The heavy noble gas composition of the depleted MORB mantle (DMM) and its implications for the preservation of heterogeneities in the mantle

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To characterize the heavy noble gas composition of MORBs we present new He, Ne, Ar, and Xe abundances and isotopic compositions from the equatorial Mid-Atlantic Ridge. Both depleted MORBs nominally devoid of plume influence and more enriched MORBs thought to represent the influence of a HIMU mantle plume are present in close geographical proximity in this region. Ne–Ar–Xe isotopic compositions in individual step-crushes are correlated, which, along with significant radiogenic excesses, allows correction for shallow-level air contamination. Based on the relationship between the noble gases and the lithophile isotopes (Sr, Nd and Pb), the depleted MORB mantle has a 21Ne/22Ne between 0.0617 and 0.0646, 40Ar/36Ar ratio of 41,500 ± 7000 and 129Xe/130Xe ratio of 7.77 ± 0.06. On the other hand, the HIMU-type MORBs are characterized by far less radiogenic Ne, Ar and Xe isotopic compositions with mantle source 21Ne/22Ne between 0.0544 and 0.0610, and 40Ar/36Ar and 129Xe/130Xe ratios of 18,100 ± 600 and 7.21 ± 0.06, respectively. The observation of less nucleogenic 21Ne/22Ne in HIMU-type MORBs is similar to observations from HIMU ocean islands and requires the HIMU plume to be comprised of both recycled and primitive material. Within the depleted MORBs we observe He and Ne to be negatively correlated. The observation suggests that along the equatorial Atlantic the most depleted MORBs are related to normal MORBs through the addition of a small proportion of a HIMU plume component.

Our new Xe isotopic measurements demonstrate distinct 129Xe/136Xe ratios in the mantle sources of depleted MORBs, HIMU-type MORBs and the Iceland plume. While substantial injection of atmospheric Xe into these mantle sources is implied, the differences in Xe isotopic composition cannot result solely from recycling of air. Rather, they require that mantle plumes sample a reservoir less degassed than the depleted MORB mantle. This conclusion is consistent with a higher proportion of Pu- to U-derived fission Xe in Iceland and HIMU-type MORBs compared to the depleted MORBs. Overall, the Xe isotopic compositions imply that mantle plumes tap a reservoir that separated from the MORB source within the first 100 million years of Earth’s history and that the two reservoirs have had limited direct mixing since then.

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1. Introduction

Noble gas isotope ratios in mantle-derived rocks are powerful tools for understanding the structure and evolution of the Earth’s mantle, the acquisition of terrestrial volatiles, and the exchange of material between the solid and fluid Earth (Allegre et al., 1987; Coltice et al., 2011; Gonnermann and Mukhopadhyay, 2009; Holland et al., 2009; Honda and McDougall, 1998; Marty, 1989; Marty and Tolstikhin, 1998; Ozima and Igarashi, 2000; Pepin, 1991; Porcelli and Wasserburg, 1995; Sarda et al., 1999; Trieloff and Kunz, 2005). Their low abundances, unreactive nature, and preponderance of radiogenic and nonradiogenic (primordial) isotopes allow them to be ideal tracers of the degassing history of mantle reservoirs and the long-term interaction between these reservoirs. This is because small amounts of radiogenic ingrowth of noble gas daughter nuclides or admixtures of different material can dramatically alter noble gas isotope ratios, often in excess of other isotope systems by orders of magnitude.

Low ratios of the radiogenic to non-radiogenic noble gas nuclides (e.g. 21Ne/22Ne, 40Ar/36Ar, 129Xe/130Xe) in ocean island basalts (OIB) compared to mid-ocean ridge basalts (MORB) are often used as evidence for a less degassed deep mantle and a more degassed, more processed upper mantle (e.g. Allegre et al., 1983, 1996; Gonnermann and Mukhopadhyay, 2009; Kurz et al., 1982; Porcelli and Wasserburg, 1995). The low ratios of 40Ar/36Ar and 129Xe/130Xe in OIBs have also been attributed to the recycling...
of atmospheric noble gases (e.g. Holland and Ballentine, 2006; Trieloff and Kunz, 2005). However, Mukhopadhyay (2012) showed that the $^{40}$Ar/$^{36}$Ar and $^{129}$Xe/$^{134}$Xe in the melting sources of the Iceland plume and the gas-rich MORB (“popping rock”) from the north Atlantic (Kunz et al., 1998; Moreira et al., 1998) cannot be related solely through recycling of atmospheric noble gases; differences in the degree of mantle degassing are required to explain the observed Ar and Xe isotopic compositions.

The extent to which the noble gas composition of popping rock is representative of the depleted MORB mantle, however, is not clear. This is because the sample is highly enriched in trace elements (La/Sm$_N$ = 2.1; Dosso et al., 1991, 1993) compared to the normal depleted MORBs (N-MORBs; La/Sm$_N$ ≤ 1) that erupt along > 70% of the mid-ocean ridge system. Furthermore, recent work suggests that atmospheric noble gases are subducted into the mantle (Holland and Ballentine, 2006; Sumino et al., 2010; Kendrick et al., 2011) although the extent to which recycling affects the mantle noble gas budget is debated (Moreira and Raquin, 2007). Nonetheless, if the recycled component is distributed heterogeneously, large variations in MORB source Ar and Xe isotopic compositions would be expected. In this regard, Sarda et al. (1999) showed that maximum measured $^{40}$Ar/$^{36}$Ar ratios in Atlantic MORBs were negatively correlated with $^{206}$Pb/$^{204}$Pb, such that the Pb isotopic evidence for recycled material was associated with maximum measured $^{40}$Ar/$^{36}$Ar ratios that were closer in composition to the atmospheric $^{40}$Ar/$^{36}$Ar. The correlation suggests recycling of atmospheric Ar to the mantle via subducting plates. Burnard (1999), however, challenged Sarda et al.’s (1999) interpretation of Ar recycling as basalts with elevated $^{206}$Pb/$^{204}$Pb are commonly found on topographic highs and those magmas would be more extensively degassed and more susceptible to atmospheric contamination, lowering their $^{40}$Ar/$^{36}$Ar closer to the atmospheric ratio. Consequently, the extent to which noble gases are recycled, whether the recycled gases are distributed homogeneously throughout the mantle, and the average and variance of the Ne, Ar and Xe isotopic composition of the MORB source all remain poorly constrained.

To better characterize the variability in the noble gas composition of the MORB source we present new He, Ne, Ar and Xe measurements. Four samples are measured nine samples from between 5°N and 5°S for heavy noble gas abundances and isotope ratios. Four samples are depleted MORBs from between the Ascension and Chain fracture zones and five samples are HIMU-type MORBs from between 5°N and the Romanche fracture zone (Fig. 1).

Fresh basaltic glass for each sample was prepared by leaching in 5% HNO$_3$ and/or in 1% oxalic acid to remove minor surface alteration. Approximately 3–5 g of glass was loaded into a piston crusher and baked under vacuum at 90–100°C for ~18 h to remove adsorbed gas. Glasses were step-crushed under vacuum, releasing the gas trapped in vesicles. Approximately 5–10 crushing steps were performed per sample, with each crushing step yielding a set of C, He, Ne, Ar, and Xe measurements.

2. Methods

2.1. Samples

The samples used in this study were dredged from the equatorial Mid-Atlantic Ridge during the cruises of R/V Endeavor EN061 and R/V Conrad RC2806. They have been previously characterized for major elements, trace elements, Sr, Nd, and Pb isotopic compositions (Hannigan et al., 2001; Schilling et al., 1994, 1995). Some samples were reanalyzed for Pb isotopes (Agranierr et al., 2005), and we use the newer measurements where applicable. For two of the samples, He isotopic compositions were previously determined by Graham et al. (1992b). We measured nine samples from between 5°N and 5°S for heavy noble gas abundances and isotope ratios. Four samples are depleted MORBs from between the Ascension and Chain fracture zones and five samples are HIMU-type MORBs from between 5°N and the Romanche fracture zone (Fig. 1).

CO$_2$ concentrations were determined by manometry using 10 Torr MKS manometers directly attached to the crusher volume. Gases were purified by sequential exposure to hot and cold SAES getters and a small split of the gas was let into a quadrupole mass spectrometer to determine the Ar abundance and an approximate $^{40}$Ar/$^{36}$Ar ratio. The noble gases were frozen onto a cryogenic cold trap and then sequentially released and let into the mass spectrometer. He, Ne, Ar, and Xe abundances and isotopic compositions were measured on a Nu Noblesse noble gas mass spectrometer operating in multicollector mode. $^{22}$Ne and $^{20}$Ne were corrected for isobaric interferences from doubly-charged $^{40}$Ar and $^{40}$Ar.

4) Are Ar and Xe being recycled back into the mantle, and if so, does this explain the differences in observed $^{40}$Ar/$^{36}$Ar and $^{129}$Xe/$^{134}$Xe ratios between MORBs and OIBs?

3) Are the HIMU-type MORBs also associated with primitive material as postulated for HIMU ocean islands (Day and Hilton, 2011; Parai et al., 2009)?

1) What is the Ne, Ar, and Xe isotopic composition of depleted MORB mantle (DMM) and of the extremely depleted MORBs (D-DMM; Workman and Hart, 2005)?

2) What is the degree of heterogeneity in Ne, Ar and Xe in the MORB source and how were these heterogeneities created?
Fig. 2. Linear arrays in $^{21}\text{Ne}/^{22}\text{Ne}$ vs. $^{20}\text{Ne}/^{22}\text{Ne}$ space for all nine samples that result from syn- to post-eruptive air contamination of bubbles containing mantle Ne. The mantle source $^{21}\text{Ne}/^{22}\text{Ne}$ was determined by $x$ and $y$ error-weighted least squares regression. Data points were translated such that the atmospheric composition was at (0,0). The $y$ data were then scaled by the square root of the ratio of the variance in $x$ to the variance in $y$ so as to put $x$ and $y$ on the same scale and fitted with an equation of the form $y = mx$. The uncertainties in the slopes (shown by the 1 error bands) were calculated using a Monte Carlo technique by randomly perturbing the data assuming normally distributed errors and refitting the slopes. The mantle value ($^{21}\text{Ne}/^{22}\text{Ne}$) was determined by extrapolation to $^{20}\text{Ne}/^{22}\text{Ne}$ of 12.5. The extrapolated values and their uncertainties are given in each panel.

$^{44}\text{CO}_2$ respectively. The Ar$^{+}$/Ar$^{-}$ was 0.061 ± 0.002 and the CO$_2$/(CO$_2$) was 0.0055 ± 0.0005. For the Ar measurements, depending upon the Ar abundance and approximate isotope ratio previously determined by the quadrupole mass spectrometer, a fraction of the gas was let into the mass spectrometer for more precise isotope ratio determination. Mass spectrometric procedure is described in Mukhopadhyay (2012). Typical procedural blanks for 4He, 22Ne, 36Ar and 130Xe prior to the starting crush were $2 \times 10^{-11}$, $5.7 \times 10^{-14}$, $3.7 \times 10^{-13}$, and $1.8 \times 10^{-16}$ cc, respectively. Blank levels were stable or decreasing during the course of crushing, with the exception of He, which increased by up to a factor of 2–4. Ne, Ar, and Xe blank isotopic ratios were statistically indistinguishable from atmosphere, and because the sample gases themselves are a mixture of mantle and shallow-level air contamination, no blank corrections were applied to the step-crushes. Mass discrimination for He was determined using the HH3 standard with a $^{3}$He/$^{4}$He ratio of 8.81.

A (Gayer et al., 2008) and for Ne, Ar, and Xe using air standards. Instrumental drift in sensitivity and mass discrimination was monitored through sample-standard bracketing with additional standards run overnight. Abundances and isotope ratios are reported in Tables S1 and S2.

### 2.3. Determination of mantle source isotopic ratios

For Ne, Ar, and Xe, each crushing step samples a population of bubbles reflecting variable mixtures of mantle-derived noble gases and syn- to post-eruptive atmospheric contaminants. As a result, noble gas compositions of bubbles define a mixing array that lies between the atmospheric end-member and the mantle end-member. In $^{20}\text{Ne}/^{22}\text{Ne}$ vs. $^{21}\text{Ne}/^{22}\text{Ne}$ space, the array is linear, while in Ne–Ar, Ne–Xe, or Ar–Xe space the array can be described by a hyperbolic mixing equation. Assuming the $^{20}\text{Ne}/^{22}\text{Ne}$ ratio of the MORB source mantle is 12.5 (see Section 3.3), a particular isotope ratio (e.g. $^{21}\text{Ne}/^{22}\text{Ne}$, $^{40}\text{Ar}/^{36}\text{Ar}$) can be regressed against $^{20}\text{Ne}/^{22}\text{Ne}$ and extrapolated to a $^{20}\text{Ne}/^{22}\text{Ne}$ of 12.5 to determine the mantle source value.

The mantle source $^{21}\text{Ne}/^{22}\text{Ne}$ was determined by $x$ and $y$ error-weighted least squares regression forced through the atmospheric composition ($y = mx$) and extrapolation to $^{20}\text{Ne}/^{22}\text{Ne}$ of 12.5 (Fig. 2). Ar and Xe isotope ratios for the mantle source were determined using an iterative nonlinear least squares hyperbolic fit constrained to go through the atmospheric composition ($Ax + Bxy + Cy = 0$). We present Ar and Xe mantle source values for two samples that show well-defined hyperbolic mixing arrays. Such hyperbolic arrays are likely to reflect either a single atmospheric contaminant or two elementally-fractionated atmospheric contaminants present in nearly fixed proportions. We denote all extrapolated mantle source isotope ratios extrapolated to a $^{20}\text{Ne}/^{22}\text{Ne}$ ratio of 12.5 with a subscript “E”.

### 2.4. Deconvolution of Pu- and U-derived fission Xe

To deconvolve U- from Pu-derived fissiogenic Xe, four mantle source Xe isotope ratios were used ($^{130,131,132,136}\text{Xe}/^{132}\text{Xe}$) following the methodology employed by Caffee et al. (1999) and Mukhopadhyay (2012). Because of the larger analytical uncertainty of $^{130}\text{Xe}$-normalized ratios, we use $^{132}\text{Xe}$-normalized ratios.
We assume the present-day mantle values represent a mixture of four end-members: initial Xe, $^{238}\text{U}$-produced Xe, $^{244}\text{Pu}$-produced Xe, and subducted atmospheric Xe. The solution $x$ to the system $A x = b$ was found by a least-squares inversion with the additional constraints $\sum x_i = 1$ and $0 \leq x_i \leq 1$. $A$ is a matrix that defines the four end-members in terms of the four isotope ratios; $x$ is a vector that defines the fraction of each of the four components in the present-day composition; and $b$ is a vector with the mantle source compositions (free of post-eruptive air contamination). To determine the elements of $b$, the mantle source $^{129}\text{Xe}/^{132}\text{Xe}$ was first determined from an Ar–Xe least-squares hyperbolic fit (Fig. S1). $^{130,131,134,136}\text{Xe}/^{132}\text{Xe}$ ratios were then linearly regressed against the $^{129}\text{Xe}/^{132}\text{Xe}$ ratio for the individual step-crushes. The linear regressions followed the same procedure as in Section 2.3 (see also Fig. 2). The slope and uncertainty in slope from the regressions were used to calculate the mantle $^{130,131,134,136}\text{Xe}/^{132}\text{Xe}$ ratios along with their uncertainties for the given mantle $^{129}\text{Xe}/^{132}\text{Xe}$ ratio. End-member and mantle source compositions ($A$ and $b$, respectively) were normalized to the standard deviations in the mantle source isotope ratios to assign equal weight to each isotope ratio. To compute the uncertainties in the deconvolution, a Monte Carlo technique was utilized; the mantle source compositions were varied at random within their $1\sigma$ uncertainties and the least-squares fit recomputed using the new values. For all simulations it was verified that convergence to a minimum was achieved.

3. Results

3.1. Depleted MORB noble gases

Samples between the Ascension and Chain fracture zones have been previously classified as depleted MORBs on the basis of isotopic and incompatible element ratios (Schilling et al., 1994). We note that the depleted MORB samples span the range of extremely depleted MORBs ($^{206}\text{Pb}/^{204}\text{Pb} = 17.68, ^{87}\text{Sr}/^{86}\text{Sr} = 0.7021$) to MORBs showing an average degree of depletion ($^{206}\text{Pb}/^{204}\text{Pb} = 18.35, ^{87}\text{Sr}/^{86}\text{Sr} = 0.7026$).

The $^{4}\text{He}/^{3}\text{He}$ ratios of the depleted MORBs vary between 80,800–86,700 with the more depleted Sr, Nd, and Pb isotopic composition associated with the lower $^{4}\text{He}/^{3}\text{He}$ ratios (Figs. 3 and S2a). $^{21}\text{Ne}/^{22}\text{Ne}$ values for the depleted MORB samples range from 0.062 to 0.065 (Table 1; Figs. 2, 4 and S2b). For the depleted MORBs, the He and Ne isotopic compositions are negatively correlated (Fig. 5).

The highest measured $^{40}\text{Ar}/^{36}\text{Ar}$ ratio in the depleted samples is $30,400 \pm 300$. We note that sample RC2806 3D-2 has a lithophile isotopic composition very similar to average N-MORB source ($^{206}\text{Pb}/^{204}\text{Pb} = 18.35, ^{87}\text{Sr}/^{86}\text{Sr} = 0.7026, ^{143}\text{Nd}/^{144}\text{Nd} = 0.513126$; cf. DMM: 18.275, 0.70263, 0.51313, Workman and Hart, 2005) and is a matrix that defines the fraction of each of the four end-members in terms of the four isotope ratios; $A$ is a vector that defines the four end-members in terms of the four isotope ratios; $b$ is a vector with the mantle source compositions (free of post-eruptive air contamination). The parameters $\alpha$ and $\beta$ are limited variations. However, given the sample's lithophile isotopic composition, and in the absence of substantial number of step-crushes for the other samples, we assume the extrapolated Ar and Xe isotopic composition for RC2806 3D-2 is broadly representative of the depleted group. The best fit hyperbola for this sample yields a mantle source $^{40}\text{Ar}/^{36}\text{Ar}$ of $41,500 \pm 9000$ (Table 1; Fig. 6a). The uncertainty in $^{40}\text{Ar}/^{36}\text{Ar}$ is related to a small number of data points with respect to the scatter in the data and suggests additional measurements on this sample in the future will be able to better constrain the hyperbolic fit. For a consistent comparison, we refit the step-crushing popping rock data (Moreira et al., 1998) using the same technique (Section 2.3), which yielded a mantle source $^{40}\text{Ar}/^{36}\text{Ar}$ of $25,200 \pm 600$ (Fig. 6a), consistent with the estimate of $27,000 \pm 4000$ from UV laser ablation of individual bubbles (Raquin et al., 2008).

The highest measured $^{129}\text{Xe}/^{130}\text{Xe}$ in the depleted sample group is $7.81 \pm 0.06$. Xe isotopes ($^{129}\text{Xe}/^{130}\text{Xe}$) correlate better with Ar isotopes than with Ne isotopes. Consequently, we obtained the best-fit hyperbola in Ar–Xe space for sample RC2806 3D-2 to constrain the mantle source $^{129}\text{Xe}/^{130}\text{Xe}$ to $7.77 \pm 0.06$ at a $^{40}\text{Ar}/^{36}\text{Ar}$ of $41,500$ (Table 1; Fig. 7a). Because of
the curvature of the best-fit hyperbola, the mantle $^{129}\text{Xe}/^{130}\text{Xe}$ is not particularly sensitive to the uncertainty in $^{40}\text{Ar}/^{36}\text{Ar}$.

### 3.2. HIMU-type MORB noble gases

Samples north of the Romanche fracture zone are classified as enriched HIMU-type MORBs and span the range of $^{206}\text{Pb}/^{204}\text{Pb}$ from 18.75 to 19.64 and $^{87}\text{Sr}/^{86}\text{Sr}$ from 0.7026 to 0.7029. Although the Pb isotopic compositions in these samples do not reach the extreme HIMU compositions observed in OIBs, we refer to them as “HIMU-type” following the identification of a HIMU component in these basalts (Schilling et al., 1994) to distinguish them from the depleted MORBs south of the equator.

$^{4}\text{He}/^{3}\text{He}$ ratios of the HIMU-type MORBs are between 93,200 and 99,000—more radiogenic than the depleted MORBs.

### Table 1: Mantle source compositions.

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<th>$^{129}\text{Xe}/^{132}\text{Xe}$</th>
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* Popping rock data is from Moreira et al. (1998) and Raquin and Moreira (2009); $^{129}\text{Xe}/^{130}\text{Xe}$ is based on an Ar–Xe hyperbolic fit using a mantle source $^{40}\text{Ar}/^{36}\text{Ar}$ ratio of 27,000.

* Iceland data is from Mukhopadhyay (2012).

* Bravo Dome data is from Holland and Ballentine (2006).

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**Fig. 4.** Relationship between $^{21}\text{Ne}/^{22}\text{Ne}$ ratios and (a) $^{206}\text{Pb}/^{204}\text{Pb}$ and (b) $^{143}\text{Nd}/^{144}\text{Nd}$. Symbols as in Fig. 3. The solid lines are illustrative two-component mixing hyperbolas for mixing between a depleted mantle (D-DMM) and the HIMU plume component. The Pb and Nd isotope ratios of the two components are as in Fig. 3. The $^{21}\text{Ne}/^{22}\text{Ne}$ ratio of D-DMM is 0.065 (this work) and the $^{21}\text{Ne}/^{22}\text{Ne}$ ratio of HIMU is 0.05 (Parai et al., 2009). The curvature of the hyperbola is controlled by the parameter $r$, defined as, e.g., $(^{21}\text{Ne}/^{22}\text{Ne})_{\text{D-DMM}}/(^{21}\text{Ne}/^{22}\text{Ne})_{\text{HIMU}}$.

**Fig. 5.** Relationship between $^{4}\text{He}/^{3}\text{He}$ ratios and $^{21}\text{Ne}/^{22}\text{Ne}$ ratios. Symbols as in Fig. 3. Note that the sample with the most depleted Pb, Sr, and Nd isotopic composition has a lower $^{4}\text{He}/^{3}\text{He}$ ratio but higher nucleogenic $^{21}\text{Ne}/^{22}\text{Ne}$. Overall, there is a general tendency for increasing $^{4}\text{He}/^{3}\text{He}$ ratios to be associated with decreasing $^{21}\text{Ne}/^{22}\text{Ne}$ within the depleted MORB group. The HIMU-type MORBs are displaced to more radiogenic He and less nucleogenic Ne than the depleted MORBs. Overall, the He–Ne relationships of the MORBs can be explained by a mixture of depleted mantle (D-DMM) and HIMU, with the requirement that HIMU is composed of both recycled and less degassed mantle material, such as FOZO (Hart et al., 1992). The solid line is an illustrative mixing hyperbola using the end-member compositions as in Figs. 3 and 4. The curvature $r = (^{21}\text{Ne}/^{22}\text{Ne})_{\text{D-DMM}}/(^{21}\text{Ne}/^{22}\text{Ne})_{\text{HIMU}} = 5.6$ is consistent with the hyperbolas in Figs. 3 and 4, e.g., $r_{\text{Pb-Pb}}/r_{\text{Ne-Pb}} = r_{\text{He-Ne}}$.

The curvature of the best-fit hyperbola, the mantle $^{129}\text{Xe}/^{130}\text{Xe}$ is not particularly sensitive to the uncertainty in $^{40}\text{Ar}/^{36}\text{Ar}$.

### 3.2. HIMU-type MORB noble gases

Samples north of the Romanche fracture zone are classified as enriched HIMU-type MORBs and span the range of $^{206}\text{Pb}/^{204}\text{Pb}$ from 18.75 to 19.64 and $^{87}\text{Sr}/^{86}\text{Sr}$ from 0.7026 to 0.7029. Although the Pb isotopic compositions in these samples do not reach the extreme HIMU compositions observed in OIBs, we refer to them as “HIMU-type” following the identification of a HIMU component in these basalts (Schilling et al., 1994) to distinguish them from the depleted MORBs south of the equator.

$^{4}\text{He}/^{3}\text{He}$ ratios of the HIMU-type MORBs are between 93,200 and 99,000—more radiogenic than the depleted MORBs.
21Ne/22Ne values for the HIMU-type MORB samples are between 0.056 and 0.061, which is less nucleogenic than the depleted MORBs (Table 1; Figs. 2, 4, 5 and S2b). The highest measured 40Ar/36Ar ratio in the HIMU-type lavas is 17,800 ± 7, significantly lower than the highest measured values in the depleted sample group. Sample RC2806 57D-1 yields the best mixing hyperbola in Ne–Ar space, which constrains the source 40Ar/36Ar ratio to be 18,100 ± 600 (Table 1; Fig. 6b) and, like for the depleted MORB group, we assume this sample’s Ar and Xe isotopic composition is broadly representative of the enriched HIMU-type MORBs. The highest measured 129Xe/130Xe in the depleted sample group is 7.26 ± 0.05. The Ar–Xe mixing array for the same sample (RC2806 57D-1) defines the mantle source 129Xe/130Xe of 7.21 ± 0.02 at a 40Ar/36Ar of 18,100 (Table 1; Fig. 7b). The curvature of the best-fit hyperbola indicates that 129Xe/130Xe is not particularly sensitive to the uncertainty in the mantle 40Ar/36Ar.

3.3. A note on the mantle 20Ne/22Ne ratio

We recognize that the 20Ne/22Ne value of the whole mantle may not be 12.5, and the source of mantle plumes may be associated with 20Ne/22Ne of 12.9, or higher (Mukhopadhyay, 2012; Yokochi and Marty, 2004). We choose a MORB source value of 12.5 based on the mantle source composition of continental well gases (Ballentine and Holland, 2008; Holland and Ballentine, 2006) and the gas-rich popping rock (Raquin et al., 2008). Furthermore, our highest measured value in both sample groups (12.36 in depleted samples; 12.43 in HIMU-type samples), and MORBs in general (Kunz, 1999; Moreira et al., 1998, 2011; Raquin and Moreira, 2009; Sarda et al., 1988) are close to 12.5. Correcting all the isotope ratios to a common 20Ne/22Ne allows for a consistent comparison between samples. Our primary results are not affected if a higher value is chosen. For example, if the mantle 20Ne/22Ne and 23Ne/22Ne varied as a mixture between the “MORB array” (12.5, 0.059; Sarda et al., 1988) and “Iceland array” (13.8, 0.0375; Mukhopadhyay, 2012) the least nucleogenic sample in our dataset would intersect this mixture at 20Ne/22Ne = 12.65, causing a negligible shift in the extrapolated 40Ar/36Ar and 129Xe/130Xe. If the mantle 20Ne/22Ne of the MORB source were higher than ~13, our results would actually be strengthened. Because of the curvature of the hyperbolas (Figs. 6 and 7), the depleted sample would extrapolate to higher 40Ar/36Ar and 129Xe/130Xe while the HIMU-type sample would be little affected, amplifying the compositional difference between them.

4. Discussion

4.1. A primitive Ne component in HIMU-type basalts

The 4He/3He ratios in the HIMU-type MORBs span the range of ~80,000–100,000, within the typical MORB range (Graham, 2002). The 4He/3He ratios of the HIMU-type lavas are, however, all more radiogenic than the depleted MORBs and are associated
with more radiogenic $^{206}\text{Pb}^{204}\text{Pb}$ (Fig. 3). Since HIMU mantle plumes are characterized by radiogenic He and Pb isotopic ratios (Chauvel et al., 1992; Day and Hilton, 2011; Graham et al., 1992a; Hanyu et al., 1999; 2011; Hauri and Hart, 1993; Hilton et al., 2000; Parai et al., 2009), the He–Pb co-variation is consistent with the injection of a HIMU-type plume into the Equatorial Mid-Atlantic Ridge (Schilling et al., 1994).

The relationship between Ne and He isotopic compositions provides insight into the genesis of the HIMU plume sampled at the equatorial Mid-Atlantic Ridge. HIMU plumes are often assigned to recycled oceanic crust (Chauvel et al., 1992; Hauri and Hart, 1993; Hofmann, 1997). In addition to radiogenic $^{206}\text{Pb}^{204}\text{Pb}$, such a component should have both radiogenic He and nucleogenic Ne isotopic ratios. This is because in highly degassed oceanic crust, nucleogenic $^{21}\text{Ne}$ will be produced from $^{18}\text{O}(\alpha,\nu)$ and $^{24}\text{Mg}(\alpha,\nu)$ reactions with the reacting alpha particles and neutrons ultimately sourced from U and Th decay. Thus, the less nucleogenic Ne in the HIMU-type lavas compared to the depleted MORBs cannot be generated solely through sampling of recycled material or mixing recycled material with DMM.

Our observation of less nucleogenic Ne and more radiogenic He have been observed in other HIMU localities, such as the Cameroon line of volcanoes (Barford et al., 1999) and the Cook–Austral islands (Parai et al., 2009). The simplest explanation for the less nucleogenic signature is the involvement of a relatively undegassed mantle, such as FOZO. In this regard, Parai et al. (2009) associated the lower radiogenic $^{40}\text{Ar}/^{36}\text{Ar}$ for the less nucleogenic He than MORBs with less nucleogenic Ne in the HIMU Cook–Austral islands to mixing between degassed recycled crust and FOZO (also see Day and Hilton, 2011). Based on the lithophile isotopes, Schilling et al. (1994) suggested that the plume-influenced MORBs north of the equator reflect mixing between a depleted mantle, recycled crust (HIMU) and FOZO. Our present study is consistent with the suggestion of a HIMU component in the north equatorial Atlantic MORBs (Schilling et al., 1994) and provides further evidence that HIMU is a hybrid of at least two different materials—a degassed recycled component and a relatively undegassed mantle.

4.2. A negative correlation between He and Ne in depleted MORBs

Our new observations from the equatorial Atlantic indicate that within the depleted group there appears to be an overall trend of more nucleogenic $^{21}\text{Ne}/^{22}\text{Ne}$ associated with less radiogenic He and more depleted Sr, Nd and Pb isotopic compositions (Figs. 3–5). This observation places constraints on the $^{4}\text{He}$/$^{3}\text{He}$ and $^{21}\text{Ne}/^{22}\text{Ne}$ variability in MORBs, and the relationship between the most depleted MORBs (D-DMM; Workman and Hart, 2005) and N-MORBs (DMM).

Helium and neon isotopic variations in the MORB source are often interpreted in terms of a flux of $^{3}\text{He}$ and $^{22}\text{Ne}$ from the OIB source into the MORB source followed by mixing with in situ radiogenic $^{4}\text{He}$ and nucleogenic $^{21}\text{Ne}$ (e.g. Allegre et al., 1987; Porcelli and Wasserburg, 1995; Tolstikhin and Hofmann, 2005). However, differences in $^{20}\text{Ne}/^{22}\text{Ne}$ and $^{4}\text{He}/^{22}\text{Ne}$ ratios between MORBs and OIBs suggests that not all the primordial He and Ne in the MORB mantle can be derived from the OIB mantle source (Honda and McDougall, 1998; Mukhopadhyay, 2012; Yokochi and Marty, 2004). Assuming $^{3}\text{He}$ is indigenous to the MORB source, variations in MORB $^{4}\text{He}/^{3}\text{He}$ ratios may be explained in the framework of a marble cake assemblage comprising pyroxenite veins in a depleted peridotite matrix (e.g. Graham et al., 2001; Hamelin et al., 2011). The $^{4}\text{He}/^{3}\text{He}$ ratio of the most depleted MORB in our dataset lies at the lower limit of the MORB range from 74,000 to 120,000 for regions removed from obvious influences of mantle plumes (Georgen et al., 2003; Graham, 2002; Graham et al., 1992b, 2001). The addition of pyroxenites (recycled crust) with radiogenic $^{4}\text{He}$ to this depleted peridotitic matrix could then produce the range of $^{4}\text{He}/^{3}\text{He}$ ratios and lithophile isotopes in non-plume influenced MORBs (Figs. 3 and 5a). Within the marble cake framework, recycled oceanic crust should be associated with nucleogenic $^{21}\text{Ne}/^{22}\text{Ne}$ ratios in addition to radiogenic $^{4}\text{He}/^{3}\text{He}$ ratios. So as $^{4}\text{He}/^{3}\text{He}$ ratios become more radiogenic in MORBs through sampling a greater proportion of pyroxenites, $^{21}\text{Ne}/^{22}\text{Ne}$ ratios should become more nucleogenic, opposite to the observed relation between He and Ne (Fig. 5). Hence, the observed isotopic relations (Figs. 3–5) cannot be generated by mixing depleted peridotites with pyroxenites that have radiogenic He and nucleogenic Ne.

Instead, we hypothesize that the observed He–Ne relationship requires the equatorial Atlantic N-MORBs to be produced by the addition of a small proportion of a HIMU plume component to a very depleted MORB source (D-DMM; e.g. Fig. 5). We note that based on lithophile isotopes Schilling et al. (1994) suggested that the depleted section of the Mid-Atlantic Ridge between the Ascension and Romanche fracture zones resulted from mixing between an extremely depleted MORB source and an enriched HIMU component. Our results are consistent with this interpretation with the additional requirement that HIMU is itself a mixture of recycled and primitive components (Section 4.1). While a marble cake mantle remains a viable way of interpreting the geochemical observations in depleted MORBs, based on our new He–Ne observations, it appears to be richer in flavor than a simple mixture of recycled pyroxenites and depleted peridotites.

4.3. Argon isotopic composition of DMM and the HIMU-type MORB source

The depleted MORB sample RC2806 3D-2 has a mantle source $^{40}\text{Ar}^{36}\text{Ar}$ ratio of 41,500 $\pm$ 9000 (Fig. 6a). This is similar to the $^{40}\text{Ar}^{36}\text{Ar}$ ratio of 41,050 $\pm$ 2670 for the Bravo Dome well gas and the maximum measured $^{40}\text{Ar}^{36}\text{Ar}$ in MORBs of 42,400 $\pm$ 7900 (Marty and Humbert, 1997). In contrast, we compute the popping rock mantle source $^{40}\text{Ar}^{36}\text{Ar}$ to be 25,200 $\pm$ 600 from the published step-crushing data (Fig. 6a) (Moreira et al., 1998), similar to the value of 25,000 reported in that work and 27,000 $\pm$ 4000 obtained by UV laser ablation of individual bubbles (Raquin et al., 2008). We note that a laser ablation study by Burnard et al. (1997) yielded a $^{40}\text{Ar}^{36}\text{Ar}$ ratio of 40,000 $\pm$ 4000 in an individual bubble in the popping rock sample, higher than the values reported by Raquin et al. (2008) and those of Moreira et al. (1998). While the mantle source composition of popping rock is debatable (see Raquin et al., 2008 for further discussion), DMM has a $^{40}\text{Ar}^{36}\text{Ar}$ ratio of at least 41,000.

The HIMU-type MORB sample RC2806 57D-1 has a mantle source $^{40}\text{Ar}^{36}\text{Ar}$ ratio of 18,100 $\pm$ 600, unequivocally lower than both DMM and the popping rock mantle source (Fig. 6b). The result demonstrates that large variations (factor of 2) in the MORB mantle source $^{40}\text{Ar}^{36}\text{Ar}$ ratio exist even when variations in He and Ne isotopic compositions are more muted. The low $^{40}\text{Ar}^{36}\text{Ar}$ ratio in the HIMU-type MORBs likely results from the injection of a HIMU mantle plume with low $^{40}\text{Ar}^{36}\text{Ar}$ into the Mid-Atlantic Ridge. Plumes feeding OIB are often characterized by low $^{40}\text{Ar}^{36}\text{Ar}$ (Honda et al., 1993; Mukhopadhyay, 2012; Raquin and Moreira, 2009; Trieloff et al., 2000, 2002; Valbracht et al., 1997). The low $^{40}\text{Ar}^{36}\text{Ar}$ in plumes of the HIMU-type MORBs of the north equatorial Atlantic could be due to sampling of a less degassed reservoir with low $^{40}\text{Ar}^{36}\text{Ar}$ ratio, an explanation that would be consistent with Ne isotopic evidence for a less degassed component in the HIMU plume. Alternatively, the low $^{40}\text{Ar}^{36}\text{Ar}$ ratio could be a result of recirculation of atmospheric Ar associated with recycled crust to the mantle source of plumes.
(Holland and Ballentine, 2006). We discuss these two separate hypotheses while discussing Xe isotopes in the next section.

4.4. Xenon isotopic differences between DMM, HIMU-type MORB and Iceland plume sources

The depleted MORB sample RC2806 3D-2 has a mantle source $^{129}\text{Xe}/^{130}\text{Xe}$ ratio of 7.77 ± 0.06 (Fig. 7a). This ratio overlaps with the $^{129}\text{Xe}/^{130}\text{Xe}$ ratio of 7.9 ± 0.14 for the mantle source of the Bravo Dome well gas (Holland and Ballentine, 2006). On the other hand, the HIMU-type MORB sample RC2806 57D-1 has a mantle source $^{129}\text{Xe}/^{130}\text{Xe}$ of 7.21 ± 0.05, significantly lower than the depleted MORB source (Fig. 7b). Like the less nucleogenic $^{21}\text{Ne}/^{22}\text{Ne}$ ratio and less radiogenic $^{40}\text{Ar}/^{36}\text{Ar}$ ratio, the lower $^{129}\text{Xe}/^{130}\text{Xe}$ in the HIMU-type MORBs could reflect sampling of a less degassed reservoir with a lower $I/^{36}\text{Xe}$ ratio than the DMM source. $^{129}\text{I}$, which produces $^{129}\text{Xe}$, became extinct at 4.45 Ga and therefore differences in the degree of degassing between DMM and the less degassed reservoir would have to have been initiated prior to 4.45 Ga and the reservoirs could not have subsequently been homogenized. Alternatively, the lower $^{129}\text{Xe}/^{130}\text{Xe}$ in the HIMU-type MORBs could reflect preferential recycling of atmospheric Xe into the HIMU plume source. These two hypotheses can be addressed by considering the $^{129}\text{Xe}/^{136}\text{Xe}$ ratio, which reflects the time-integrated $^{239}\text{Pu}/(^{238}\text{U} + ^{239}\text{U})$ ratio (Fig. 8).

Given the uncertainties in Xe isotopic ratios, we cannot yet identify possible Xe isotopic differences within the depleted or HIMU-type groups. As a result, in Fig. 8, we combine the step-crushes for each sample group to investigate whether differences in Xe isotopic composition exist between the depleted and HIMU-type groups and whether the depleted MORBs are isotopically similar to or different from the slope calculated for all the data taken together.

The distinct $^{129}\text{Xe}/^{130}\text{Xe}$ ratios for the two MORB groups can be clearly seen in $^{129}\text{Xe}/^{130}\text{Xe}$ vs. $^{136}\text{Xe}/^{130}\text{Xe}$ space when the error-weighted average compositions of the crushing steps are plotted (Fig. 8b). The data are not corrected for post-eruptive contamination, so the true mantle source (extrapolated) values lie further along the particular mixing line with air. But extrapolation is unnecessary to demonstrate the simple conclusion that the HIMU-type MORBs have a $^{129}\text{Xe}/^{136}\text{Xe}$ ratio in excess of air while the depleted MORBs have a ratio lower than air. The data in Fig. 8, therefore, clearly demonstrate that the depleted MORBs and the HIMU-type MORBs have distinct $^{129}\text{Xe}/^{136}\text{Xe}$ ratios. Furthermore, a clear distinction is observed between the Iceland plume composition (Mukhopadhyay, 2012) and the depleted MORBs, with the HIMU-type MORBs intermediate between the two.

In both of the Xe–Xe isotopic spaces (Fig. 8a and b), recycling of atmospheric Xe would shift the composition of a reservoir linearly towards air. Thus, neither the small difference in slope (Fig. 8a) nor the measured difference in the $^{129}\text{Xe}/^{136}\text{Xe}$ ratio (Fig. 8b) between the depleted and HIMU-type MORBs can be produced solely through recycling of atmospheric Xe. We conclude that $^{129}(^{244}\text{Pu} + ^{238}\text{U})$ ratios are different between the two groups of MORBs, reflecting different degrees of mantle outgassing. The lower $^{125}\text{Xe}/^{136}\text{Xe}$ ratio of DMM is consistent with a more degassed source than the mantle source of HIMU-type MORBs (Fig. 8). This is because a more degassed source will have lower concentrations of primordial Xe isotopes ($^{130}\text{Xe}$), and lower concentrations of $^{129}\text{Xe}$ from decay of extinct $^{129}\text{I}$ and $^{136}\text{Xe}$ from fission of extinct $^{244}\text{Pu}$. Adding $^{129}\text{Xe}$ from $^{238}\text{U}$ fission or $^{244}\text{Pu}$ fission while $^{244}\text{Pu}$ is still alive to such a degassed source will lead to a decrease in $^{129}\text{Xe}/^{130}\text{Xe}$ ratio. The higher $^{125}\text{Xe}/^{136}\text{Xe}$ ratio of the HIMU-type MORBs likely reflects mixing between a less degassed source, such as the reservoir supplying primordial noble gases to the Iceland plume (Mukhopadhyay, 2012), and more degassed sources, such as recycled crust and DMM. In this regard, the $^{129}\text{Xe}/^{136}\text{Xe}$ evidence suggests that the lower $^{40}\text{Ar}/^{36}\text{Ar}$ ratio in the HIMU-type MORBs also cannot originate solely from recycling of atmospheric Ar; sampling of a less degassed mantle reservoir with low $^{40}\text{Ar}/^{36}\text{Ar}$ ratio is required.

4.5. $^{239}\text{Pu}–^{238}\text{U}$ derived Xe in the depleted and the HIMU-type MORB mantle source

The proportion of Pu- to U-derived Xe provides powerful independent constraints on the degree of outgassing of a mantle source that augments arguments from the other noble gas isotope ratios. Because fission Xe yields from Pu are significantly larger than from U, a reservoir that has remained completely closed over Earth’s history will have $^{136}\text{Xe}/^{138}\text{Xe}$ of ~27, where *refers to fissogenic Xe. Because $^{244}\text{Pu}$ became extinct at ~4 Ga, reservoirs open to substantial volatile loss over the past 4 billion years.

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**Fig. 8.** (a) $^{129}\text{Xe}/^{130}\text{Xe}$ vs. $^{136}\text{Xe}/^{130}\text{Xe}$ ratios of step-crushes for depleted and HIMU-type MORB. Symbols as in Fig. 3. The solid lines represent the x–y error weighted best fit line and the gray bands denote the 1σ error envelopes. The methodology for the least squares regression was outlined in Section 2.3 and in Fig. 2. The slope of the line is a function of the time integrated $^{(244}\text{Pu} + ^{238}\text{U})/^{129}\text{Xe}$ ratio. The slopes are 0.375 ± 0.009 and 0.334 ± 0.009 for depleted and HIMU-type MORBs, respectively. The slope with all data taken together is 0.338 ± 0.006, which does not overlap the slope of either group. The difference in slopes demonstrates a difference in the time integrated $^{(244}\text{Pu} + ^{238}\text{U})/^{129}\text{Xe}$ ratio within MORBs. (b) $^{129}\text{Xe}/^{130}\text{Xe}$ vs. $^{136}\text{Xe}/^{130}\text{Xe}$ isotope plot showing the weighted average composition of DICE 10 (Iceland; Mukhopadhyay, 2012), depleted MORB and HIMU-type MORB step-crushes. Symbols as in Fig. 3. The air-Iceland mixing line is distinct from the air-depleted MORB mixing line. Consequently, the plume Xe signal cannot be related to MORBs solely through addition of subducted atmospheric Xe or through addition of $^{238}\text{U}$-produced fission $^{136}\text{Xe}$. Sampling of a reservoir that is less degassed than DMM is required to explain the Xe isotopic composition of the HIMU-type MORBs and the Iceland plume.
In Earth’s history (Moreira and Raquin, 2007). However, since the recycling of atmospheric Xe to the mantle is required. We find that on order 80–90% of the $^{132}\text{Xe}$ in the HIMU-type MORB source, and (d) I/Pu ratios from Pu fission is 0.52$^{+0.13}_{-0.09}$ (solar) or 0.63$^{+0.18}_{-0.12}$ (chondritic) (Table 2). Although the uncertainties are large, the result suggests that the HIMU-MORB sources may have a higher proportion of Pu- to U-derived fission Xe than the depleted MORBs but lower than the Iceland plume (Table 2; see also Fig. S3). The high proportion of Pu-derived Xe in the HIMU-type MORBs compared to the depleted MORBs is inconsistent with the HIMU component being comprised only of recycled crust or a mixture of recycled crust and DMM. This is because recycled crust should be highly degassed and will have Xe fission from 238U; i.e., $^{136}\text{Xe}_{\text{Pu}}$/$^{136}\text{Xe}_{\text{U}}$ in degassed reservoirs will be $\ll 27$ (Kunz et al., 1998; Ozima et al., 1985; Tolstikhin and O’Nions, 1996; Yokochi and Marty, 2005). In the present day mantle, $^{131,132,134,136}\text{Xe}$ isotopes reflect a mixture of the initial mantle Xe, Pu- and U-produced fission Xe, and subducted atmospheric Xe, if any. For the initial mantle composition we investigated solar and chondritic Xe (AVCC) (Caffee et al., 1999; Holland and Ballentine, 2006; Holland et al., 2009; Pujol et al., 2011). The deconvolution of the four Xe components was performed with the mantle source ratios listed in Table 1 and the methodology discussed in Section 2.4. Here we will note four major implications of the fission deconvolution: (a) recycling of atmospheric Xe to the mantle, (b) the ratio of Pu- to U-derived Xe in the depleted MORB source, (c) the ratio of Pu- to U-derived Xe in the HIMU-MORB source, and (d) I/Pu ratios in the depleted and HIMU-MORB sources.

For a mantle starting composition of either solar Xe or chondritic Xe, substantial injection of atmospheric Xe back into the mantle is required. We find that on order 80–90% of the $^{132}\text{Xe}$ in the mantle source is derived from recycled atmospheric Xe (Table 2). The proportion of recycled Xe in the HIMU-type MORBs, however, does not appear to be larger than in the depleted MORBs (Table 2). Thus, we suggest that the lower $^{129}\text{Xe}/^{130}\text{Xe}$ ratios in the enriched HIMU-type MORBs do not result from preferential recycling of atmospheric Xe into the plume source. Rather the HIMU-type MORBs must sample a reservoir with a lower $^{129}\text{Xe}/^{130}\text{Xe}$ than the depleted MORBs.

The requirement that a significant fraction of the fissionogenic Xe isotopes in the mantle is recycled from the atmosphere (Table 2) is consistent with previous work that suggests Xe can be recycled back into the mantle (Holland and Ballentine, 2006; Kendrick et al., 2011; Sumino et al., 2010). The recycling of atmospheric Xe could have happened all through Earth’s history associated with plate subduction (Holland and Ballentine, 2006), or very early on in Earth’s history (Moreira and Raquin, 2007). However, since the $^{129}\text{Xe}/^{130}\text{Xe}$ ratio can only evolve during the first 100 Myr of solar system history, the distinct atmospheric and mantle compositions require that recycling efficiency of atmospheric Xe must be relatively low so as not to homogenize the mantle and atmospheric compositions over 4.45 Ga. In this regard, since at most 80–90% of the Xe in the mantle is recycled (Table 2), the mantle $^{129}\text{Xe}/^{130}\text{Xe}$ ratio can still be distinctly different from the atmospheric ratio.

## Table 2

Xenon components in the depleted MORB and HIMU-type MORB mantle sources.

<table>
<thead>
<tr>
<th>Starting mantle composition</th>
<th>Sample</th>
<th>Fraction $^{129}\text{Xe}$ from recycled air</th>
<th>Fraction $^{129}\text{Xe}$ from SWAVCC</th>
<th>Fraction $^{132}\text{Xe}$ from $^{238}\text{U}$</th>
<th>Fraction $^{132}\text{Xe}$ from $^{244}\text{Pu}$</th>
<th>Fraction $^{136}\text{Xe}$ from $^{244}\text{Pu}$</th>
<th>$^{129}\text{Xe}^{132}\text{Xe}$/$^{136}\text{Xe}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solar wind</td>
<td>Depleted MORBs</td>
<td>0.896 ± 0.054</td>
<td>0.058 ± 0.049</td>
<td>0.027 ± 0.004</td>
<td>0.019 ± 0.008</td>
<td>0.29$^{+0.14}_{-0.12}$</td>
<td>8.9$^{+6.5}_{-3.1}$</td>
</tr>
<tr>
<td></td>
<td>HIMU-type MORBs</td>
<td>0.778 ± 0.041</td>
<td>0.188 ± 0.038</td>
<td>0.013 ± 0.003</td>
<td>0.022 ± 0.006</td>
<td>0.52 ± 0.13</td>
<td>4.6$^{+1.9}_{-1.0}$</td>
</tr>
<tr>
<td>AVCC</td>
<td>Depleted MORBs</td>
<td>0.910 ± 0.052</td>
<td>0.045 ± 0.049</td>
<td>0.028 ± 0.005</td>
<td>0.017 ± 0.008</td>
<td>0.25$^{+0.17}_{-0.10}$</td>
<td>10.7$^{+4.0}_{-2.5}$</td>
</tr>
<tr>
<td></td>
<td>HIMU-type MORBs</td>
<td>0.767 ± 0.047</td>
<td>0.201 ± 0.044</td>
<td>0.009 ± 0.004</td>
<td>0.023 ± 0.007</td>
<td>0.63 ± 0.18</td>
<td>4.1$^{+1.9}_{-1.0}$</td>
</tr>
</tbody>
</table>

Note: $^{136}\text{Xe}/^{138}\text{Xe}$ is the ratio of radiogenic $^{136}\text{Xe}$ from decay of $^{136}\text{I}$ to fissionogenic $^{136}\text{Xe}$ derived from $^{244}\text{Pu}$ fission. The distributions of $^{129}\text{Xe}^{132}\text{Xe}$ are skewed and hence, the median and the 67% confidence intervals are presented. We use the following method to determine $^{129}\text{Xe}$ ratios: Depending on whether the initial mantle Xe composition is solar or chondritic, the fraction of $^{136}\text{Xe}$ derived from Pu fission in the depleted MORB source is 0.29$^{+0.10}_{-0.09}$ or 0.29$^{+0.12}_{-0.10}$, respectively (Table 2). The corresponding $^{136}\text{Xe}^{136}\text{Xe}$ ratio is 0.33 ± 0.12 or 0.41 ± 0.1. The $^{136}\text{Xe}^{136}\text{Xe}$ ratios are ~1% of the closed system value of 27 and suggest that the depleted MORB source has lost 99% of its Pu-produced Xe as a result of mantle outgassing from a combination of giant impacts and mantle convection.

In the HIMU-type MORB source, the fraction of $^{136}\text{Xe}$ derived from Pu fission is 0.52 ± 0.13 (solar) or 0.63 ± 0.18 (chondritic) (Table 2). Although the uncertainties are large, the result suggests that the HIMU-MORB sources may have a higher proportion of Pu- to U-derived fission Xe than the depleted MORBs but lower than the Iceland plume (Table 2; see also Fig. S3). The high proportion of Pu-derived Xe in the HIMU-type MORBs compared to the depleted MORBs is consistent with the HIMU component being comprised only of recycled crust or a mixture of recycled crust and DMM. This is because recycled crust should be highly degassed and will have Xe fission from 238U fission during its residence in the mantle. Pu-produced Xe could be present in recycled crust only if the crust were produced and subducted back into mantle prior to 4 Ga when $^{244}\text{Pu}$ was still alive. However, Re–Os and Pb isotopic systematics indicate an age of 1–2 Ga for the HIMU mantle (Chauvel et al., 1992; Hauri and Hart, 1993). Therefore, the relatively high proportion of Pu- to U-derived fission Xe in the HIMU-type mantle provides further evidence that HIMU is a hybrid of a recycled and a less degassed mantle component and requires the Xe contribution from the less degassed mantle component to dominate over the 238U-produced fission Xe in the recycled crust. Such a conclusion is consistent with the HIMU-type MORBs having a $^{129}\text{Xe}^{136}\text{Xe}$ ratio that is intermediate between Iceland and the depleted MORBs.
Interpreting these values as closure ages for a mantle with an initially homogenous I/Pu ratio, the higher $^{129}Xe/^{130}Xe_E$ ratio in the depleted MORB source would imply that the shallow upper mantle became closed to volatile loss prior to the deep mantle reservoir supplying noble gases to the HIMU plume. Such a conclusion appears paradoxical. Rather, a simpler explanation is that the lower $^{129}Xe/^{130}Xe_E$ in the HIMU-type MORB source reflects a lower initial I/Pu ratio for the plume source compared to the MORB source. This is consistent with the conclusions inferred from the Xe data in the Iceland plume (Mukhopadhyay, 2012) and suggests that the initial phase of Earth’s accretion was volatile poor compared to the later stages of accretion because Pu is a refractory element while I is a volatile element.

Differences in I/Pu ratios and I/Xe ratios in the mantle are relicts of accretion and early Earth processes. The preservation of these differences in the present day mantle reflects the relatively low recycling efficiency of the noble gases with respect to the lithophile and siderophile elements. Importantly, since $^{129}$I became extinct 100 Myr after the start of the solar system, the reservoir supplying noble gases to the HIMU plume. Such a depleted MORB source would imply that the shallow upper mantle is the depleted MORB source. This is consistent with the conclusions inferred from the Xe data in the Iceland plume (Mukhopadhyay, 2012) and suggests that the initial phase of Earth’s accretion was volatile poor compared to the later stages of accretion because Pu is a refractory element while I is a volatile element.

5. Conclusions

We present new heavy noble gas data in MORBs from the equatorial Mid-Atlantic Ridge. Based on our observations we conclude:

1. He–Ne are tightly coupled to variations in the lithophile isotopes. The depleted end of the MORB array (D-DMM) is less radiogenic in He but more nucleogenic in Ne than N-MORBs (DMM). We conclude that N-MORBs in the equatorial Atlantic are produced by mixing a small proportion of a HIMU plume component to a very depleted MORB matrix.

2. Despite having more radiogenic Pb and He isotopic compositions, the HIMU-type MORB mantle source is less radiogenic in $^{21}$Ne/$^{22}$Ne and less radiogenic $^{40}$Ar/$^{36}$Ar and $^{129}$Xe/$^{130}$Xe ratios. D-DMM $^{129}$Xe/$^{130}$Xe ratios preclude air or seawater subduction as the sole cause of the lower $^{129}$Xe/$^{130}$Xe in the HIMU-type MORB source compared to the depleted MORB source; a less degassed component is required in the HIMU source. Consequently, the lower $^{21}$Ne/$^{22}$Ne, $^{40}$Ar/$^{36}$Ar and $^{129}$Xe/$^{130}$Xe are also interpreted as evidence for a less degassed mantle reservoir contributing to the HIMU plume. We emphasize that our results do not imply that air or seawater subduction do not occur; rather by itself it cannot account for the differences between the two groups of MORBs.

3. The ratio of Pu- to U-derived fission $^{136}$Xe in the depleted MORB source is ~0.4, suggesting it has lost ~99% of its Pu-derived Xe, and thus at least ~99% of its original primordial volatile inventory. The HIMU-type MORBs have a higher ratio of Pu- to U-derived fission Xe than the depleted MORB group, consistent with the sampling of a less degassed mantle reservoir in the HIMU plume.

4. Differences in $^{129}$Xe/$^{130}$Xe suggest a lower I/Pu ratio for the relatively undegassed mantle reservoir compared to the depleted MORB source and thus supports heterogeneous volatile accretion with early accretion being drier than later accreting material. Overall the Xe isotopic compositions require the preservation of early (pre-4.45 Ga) formed heterogeneities in the present day mantle.

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Appendix A. Supplementary materials

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.epsl.2012.08.025.

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