We report some of the first noble gas data for in situ lower oceanic crust and shallow mantle. From a suite of gabbros and peridotites recovered from the Atlantis Bank oceanic core complex on the Southwest Indian Ridge, we measured He, Ne, Ar, Kr and Xe concentrations as well as $^{3}$He/$^{4}$He and $^{40}$Ar/$^{36}$Ar ratios, thereby documenting the noble gas content and signature of oceanic lithosphere. Except for a single ultramylonite, the gabbros have higher $^{3}$He/$^{4}$He ratios than atmospheric. Three gabbros have MORB-like bulk $^{3}$He/$^{4}$He ratios higher than 6R$_A$ despite variable helium concentrations, as much as two to three orders of magnitude lower than in MORB glasses. One of these is mylonitized, demonstrating that magmatic helium can be retained despite intense high-temperature crystal-plastic deformation in the lower crust. Of the gabbros measured, green amphibole-bearing samples show relatively high helium abundances. Peridotite noble gas concentrations measured in clinopyroxene separates are dominantly lower than gabbros. Specifically, He abundances are similar to or greater than gabbros with MORB-like $^{3}$He/$^{4}$He isotopic ratios. All the gabbros and peridotite clinopyroxenes show severely contaminated $^{40}$Ar/$^{36}$Ar values up to 1300. Magmatic $^{40}$Ar is enriched in the oxide-olivine gabbro with the highest $^{40}$Ar/$^{36}$Ar in the entire sample suite. These results suggest as an actual data that the recycling of the lower oceanic crust and shallow mantle should be considered in modeling mantle evolution at least for helium. Measured helium abundances, though lower than in basalt glasses, are greater than those in crystalline MORB. Even if entire upper crust retains primary magmatic signature, oceanic lower crust and lithospheric mantle may impact larger by recycling due to their large volumes.
1. Introduction

Noble gases have provided vital information on mantle evolution, both as a constraint on early mantle evolution and the formation of the continents, and as evidence for deep mantle plumes [Porcelli et al., 2002]. Noble gases in the upper mantle are thought to be degassed, as mid-ocean ridge basalts (MORB) have higher $^{40}\text{Ar}/^{36}\text{Ar}$ ratios compared to those of ocean island basalt (OIB), showing that the latter have retained a more primitive noble gas signature. MORB $^{3}\text{He}/^{4}\text{He}$ values of $8R_A$ lie between those of OIBs and continental materials [cf. Graham, 2002] ($R_A = \text{atmospheric } ^3\text{He}/^4\text{He} \text{ ratio of } 1.4 \times 10^{-6}$ [Ozima and Podosek, 1983]). The relatively uniform MORB helium isotopic ratio is believed to reflect the well-mixed nature of their source region, which appears to have undergone a prior melting event [Graham, 2002].

There are very limited numbers of noble gas studies for in situ deep crust or mantle rocks [e.g., Ozima and Podosek, 1983], and no published data for abyssal gabbros and peridotites. This is mainly due to sampling difficulties and a dearth of fresh, unaltered material, especially abyssal peridotite. Even in the continental environment, limited studies have been made on in situ samples [e.g., Drescher et al., 1998]. Instead, mantle-xenolith studies have been used to infer the noble gas signature of lithospheric mantle. A recent review by Dunai and Porcelli [2002] shows that the mean helium isotope ratio of subcontinental lithosphere determined from xenoliths is $6R_A$ - significantly different than that inferred for the oceanic lithosphere. However, they point out that this ratio may have been modified during transport from the mantle by contamination from the host magma as the host lavas could be derived from a different mantle source than the peridotite xenoliths.

To better constrain the noble gas budget of the mantle, we have made some of the first noble gas measurements on gabbros and peridotites sampled in situ in the oceans. Surprisingly, despite the altered state of these rocks, they retain a mantle signature, at least for helium, and contain much higher concentrations than anticipated. Thus recycled lower ocean crust and shallow mantle can be a significant contributor to the lithospheric noble gas budget.

2. Sample Locality

The Atlantis Bank core complex is the most extensively surveyed oceanic plutonic complex to date [Dick et al., 1999a; Hosford et al., 2003; Matsumoto et al., 2002]. The complex consists of a roughly 800 km$^2$ gabbro massif exposed along a major north-south lineated bathymetric high flanking the Atlantis II Transform Fault on the Southwest Indian Ridge (SWIR, Figure 1). The gabbro massif is in contact at variable depths with mantle peridotite on its western flank along the transform wall.

ODP Hole 735B is located at the top of the bank in the center of the gabbro complex. It penetrated 1500 m of intercalated coarse-grained olivine gabbro and subordinate ferrogabbro. Most of the section is undeformed or very weakly deformed with only a few percent alteration. Near the top of the Hole, however, the section is often mylonitized, and extensively altered at granulite to middle amphibolite facies conditions. Few primitive gabbros were drilled, and the average composition of the hole is too differentiated to represent in situ crystallization of primary MORB [Dick et al., 2000]. U-Pb zircon ages of $\sim$12 Ma were reported for Hole 735B by John et al. [2002].

Our samples were collected with the submersible Shinkai 6500 and the ROV Kaiko on three separate cruises to Atlantis Bank during a joint investigation by the Japan Marine Science and Technology Center (JAMSTEC) and the Woods Hole Oceanographic Institution from 1998 to 2001. Locations are summarized in Table 1 and Figure 1. Five representative gabbros were selected at $\sim$300 m depth intervals from a 4.8 km traverse that sampled 18 olivine and 11 intercalated oxide gabbros from a massive section of lower crust exposed on the transform wall at $32^\circ38'S$ between 4001 and 2528 meters water depth. Two peridotites were also selected 50 m apart from the base of a 4.2 km traverse that sampled mixed plagioclase-bearing and plagioclase-free lherzolite and harz-
burgite from a massive mantle section exposed on the transform wall 3–4 km west of the main gabbro massif at 32°50’S between 5000 to 3000 m water depth.

3. Sample Description

3.1. Gabbros

Two oxide gabbros (6K466R10 and 6K467R2) and three olivine gabbros (6K466R5 and 6K467R7, R10) were selected for analysis (Table 2). These rock names follow those of ODP Leg 176 [Dick et al., 1999b].

The oxide gabbros include an amphibolitized oxide gabbro ultramylonite (6K466R10) and an oxide-olivine gabbro mylonite (6K467R2). They consist largely of plagioclase and clinopyroxene with 3–15% ilmenite and titanomagnetite, though sample 6K467R2 contains a small amount of olivine now pseudomorphed by clay, and 2% brown
amphibole. Like many Atlantis Bank ferrogabbros, both exhibit strong crystal-plastic deformation. Plagioclase is 90–95% and clinopyroxene is 50–65% dynamically recrystallized. Alteration is mostly in the amphibolite facies, and is light in the oxide-olivine gabbro, but very heavy in the oxide gabbro. In the latter, clinopyroxene is entirely replaced by green amphibole and plagioclase is often turbid.

[10] The olivine gabbros contain ~60% plagioclase, 22–30% clinopyroxene, 10–18% olivine, <1% brown amphibole and <1% iron oxides. Olivine gabbro 6K466R5 shows no crystal-plastic deformation, 6K467R7 weak deformation and 6K467R10 moderate deformation with a good foliation. All three are quite fresh with olivine unaltered to 2% replaced by low temperature clays and chlorite, and plagioclase and pyroxene only 1–5% replaced by green amphibole and/or chlorite. Owing to the overall freshness of the coarser gabbros and the fine grain size of the mylonites, we analyzed only whole rock gabbros for this study.

3.2. Peridotites

[11] The peridotites include a granular websterite (10K173R19) and a protogranular lherzolite (10K173R23, Table 2). These textures are characteristic of the earliest high-temperature fabrics in abyssal peridotites, often associated with mantle melting beneath an ocean ridge. Textural elements, associated with lower temperature crystal-plastic deformation or cataclasis are minor. Plagioclase, indicative of crystallized trapped melt [Dick, 1989], was not observed in either peridotite. The websterite likely represents a mantle vein produced by melt impregnation at pressures above the plagioclase stability field, while the lherzolite is probably a simple residue of low degree mantle melting.

[12] The extent of alteration of the peridotites is substantial. Most olivine is replaced by serpentine, principally lizardite, though a significant proportion of primary enstatite and clinopyroxene has survived. In the lherzolite, 85% of the olivine is weathered to brown-colored clays and other secondary minerals. Trace sulfides are also observed, while pyroxene has few serpentine veins but abundant inclusions too small to identify. In the websterite abundant fibrous serpentine veins crosscut pyroxene, and carbonate and zeolite veins are also present. Therefore to avoid alteration effects, whole rock analyses were not made, and fresh clinopyroxene was separated from both samples and used in this study.

4. Analytical Procedures

[13] Gabbro fragments of ~1 g were cut and ultrasonically cleaned for 5 min with 1N-HNO₃ and rinsed with pure water, acetone and ethanol. To make the mineral separates the peridotites were crushed to less than one millimeter in size, sieved, and relatively clear and less-fractured clinopyroxene grains were hand picked. These grains were then gently etched at 70°C for 30 min with 2N-HNO₃. After etching, discolored and fragmented clinopyroxene grains were removed by hand picking using a binocular microscope, reducing the volume of the lherzolite separate by 65% and the websterite separate by 25%. The remaining clear clinopyrox-
Table 2. Petrography of Analyzed Gabbros and Peridotites

<table>
<thead>
<tr>
<th>Sample Number</th>
<th>Lithology</th>
<th>Gr. Size, mm</th>
<th>Gr. Size, mm</th>
<th>Primary Igneous Mineralogy</th>
<th>Gr. Size, mm</th>
<th>Primary Igneous Mineralogy</th>
<th>Gr. Size, mm</th>
<th>Primary Igneous Mineralogy</th>
<th>Gr. Size, mm</th>
<th>Primary Igneous Mineralogy</th>
<th>Crystal-Plastic Recrystallization, %</th>
<th>Alteration, %</th>
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<td></td>
<td></td>
<td></td>
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<tr>
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<td>60</td>
<td>30</td>
<td>10</td>
<td>&lt;1</td>
<td>Tr</td>
<td>90</td>
<td>80</td>
<td>95</td>
<td>3.5</td>
<td>1 Chl, Cl</td>
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<tr>
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<td>22</td>
<td>-</td>
<td>2</td>
<td>2</td>
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<td>90</td>
<td>50</td>
<td>100</td>
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<td>29</td>
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</tr>
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<td>- 1 Tr</td>
<td>-</td>
<td>-</td>
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<td>-</td>
</tr>
</tbody>
</table>

Mineral abbreviations: Plag, plagioclase; Cpx, clinopyroxene; Opx, orthopyroxene; Ol, olivine; Br Am, brown amphibole; Tmt, titanomagnetite; Ct, chromite; Sulf, sulfide. Deformation grade runs from 0 to 5, with 0 undeformed to 5 ultramylonite. Alteration minerals: grAm, green amphibole; Chl, chloride; Cl, clay; Serp, serpentine.

b Sample contains a few thin chlorite and smectite veins and about 2% clay. Well foliated.
c Thin late magmatic vein of recrystallized gabbro impregnated with brown amphibole.
d A few zircons present.
e Contains dark green amphibole veins perpendicular to the well developed foliation. Coarse to medium pyroxene porphyroclasts abundant. Locally has 1–2% iron-oxide stained thin clay filled cracks. Trace of apatite. Plagioclase is clear, but locally very turbid filled with 5–20% small micro-inclusions. Could be fine clay or chlorite.
f Sample contains some thin brown clay stained cracks and veins.
g Some small irregular interstitial clinopyroxenes that look like they were melt-impregnated.
h Contains cross fiber serpentine veins up to 2 mm, a 1 mm carbonate vein with mineral fragments, and a 1 mm zeolite vein running along side the carbonate vein.
enes were then ultrasonically cleaned with distilled water, acetone and ethanol. Prior to gas extraction, all samples were baked at 150°C in vacuum.

Noble gas analyses were performed with a VG-5400 noble gas mass spectrometer installed in Earthquake Research Institute of the University of Tokyo. Stepwise heating was used with two temperature steps for gabbros and three for clinopyroxene separates to extract the gases (Table 3). Details of analysis are described in Hanyu et al. [1999]. Major elements as well as Ba, Sr, Y, Sc, Zr, Be and V were obtained by lithium metaborate/tetraborate fusion ICP analysis, while REE and 29 additional trace elements analyses were obtained by ICP-MS at Activation Laboratories of Ancaster, Canada. Details are described in Appendix.

5. Helium

5.1. MORB-type 3He/4He

Concentration weighted ‘total’ He compositions of clinopyroxenes separated from abyssal peridotites fall within the commonly held MORB-type benchmark of 8 ± 1 Ra. In particular, lherzolite 10K173R23 clinopyroxene shows uniform 3He/4He within analytical uncertainty (ca.
Olivine gabbro 6K467R7 also shows MORB-type $^{3}$He/$^{4}$He for both 900°C and 1700°C fractions (8.1 and 7.9 RA). Except for the extremely altered oxide gabbro ultramylonite (6K466R10), the remaining three gabbros had MORB-like $^{3}$He/$^{4}$He ratios of almost 8 RA or above in the 1700°C fraction, regardless of helium concentration (Figure 2). Hatched area shows the typical ranges of MORB glasses. As $^{3}$He in crystalline MORBs frequently are under the detection limit, an arrow just above the upper horizontal axis shows their range of $^{4}$He amount. The filled square represents helium composition of seawater.

5.2. Radiogenic $^{3}$He/$^{4}$He in Oxide Gabbro Ultramylonite 6K466R10

Oxide gabbro ultramylonite 6K466R10 is unique in having a very low bulk $^{3}$He/$^{4}$He of 0.22 RA. The 1700°C gas fraction is atmospheric ($^{3}$He/$^{4}$He = 1 RA), with a $^{4}$He concentration of $4 \times 10^{-9}$ cm$^{3}$STP/g. This apparent atmospheric helium ratio, however, does not likely represent seawater helium incorporated into metamorphic amphibole. The $^{4}$He/$^{20}$Ne ratio of the 1700°C gas fraction reaches 100, while the ratio of seawater is only 0.4. Thus this ‘atmospheric’ $^{3}$He/$^{4}$He ratio seems to be generated by a chance mixture of mantle He and ingrowth of radiogenic He. The 900°C fraction has a much more radiogenic ($^{4}$He-rich) isotopic composition of 0.165 RA, supporting this interpretation. Normal in situ decay of U and Th gives pure $^{4}$He, which, combined with associated very small $^{3}$He contributions by following nuclear reactions, gives values of ~0.02 RA [Graham, 2002] for helium isotopic ingrowth. The Th and U contents of this sample (0.17 and 0.23 ppm respectively) would produce the observed accumulation of $^{4}$He ($6.7 \times 10^{-8}$ cm$^{3}$ STP/g) in 2 Ma. If all the $^{4}$He produced by decay of the observed Th and U in the sample were preserved over the ~11 Ma age of the sample locality, the accumulation of $^{4}$He would have reached $3.5 \times 10^{-7}$ cm$^{3}$ STP/g. An equivalent accumulation of $^{40}$Ar, as discussed later, has also been found, and can be similarly explained by decay of $^{40}$K.

5.3. Helium Abundances

Although the total helium abundances in all our samples are lower than those of MORB glasses by two to three orders of magnitude, these abundances are still significantly higher than those of crystalline MORBs (Figure 2). Bulk helium in the lherzolite clinopyroxene is also 5 times higher than in the websterite clinopyroxene. Although the degree of alteration is similar, clinopyroxene in the websterite is dominantly recrystallized and contains abundant fibrous serpentine veins. These textural differences may greatly impact the helium content of the respective clinopyroxene, though one must also consider the petrogenetic origin of host peridotite as well.

In either case, however, clinopyroxene helium abundances are much higher than those anticipated from partitioning of helium with MORB magmas as represented by glassy MORBs with typical helium contents of $10^{-5}$ cm$^{3}$ STP/g [e.g., Honda...
and Patterson, 1999]. Even in natural samples, the presence of inclusions does not guarantee that noble gas abundance is completely controlled by partitioning; helium is believed to be a highly incompatible element [e.g., Graham, 2002]. This view is supported by both laboratory high-pressure experiments and by comparisons between coexisting glass and phenocryst compositions in natural samples. For example, Brooker et al. [2003] reports extremely low partition coefficients between clinopyroxenes and coexisting glass for Ne with the value lower than $10^{-3}$. Their data implies a helium mineral/melt partition coefficient close to $10^{-4}$, under inclusion-free condition. However, glasses from the unique popping rocks dredged on the MAR contain great amounts of helium up to almost $10^{-4} \text{ cm}^3 \text{ STP/g}$ [Moreira et al., 1998]. While trace element signatures are enriched relative to N-MORB, these He concentrations for undegassed MORB melts (‘popping rock’) would suggest that most MORB magmas lose 90% of their helium prior to glass solidification as an upper limit [Sarda and Moreira, 2002]. Even if this is the case, the partition coefficient between MORB melt and residual clinopyroxene will be $10^{-3}$. The required value, however, would still be higher. If, on the other hand, their model provides a lower limit for He concentrations in primary MORB magmas, the He abundance in the peridotite clinopyroxenes can be explained by equilibrium with MORB magma prior to losing their primordial volatiles.

Our gabbros contain the equivalent or smaller amounts of helium than the peridotite pyroxenes. The most abundant MORB-type helium was found in a lightly deformed and weakly altered olivine gabbro, 6K467R7. The oxide-olivine gabbro mylonite 6K467R2, however, retains significant amounts of MORB-like helium despite the degree of deformation. The Atlantis Bank oxidized-olivine gabbros crystallized from highly differentiated iron-titanium-rich melts (∼70% plus crystallization, Natland and Dick [2001]). Thus the high retained-helium abundance in this sample likely relates to a high initial content due to enrichment of the melt in volatiles during differentiation.

Within the gabbros, the green amphibole-bearing samples (6K466R10, 6K467R2 and R7) show significantly higher helium contents regardless of isotopic composition. This suggests that amphiboles could be a principal host for noble gases, at least for helium. Green amphibole may form either deuterically or due to seawater alteration, and thus could host either magmatic or seawater volatiles. The remaining olivine gabbros are characterized by low helium concentration and low $^3\text{He}/^4\text{He}$ in the lower temperature fraction. One (6K467R10) is highly deformed while the other (6K466R5) is undeformed, and both are relatively fresh. They do have small amounts of brown magmatic amphibole, but neither has green amphibole, containing small amounts of chlorite instead. The low overall helium concentration, then, could relate to the absence of green amphibole, while magmatic helium retained in brown amphibole could account for greater than atmospheric $^3\text{He}/^4\text{He}$.

Although our sample set is small, and a larger data set would be more definitive, the helium abundances of gabbros in the present study can be interpreted as a result of equilibration with MORB magma during their precipitation and solidification, modified to some extent by later alteration. Evidence for a correlation to degree of deformation is weak at best, though relative amphibole abundances suggest the possibility that this phase may be a host for helium in the gabbros.

6. Other Noble Gases

6.1. $^{40}\text{Ar}/^{36}\text{Ar}$

$^{40}\text{Ar}/^{36}\text{Ar}$ ratios are low compared to typical MORB and range from 1290 down to 305 near the atmospheric value of 295.5, indicating only a little non-atmospheric $^{40}\text{Ar}$ (Figure 3). While many MORBs have $^{40}\text{Ar}/^{36}\text{Ar}$ ratios in the range of a few hundred to a few thousand [e.g., Sarda et al., 2000; Graham, 2002], pristine MORB magmas are widely regarded as those with $^{40}\text{Ar}/^{36}\text{Ar}$ values exceeding 10,000. As all these samples have a high $^{36}\text{Ar}$ concentration, they appear to be severely contaminated by an atmospheric component.

Non-atmospheric $^{40}\text{Ar}$ ($^{40}\text{Ar}^*$), calculated by subtraction of atmospheric $^{40}\text{Ar}$ (295.5 times of $^{36}\text{Ar}$) from total $^{40}\text{Ar}$ is highly variable in the five
gabbros. Measured K₂O values in the gabbros range from 0.08 to 0.21 wt.% (Table 4). Except for olivine gabbro 6K467R10 with the lowest ⁴⁰Ar*, both the remaining olivine gabbros show significantly higher ⁴⁰Ar* than that expected for in situ accumulation by decay of ⁴⁰K in 12 Ma. Oxide-gabbro ultramylonite 6K466R10 has an amount of ⁴⁰Ar* roughly equivalent to that expected for in situ accumulation. Also lacking excess ⁴He, this sample possibly lost its magmatic noble gases during alteration.

Oxide-olivine gabbro 6K467R2 has a unique signature: while it contains the highest K₂O content (0.21 wt.%), its ⁴⁰Ar* content reaches almost 1 × 10⁻⁶ cm³ STP/g—11 times higher than the possible accumulation of ⁴⁰Ar* over 12 Ma. This gabbro also shows the highest ⁴⁰Ar/³⁶Ar, and has the second highest helium content measured. This amount of ⁴⁰Ar* is equivalent to the excess argon found in zero age MORB glass, and the elevated ⁴⁰Ar/³⁶Ar in sample 6K467R2 likely comes from its parental magma.

[25] K₂O in abyssal peridotites is generally below detection limits for EPMA, however, assuming a K₂O content of 30 ppm up to 1.4 × 10⁻⁹ cm³ STP/g of ⁴⁰Ar* could accumulate in the pyroxene over 12 Ma. This is lower by an order of magnitude than observed ⁴⁰Ar* in the pyroxene separates. Therefore the large observed differences in ⁴He/⁴⁰Ar* in the lherzolite and websterite clinopyroxene separates could be primary mantle or magmatic signatures reflecting the differences in their petrogeneses.

6.2. Elemental Abundances and Atmospheric Neon, Krypton and Xenon

[26] The noble gas elemental abundance patterns, with the exception of He and Ne, have similar shapes to those of glassy MORBs, and equivalent or higher heavier noble gas contents (Figure 4). Helium has concentrations lying between crystalline and glassy MORB, departing from both patterns. Neon shows a variable depletion relative to the ‘smooth’ MORB pattern, ranging from almost none for oxide-olivine gabbro mylonite 6K467R2 to one order of magnitude in olivine gabbro protot-

![Figure 3.](image)

**Figure 3.** ⁴⁰Ar/³⁶Ar versus ⁴He/³He diagram. Symbols of samples are the same as Figure 1. The bulk values of stepwise heating are shown. The high end of MORB ⁴⁰Ar/³⁶Ar values is extended beyond the diagram, as its maximum is shown to be 40,000 [Burnard et al., 1997]. The large filled square represents the value for seawater.

<table>
<thead>
<tr>
<th>Sample</th>
<th>U, ppm</th>
<th>Th, ppm</th>
<th>In Situ ⁴He (10⁻⁸ cm³ STP/g)</th>
<th>Measured ⁴He (10⁻⁸ cm³ STP/g)</th>
<th>In Situ ⁴⁰Ar (10⁻⁸ cm³ STP/g)</th>
<th>Measured ⁴⁰Ar* (10⁻⁸ cm³ STP/g)</th>
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* N.A., not analyzed.
mylonite 6K467R10. Xenon contents are as high as the most xenon-rich glassy MORBs, while the abundances of $^{36}\text{Ar}$ and $^{84}\text{Kr}$ are lower in the olivine gabbros than the highly evolved oxide gabbros. The latter have higher absolute abundances for most of the noble gases, and significantly less depletion in $^{20}\text{Ne}$ relative to MORB glasses. The clinopyroxene separates have slightly lower neon and argon abundances than those of the lowest gabbro value (Figure 4b), while their $^{132}\text{Xe}$ contents are virtually identical. Lacking of comprehensive information on the partitioning of noble gases between pyroxene and olivine, it is difficult to interpret differences in the absolute elemental abundances between the gabbro whole rock analyses and the peridotite mineral separates.

[28] Although not shown in Table 3, the individual isotopic compositions measured for neon, krypton and xenon in our samples are all atmospheric as in MORB with low $^{40}\text{Ar}/^{36}\text{Ar}$. Considering this, the departures in the noble gas patterns in Figure 4 from seawater must relate to contamination or hydrothermal alteration processes.

7. Discussion

7.1. Noble Gas Budgets in Lithospheric Mantle

[29] Our results show that the contribution of gabbroic lower crust to the lithospheric helium inventory is equal to or greater than that of pillow lavas and diabase, while the shallow mantle may potentially provide a reservoir with an even larger amount of helium (Table 5). This is because seafloor basalts usually lose most of their magmatic helium except for a thin quenched glassy rind [Dymond and Hogan, 1973; Kumagai and Kaneoka, 1998]. Typically, the rinds are a few millimeters or less in thickness and likely comprise only 1% of a typical lava flow. Given that roughly half the earth’s surface is normal oceanic crust ($2.6 \times 10^8 \text{ km}^2$), these rocks may contribute significantly to the lithospheric helium budget. Presuming a uniform 5 km thick gabbroic lower crust gives a volume exceeding $1.3 \times 10^9 \text{ km}^3$, and a mass of $\sim 4 \times 10^{24} \text{ g}$. Our measured gabbro He abundances, exceed $1 \times 10^{-7} \text{ cm}^3 \text{ STP/g}$. If this is taken as an average estimate, as much as $5 \times 10^{17} \text{ cm}^3 \text{ STP}$ of MORB-type helium could be contained in the oceanic lower crust, equivalent to 2.5% of atmospheric helium.
The peridotites analyzed provide a first look at the possible contribution of the oceanic upper mantle to the lithospheric noble gas budget. Both were emplaced to the seafloor at the rift valley wall of the paleo-SWIR on the western flank of the core complex close to 12 Ma. Trace element abundances in lherzolite 10K173R23, specifically depleted LREE patterns (Y. Nishio, personal communication, 2003), indicate generation from low to moderate degrees of mantle melting similar to those reported by Johnson and Dick [1992]. Websterite 10K173R19 likely represents late-stage impregnation by ambient melt near the base of the lithosphere. Because they are the only available helium abundance of clinopyroxene in abyssal peridotite, we assume that they could represent possible upper and lower bounds for helium contents for the shallow oceanic mantle far from a hot spot. They span nearly the entire range of helium contents found in the gabbros, and both exhibit high $^3$He/$^4$He ratios essentially identical to MORB. Given that MORB is thought to represent $\sim$8–20% mantle melting [Klein and Langmuir, 1987], by inference the residual mantle is 5 to 13 times the volume of the entire crust. Thus the shallow oceanic mantle is potentially a major component of the lithospheric helium budget, and may contain a volume of helium equal to half that found in the earth’s atmosphere. However, undegassed upper mantle should have as $10^3$ times much helium as lithospheric and atmospheric inventories (Table 5).

Unlike helium, with the exception of xenon, our measurements do not suggest a major reservoir for the other noble gases in the shallow earth, as their abundances are too low. For example, the undegassed MORB source has been estimated to contain $1 \times 10^{-10}$ cm$^3$ STP/g of $^{36}$Ar [Porcelli and Ballentine, 2002], which seems to be possible maxima results in $1 \times 10^{-9}$ cm$^3$ STP/g of $^{36}$Ar in glassy MORB. If this is the case, $1 \times 10^{17}$ cm$^3$ STP of $^{36}$Ar resides in the MORB source region at most. This value is three orders of magnitude lower than that of atmospheric $^{36}$Ar.

Our peridotites contain on the order of $10^{-12}$ cm$^3$ STP/g of $^{132}$Xe with an atmospheric isotopic composition. Using this value, lithospheric oceanic mantle could contain up to $2.5 \times 10^{14}$ cm$^3$ STP of $^{132}$Xe, equivalent to 0.25% of the total atmospheric budget. If the oceanic lithosphere is recycled back into the mantle every $\sim$200 Ma, $5 \times 10^{15}$ cm$^3$ STP of $^{132}$Xe, roughly 5% of total atmospheric inventory, could have been recycled back into the earth. This impact is similar to that by the upper crust including sediments.

### 7.2. $^3$He/$^4$He of the Lower Temperature Fractions

With the exception of one olivine gabbro with abundant helium, and the lherzolite clinopyroxene separate, our samples show lower $^3$He/$^4$He in the 900°C and 700°C low-temperature fractions respectively. Generally, samples with smaller helium contents have lower $^3$He/$^4$He in the low-temperature fraction. These values spread downward from MORB-like ratios toward the value of the 1700°C oxide gabbro ultramylonite fraction (Figure 2). This is interpreted as the result of several likely effects including loss of magmatic helium and in situ radiogenic $^4$He ingrowth.

Table 4 shows the possible in situ accumulation of $^4$He based on the measured U and Th
concentrations in our gabbro samples. Although some uncertainties remain as half of U and Th data are reported as upper limit, possible radiogenic $^4\text{He}$ ingrowth in 12Ma is enough to decrease their $^3\text{He}/^4\text{He}$ from that of MORB to the observed values. The helium composition of clinopyroxene in websterite 10K173R19 can be similarly explained. In the 700°C fraction it has $^3\text{He}/^4\text{He}$ 4.7 R$_A$ and only $3 \times 10^{-9}$ cm$^3$ STP/g of $^4\text{He}$. This decrease of the $^3\text{He}/^4\text{He}$ ratio requires a $^4\text{He}$ accumulation of only $1.3 \times 10^{-9}$ cm$^3$ STP/g over 12 Ma. Assuming that its Th and U concentrations are 10 ppb and 3 ppb with an acceptable mantle Th/U ratio (typically 3.5; e.g., Ozima and Podosek [1983]), possible in situ production of $^4\text{He}$ over 12 Ma reaches $7.8 \times 10^{-9}$ cm$^3$ STP/g. Thus after at least partial escape of primordial helium, in situ ingrowth of radiogenic $^4\text{He}$ could explain the all observed $^3\text{He}/^4\text{He}$ ratio.

7.3. Atmospheric Contamination Processes

[35] The $^{36}\text{Ar}^{22}\text{Ne}$-$^{132}\text{Xe}^{22}\text{Ne}$ diagram may be a useful tool for discriminating among possible contaminants [Kumagai and Kaneoka, 2003] as three potential contaminants with atmospheric isotope ratios are clearly separated in the diagram: air, seawater, and seawater-equilibrated tholeiitic melt (Figure 5). The latter is a theoretical component calculated using noble gas solubilities in a tholeiitic magma [Carroll and Webster, 1994, Figure 13] equilibrated with seawater using Henry’s law [Ozima and Podosek, 1983, Table 7.6]. Although this component has an atmospheric isotopic composition it has elemental abundances quite different than either the atmosphere or seawater. In addition, simple two component mixing makes a linear trend in this diagram.

[36] All our gabbro and peridotite data follow a trend between the seawater-equilibrated tholeiitic magma component and another component represented by the lherzolite clinopyroxene separate. Therefore we conclude that the noble gas components of our samples were derived from mixing between these end-members. Three of the gabbros lie to the right of the mixing line offset toward the seawater component, and therefore also show some effects that can be ascribed to seafloor weathering. The low $^{40}\text{Ar}/^{36}\text{Ar}$ of the lherzolite clinopyroxene separate also supports some seawater contamination.

[37] The origin of the ‘exotic’ Xe enriched component has several possible explanations. This could be a mantle component, though the atmospheric isotope signatures of Ne and Xe do not support this. It could represent a severe fractionation of atmospheric noble gases occurring during alteration, but this component has also been seen in fresh hot spot lavas [e.g., Patterson et al., 1990; Harrison et al., 1999; Althaus et al., 2003]. Accordingly the origin of this component remains undetermined.

[38] The direct equilibration of seawater and a MORB magma seems mechanically improbable. MORB melts can digest altered gabbro in the lower crust, however, this process seems to be most robust at fast spreading not slow-spreading ridges [Michael and Schilling, 1989]. A more likely explanation for this component in our gabbros is simply high-temperature hydrothermal alteration.
by seawater circulating into the lower crust beneath the ridge axis. Our principle reservoir for helium and presumably the other noble gases appears to be green amphibole, which dominantly forms at middle amphibolite facies or higher in the Hole 735B gabbros [Stakes, 1991; Stakes et al., 1991].

7.4. $^{4}\text{He}/^{40}\text{Ar}^{*}$

[39] $^{4}\text{He}/^{40}\text{Ar}^{*}$ ratios are frequently utilized as a tracer of magmatic components. Possible accumulation of in situ radiogenic $^{4}\text{He}$ and/or $^{40}\text{Ar}$ during 12 Ma, however, obscures the original $^{4}\text{He}/^{40}\text{Ar}^{*}$ ratios in most of the gabbros analyzed (Table 4). One exception is olivine gabbro 6K467R7, which contains the most abundant MORB-like helium in our suite. The $^{4}\text{He}/^{40}\text{Ar}^{*}$ ratio of this gabbro is 2.01, which is almost equivalent to the integrated production ratio of $^{4}\text{He}/^{40}\text{Ar}$ for 4.5 Ga of 2.1. In addition, this value is very close to the popping rock value of 1.6 [Moreira et al., 1998]. Even taking into account possible in situ accumulation of $^{40}\text{Ar}$ for 12 Ma, the $^{4}\text{He}/^{40}\text{Ar}^{*}$ value does not exceed 4. This is still lower than the present-day production ratio of $^{4}\text{He}/^{40}\text{Ar}$, 5.5.

[40] $^{4}\text{He}$ and $^{40}\text{Ar}$ ingrowth in the peridotites is negligible. Websterite 10K173R19 clinopyroxene shows almost identical $^{4}\text{He}/^{40}\text{Ar}^{*}$ with that of popping rock. This is consistent with an interpretation that it was produced by melt impregnation containing mantle noble gases. The high $^{4}\text{He}/^{40}\text{Ar}^{*}$ value of lherzolite (8.5) is difficult to explain at this point without more data.

8. Conclusions

[41] Oceanic gabbros and peridotites can preserve the primary magmatic helium signature of the MORB source, even within highly deformed and/or weathered rocks. Their maximum helium abundances exceed $1 \times 10^{-7}$ cm$^3$ STP/g. Helium concentrations up to $10^{-6}$ cm$^3$ STP/g with MORB-like $^{3}\text{He}/^{4}\text{He}$ have also been estimated for oceanic lithosphere from ultramafic xenoliths [e.g., Moreira and Kurz, 2001]. Although those values are lower than those of MORB glasses by two to three orders of magnitude, these abundances are still significantly higher than those of crystalline

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Detection limit 2 2 2 2 1 5

Fusion ICP measurements with a flux of lithium metaborate/tetraborate were performed by Activation Laboratories Ltd., Ancaster, Canada. Report number 27067. Dashes indicate under detection limits.
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*ICP-MS measurements were performed by Activation Laboratories Ltd., Ancaster, Canada: Report No 27067. Dashes indicate under detection limit.

*b Given in ppm.
MORBs. Considering the large volumes of the lower crust and residual mantle, then, they are both important reservoirs of helium with the lower crust contributing equally or more than the upper crust to the helium budget. Both, however, are likely minor contributors (10% or less) compared to the residual mantle to the lithospheric helium budget. Given the assumption that the helium is highly incompatible, this is a surprising result. Recycling the oceanic lithosphere at subduction zones then should be considered in modeling mantle evolution.

[42] The noble gas abundances in our gabbros and peridotites are dominated by a component reflecting high temperature alteration and an exotic atmospheric component, though some of samples partly preserve a mantle derived noble gas signature indicated by $^{4}\text{He}/^{40}\text{Ar}^*$ ratios.

Appendix A. Chemical Analyses of Gabbros

[43] Chemical analyses of gabbros are shown in Tables A1 and A2.

Acknowledgments

[44] We would like to express our appreciation to Captain Ishida and the crews of the R/V Kairei and Yokosuka, operation teams of Shinkai 6500 and Kaiko ROV for their assistance throughout cruises. On-board and shore-based scientific teams are also appreciated owing to their contribution. We also thank to Philippe Sarda and Samuel Niedermann for their constructive reviews for early version of manuscript, and to Mark Kurz and Wolfgang Bach for their constructive discussions and useful comments. Yoshio Nishio kindly provided his unpublished data. This study was supported by the following grants: the Grant-in-Aid for Scientific Research by the Ministry of Education, Culture, Sports, Science and Technology, Japan, (14540457), the Earthquake Research Institute cooperative research program (2001-G-06 and 2002-G-14) and the Grant C32 for Deep-Sea Research and Fellowship for Research Abroad from JAMSTEC to H.K., and NSF Grant OCE-9907630 to the Woods Hole Oceanographic Institution (WHOI). The expeditions were conducted as part of the Ocean Floor Geodynamics Program in JAMSTEC under MOU with WHOI.

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