleted mantle†</td>
<td>0.37</td>
<td>1.1</td>
<td>0.5</td>
<td>18</td>
<td>0</td>
<td>3.3</td>
<td>0</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Data taken from Jacobsen and Wasserburg [1979a] unless otherwise indicated.

* Assuming $Rb_{upper crust} = 2 \times Rb_{lower crust}$ and $Mass_{upper crust} = 1.5 Mass_{upper crust}$.
† Result of the data and assumptions used by Jacobsen and Wasserburg [1979a, b].
‡ Average continental runoff [Veizer and Compston, 1974].
ments as a major component in the source of oceanic volcanics. The connection between the alnoite source and that of ocean islands cannot be made; however, the trend of oceanic island basalts toward enriched (or at least undepleted) parent reservoirs is the same as found by us for the sources of the OIP.

From this cursory investigation it appears that intensive studies of isotopic characteristics at localities where some knowledge of mantle "lithic stratigraphy" is known may permit a better understanding of the mantle "isotopic stratigraphy."

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REFERENCES


Basu, A. R., and M. Tatsumoto, Nd-isotopes is selected mantle…

Boetcker, A. L., J. R. O'Neil, K. E. Windsor, D. C. Stewart, and H. G. Wilshire, Metasomatism of the upper mantle and the genesis of kimberlites and alkaline basalt, in The Mantle…


Shimizu, N., Rare earth elements in garnets and clinopyroxenes from


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