

Identification of chondritic krypton and xenon in Yellowstone gases and the timing of terrestrial volatile accretion

Michael W. Broadley^{a,1}, Peter H. Barry^b, David V. Bekaert^a, David J. Byrne^a, Antonio Caracausi^c, Christopher J. Ballentine^d, and Bernard Marty^a

^aCentre de Recherches Pétrographiques et Géochimiques, UMR 7358 CNRS—Université de Lorraine, BP 20, F-54501 Vandoeuvre-lès-Nancy, France; ^bMarine Chemistry and Geochemistry Department, Woods Hole Oceanographic Institution, Woods Hole, MA 02543; ^cInstituto Nazionale di Geofisica e Vulcanologia, 90146 Palermo, Italy; and ^dDepartment of Earth Sciences, University of Oxford, OX1 3AN Oxford, United Kingdom

Edited by Norman H. Sleep, Stanford University, Stanford, CA, and approved May 6, 2020 (received for review February 29, 2020)

Identifying the origin of noble gases in Earth's mantle can provide crucial constraints on the source and timing of volatile (C, N, H₂O, noble gases, etc.) delivery to Earth. It remains unclear whether the early Earth was able to directly capture and retain volatiles throughout accretion or whether it accreted anhydrously and subsequently acquired volatiles through later additions of chondritic material. Here, we report high-precision noble gas isotopic data from volcanic gases emanating from, in and around, the Yellowstone caldera (Wyoming, United States). We show that the He and Ne isotopic and elemental signatures of the Yellowstone gas requires an input from an undegassed mantle plume. Coupled with the distinct ratio of ¹²⁹Xe to primordial Xe isotopes in Yellowstone compared with mid-ocean ridge basalt (MORB) samples, this confirms that the deep plume and shallow MORB mantles have remained distinct from one another for the majority of Earth's history. Krypton and xenon isotopes in the Yellowstone mantle plume are found to be chondritic in origin, similar to the MORB source mantle. This is in contrast with the origin of neon in the mantle, which exhibits an isotopic dichotomy between solar plume and chondritic MORB mantle sources. The co-occurrence of solar and chondritic noble gases in the deep mantle is thought to reflect the heterogeneous nature of Earth's volatile accretion during the lifetime of the protosolar nebula. It notably implies that the Earth was able to retain its chondritic volatiles since its earliest stages of accretion, and not only through late additions.

origin of Earth's volatiles \mid accretion \mid mantle plume \mid noble gases \mid Yellowstone

Noble gas isotopes exhibit resolvable isotopic differences between chondritic (1), solar (2), and cometary reservoirs (3) and can be used to reveal important clues as to how the Earth acquired its early volatile budget (3-6). Differences in noble gas signatures between the deep plume and shallower mid-ocean ridge basalt (MORB) mantle sources (4, 5), as well as the atmosphere (3, 6), indicate that Earth experienced a heterogeneous history of volatile accretion. The highest ²⁰Ne/²²Ne ratio (13.03 ± 0.04) measured in plume-influenced samples is higher than that of solar wind-irradiated material (12.5 to 12.7) (7) and CI chondrites (8.2 ± 0.4) (8), therefore requiring the deep mantle to contain a nebular Ne component (13.36 \pm 0.18) (4, 9). This nebular Ne component was most likely acquired by dissolution of a nebular atmosphere into the mantle during periods of global magma ocean (10, 11). The protosolar nebula is a transient feature in the solar system considered to have dispersed ~4 My after the formation of the solar system (taken as the formation age of Calcium-Aluminum Inclusions) (12). The nebular noble gas component present in the deep mantle must therefore have been added early to the growing proto-Earth, within the lifetime of the protosolar nebula (4, 13). Despite there being several plume-influenced samples with ²⁰Ne/²²Ne values greater than solar wind-implanted material and CI chondrites (4, 13, 14), as of yet no plume-influenced sample has reached the nebular 20 Ne/ 22 Ne value. Therefore, the plume source mantle still requires the addition of a component with a 20 Ne/ 22 Ne lower than that of the solar nebula. It has been previously suggested that neon in the plume mantle source could represent a mixture between a nebular and solar wind-implanted component (14) or, alternatively, that the subduction of atmospheric neon in the mantle could have lowered the 20 Ne/ 22 Ne of the mantle over time (4)

While Ne in the deep mantle may have, at least in part, originated from the solar nebula, the ²⁰Ne/²²Ne of the upper mantle as measured in MORB (15) and continental well gases (5) is within the range of solar wind-irradiated material found in chondrites. This dichotomy in Ne isotopes between the depleted upper mantle and deep plume source mantle has been suggested to be the result of late additions of chondritic material to the upper mantle with an original solar-like composition (4). Preferential late additions of chondritic material to the Earth's surface and upper mantle may therefore have overprinted the original solar noble gas signature in the MORB reservoir, while the deep plume source mantle was able to better preserve its original solar noble gas signature (4).

Whether this apparent dichotomy in accretionary noble gas signatures can be extended to heavy noble gases (Kr and Xe)

Significance

Volatile elements play a critical role in the evolution of Earth. Nevertheless, the mechanism(s) by which Earth acquired, and was able to preserve its volatile budget throughout its violent accretionary history, remains uncertain. In this study, we analyzed noble gas isotopes in volcanic gases from the Yellowstone mantle plume, thought to sample the deep primordial mantle, to determine the origin of volatiles on Earth. We find that Kr and Xe isotopes within the deep mantle have a similar chondritic origin to those found previously in the upper mantle. This suggests that the Earth has retained chondritic volatiles throughout the accretion and, therefore, terrestrial volatiles cannot not solely be the result of late additions following the Moon-forming impact.

Author contributions: M.W.B., P.H.B., and B.M. designed research; M.W.B., D.V.B., D.J.B., A.C., C.J.B., and B.M. performed research; M.W.B., P.H.B., D.V.B., D.J.B., A.C., C.J.B., and B.M. analyzed data; and M.W.B. wrote the paper.

The authors declare no competing interest.

This article is a PNAS Direct Submission.

This open access article is distributed under Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND).

¹To whom correspondence may be addressed. Email: broadley@crpg.cnrs-nancy.fr.

This article contains supporting information online at https://www.pnas.org/lookup/suppl/doi:10.1073/pnas.2003907117/-/DCSupplemental.

First published June 8, 2020.

remains unknown. Krypton and xenon in the mantle are dominated by recycled atmosphere (16), rendering any evidence of accretionary signatures difficult to resolve. In order to circumvent this issue, large quantities of magmatic gas have been previously analyzed (6, 16, 17) to determine the primordial Kr and Xe signature of the mantle. Furthermore, a new protocol has been developed to accumulate mantle noble gases from highly vesicular basaltic MORB glass samples (18), which has enabled for the first time the primordial Kr and Xe signature from basaltic glass samples to be determined. The Kr and Xe isotopic data obtained from these studies supports a chondritic origin for heavy noble gases in the upper MORB source mantle (6, 18).

To date, no primordial heavy noble gas signatures have been identified in plume-influenced basaltic glass and olivine samples, with the light nonradiogenic ^{124,126,128}Xe isotopes being, for the most part, isotopically indistinguishable from atmosphere (19, 20). The lack of isotopic anomalies in these samples is most likely a function of the lower noble gas concentrations and therefore higher degree of atmospheric contamination compared with highly vesicular MORB glass (18). Recent attempts to determine the primordial heavy noble gas signature of the deep mantle have therefore been focused on the analysis of Xe in magmatic gases from the Eifel volcanic area in Germany (21). However, a recent reevaluation of Xe isotopes in Eifel has called into question whether it can be considered to originate from a deep, undegassed mantle source (17). Indeed, there is no supporting He isotope evidence, as ${}^{3}\text{He}/{}^{4}\text{He}$ values (<6 R_A, where R_A = 1.39 × 10⁻⁶, the ³He/⁴He ratio of air) in Eifel gas samples are lower than canonical MORB ($8 \pm 1 R_A$) values and Ne isotopes follow the same trajectory as the air-MORB mixing line (17). The accretionary origin of heavy noble gases (Kr and Xe) in the deep mantle is therefore yet to be satisfactorily resolved.

Determining the origin of heavy noble gases in the deep mantle is key to understanding whether the deep mantle accreted volatiles from a fundamentally different reservoir from that found in the upper MORB mantle source. A change from solar toward chondritic-dominated accretion would have important implications regarding the timing of volatile delivery to Earth, potentially indicating that volatiles in the upper mantle and at Earth's surface were added late in the accretionary process (22). In order to address these questions, we present highprecision noble gas isotope measurements in CO₂-rich magmatic gases (n = 13) collected from three localities: Mud Volcano, Turbid Lake, and Brimstone Basin (Fig. 1), within Yellowstone National Park, where geochemical (e.g., ³He/⁴He up to 15R_A) (23) and geophysical (24) data indicate a relatively undegassed, deep-seated source of volcanism.

Results and Discussion

The ³He/⁴He of samples analyzed here range between 2.3 and 16.3 R_A ($R_A = {}^{3}He/{}^{4}He$ of atmosphere), which is within the range of previously reported values for Yellowstone magmatic gases (23, 25). The large range of ³He/⁴He is attributed to assimilation of crustal-derived ⁴He from the surrounding Archean crust during magmatic gas migration (25). Assuming that magmatic gases originated from a mantle source with ³He/⁴He of 16.3 R_A (highest value measured in this study at Mud Volcano), then the crustal component would contribute up to 85% of the total He within the samples originating from Turbid Lake and Brimstone Basin (*SI Appendix*). Samples from Mud Volcano which exhibit high ³He/⁴He (>13 R_A), show a strong linear correlation for Ne isotopes, reflecting two-component mixing between the atmosphere and a high ²⁰Ne/²²Ne mantle component (Fig. 2). The steep slope of this mantle-air mixing line is characteristic of a mantle source enriched in primordial 22Ne relative to nucleogenic ²¹Ne. The trend defined by the samples is statistically distinct from the mass-dependent fractionation line, falling along a mixing line similar to that defined by samples

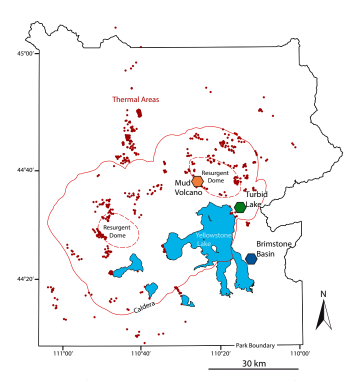


Fig. 1. Map of Yellowstone National Park. The locations of the three sampling sites that make up this study (Mud Volcano, Turbid Lake, and Brimstone Basin) are denoted as hexagons. Other thermal sites within the National Park are shown as red circles.

from the Loihi Seamount, Hawaii (26). The Mud Volcano sample with the highest measured ²⁰Ne/²²Ne ratio (Giggenbach sample 1B) is also offset from the mass-dependent fractionation line when 20 Ne/22 Ne is plotted against 38 År/36 Ar (SI Appendix, Fig. S5), indicating that measured Ne isotopic ratios are unlikely to be the result of mass fractionation and likely representative of the Yellowstone mantle source. This, taken together with high ³He/⁴He values, indicates the Yellowstone mantle source has preserved a high volatile to U+Th ratio and has therefore undergone less degassing of primordial noble gases compared with the MORB mantle source (30).

Samples from Brimstone Basin, which are characterized by the lowest ³He/⁴He, do not follow the same Ne isotopic trend as the other samples (Fig. 2). They exhibit higher ²¹Ne/²²Ne and plot near the MORB-air mixing line, potentially indicating that the Brimstone Basin samples contain a secondary component in addition to that sampled at Mud Volcano. Brimstone Basin is situated outside the Yellowstone Caldera (Fig. 2), where low ³He/⁴He have previously been attributed to the incorporation of Archean-aged crustal material, rich in radiogenic ⁴He, to an original high ³He/⁴He plume mantle source (25). This would also have the effect of increasing the ²¹Ne/²²Ne through the addition of crustal-derived nucleogenic ²¹Ne, likely accounting for why the Brimstone Basin samples are offset from the Ne isotopic trend defined by the Mud Volcano samples (Fig. 2).

The mantle noble gases from Brimstone Basin could also have originated from the MORB mantle reservoir, the subcontinental lithospheric mantle (SCLM), or a mixture between these reservoirs and the Yellowstone plume. However, the ³He/⁴He of Brimstone Basin (2.34 to 3.04 R_A) is lower than the average values for both MORB (8 \pm 1 R_A) (30) and SCLM (6 \pm 2 R_A) (31, 32) reservoirs, and, therefore, regardless of the original mantle source, the low ³He/⁴He measured in Brimstone Basin requires an additional crustal noble gas component enriched in radiogenic ⁴He (25). As the addition of crustal ⁴He should also

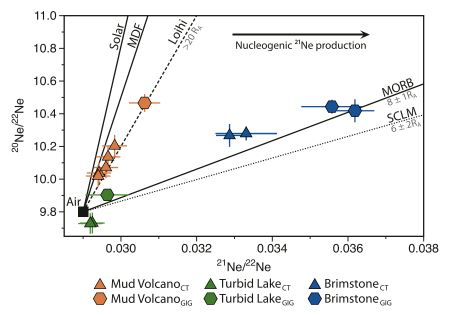


Fig. 2. Neon isotopic composition of Yellowstone volcanic gas. Data from samples collected (n = 13) at three localities are shown. Samples from Mud Volcano and Brimstone Basin show clear excesses from atmosphere. Gas collected in Giggenbach bottles (GIG) are less air-contaminated than samples collected in copper tubes (CT). Mud Volcano samples show a strong linear correlation with a slope approximate to that defined by the Loihi Seamount samples (26), confirming the Yellowstone plume samples a deep undegassed mantle reservoir. Brimstone Basin samples are offset toward the MORB–air mixing line (27) from the addition of crustal-derived nucleogenic 21 Ne, while samples from Turbid Lake are similar to atmosphere. The SCLM–air mixing line (28, 29) and mass-dependent fractionation line are shown for reference. Uncertainties are displayed at 1 σ .

be associated with the addition of nucleogenic ²¹Ne, the slope of the air—mantle mixing line that passes through the Brimstone Basin samples should represent a minimum, as the addition of ²¹Ne would serve to lower the steepness of the slope of the mixing line. Given that the Brimstone Basin samples plot along or slightly above the MORB—air mixing line in Fig. 2, this would rule out the SCLM as the primary mantle source for Brimstone Basin, despite the fact that volcanic products attributed to the Yellowstone plume are known to contain significant SCLM contributions (33). To reconcile the MORB-like Ne isotopes with the presumed high levels of crustal He present within Brimstone Basin requires the mantle source to contain a third, less-radiogenic component, which we surmise is the Yellowstone mantle plume component (as sampled at Mud Volcano).

To further determine whether the low ³He/⁴He Brimstone Basin and high ³He/⁴He Mud Volcano samples share a similar mantle component, we evaluate the relationship between primordial He and Ne isotopes in the samples. When ²⁰Ne/²²Ne is plotted against ³He/²²Ne (Fig. 3), all of the samples in excess of atmosphere, with the exception of one outlier, form a clear linear correlation. Crucially, the isotopes that form this correlation are not produced in significant quantities by radiogenic and/or nucleogenic processes and therefore provide information on the mantle source, without being influenced by crustal assimilation, as can be the case for ³He/⁴He and ²¹Ne/²²Ne. However, ³He/²²Ne can be affected by different physical processes such as degassing and dissolution, which can result in elemental fractionation altering the original mantle signature. If the ³He/²²Ne of Yellowstone has been modified, then this should also be apparent in other noble gas elemental ratios such as ⁴He/⁴⁰Ar¹*. The ⁴He/⁴⁰Ar* of the samples defining the He–Ne correlation in Fig. 3 (0.6 to 1.5) is similar to, if somewhat lower than, the mantle production ratio (1.6 to 4.2) (30), with the Turbid Lake and Brimstone Basin samples being corrected for crustal ⁴He and 40Ar contributions. The similarity between the measured ⁴He/⁴⁰Ar* in Yellowstone and the mantle production ratio

suggests that any elemental fractionation is likely to be minimal and the ${}^{3}\text{He}/{}^{22}\text{Ne}$ is likely representative of the Yellowstone mantle source (Fig. 3).

The linear correlation exhibited in Fig. 3 can therefore be considered to represent a mixing line between atmosphere and a common mantle component. Extrapolating the correlation line to the plume $^{20}\text{Ne/}^{22}\text{Ne}$ endmember value of ~ 13.4 (4) gives a $^{3}\text{He/}^{22}\text{Ne}$ ratio of 1.4 to 2.5 (Fig. 3). The $^{3}\text{He/}^{22}\text{Ne}$ of the Yellowstone mantle source therefore falls within the range previously published for plume-influenced samples (1.5 to 3) (14, 20) but is clearly distinct from MORB and SCLM values (>4.6) (34–36). Extrapolating to the MORB $^{20}\text{Ne/}^{22}\text{Ne}$ endmember (5) would result in an even lower $^{3}\text{He/}^{22}\text{Ne}$, therefore further distinguishing the Yellowstone mantle source from MORB. These observations suggest that the noble gases in Brimstone Basin and Mud Volcano originate from the same plume-derived mantle source and that the higher $^{21}\text{Ne/}^{22}\text{Ne}$ measured in Brimstone Basin is simply the consequence of higher crustal ^{21}Ne contributions (25).

The 40Ar/36Ar of the Brimstone Basin samples reach values greater than 1,400, which are the highest values yet measured in Yellowstone magmatic gas samples (37) and significantly higher than that measured in Mud Volcano and Turbid Lake (SI Appendix, Table S2). Importantly, crustal Ar contributions in the Brimstone Basin samples can be accounted for to derive the underlying mantle argon compositions (*SI Appendix*). Following correction for crustal 40 Ar* contribution, the 40 Ar/ 36 Ar of Brimstone Basin sample 4B is 1.001 ± 42 , which is still in excess of atmosphere. The corrected 40Ar/36Ar of Brimstone Basin remains higher than the other samples, suggesting that Brimstone Basin has the lowest degree of air contamination and greatest contribution of mantle-derived Ar. Unlike Mud Volcano and Turbid Lake, Brimstone Basin is not a thermal area but is still a significant degassing site (38). The low temperature of the Brimstone Basis system is considered to have limited boiling and therefore gas loss during transit toward the surface, potentially explaining why the diffusive magmatic CO₂ flux is so large for a

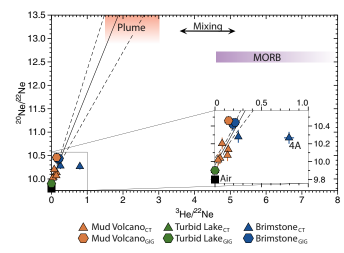


Fig. 3. ²⁰Ne/²²Ne versus ³He/²²Ne of Yellowstone volcanic gas. The He and Ne isotopic signature of the Yellowstone samples, with the exception of one outlier, are correlated, yielding a linear mixing array between atmosphere and the Yellowstone mantle source. Extrapolating the correlation line to the primitive mantle 20Ne/22Ne value of 13.4 gives the 3He/22Ne ratio of the Yellowstone mantle source to be between 1.4 and 2.5. A similarly low ³He/ ²²Ne for the Yellowstone mantle source (~3.6) is found following the method set out in ref. 34. The ³He/²²Ne of the Yellowstone mantle source is within the range previously determined for other plume influenced samples (1.5 to 3.0) (14, 20) but clearly distinct from that of MORB (>4.6) (34, 35). The higher 3He/22Ne ratio in the copper tube (CT) sample 4A is the result of He and Ne not being adequately cryoseparated during sample preparation. The Giggenbach (GIG) sample of the same gas falls along the same mixing line as the other samples, and we therefore do not consider the higher ³He/²²Ne measured in 4A to be representative of the mantle source. The dashed lines are the 1 σ confidence intervals fitted through all of the data minus the outlier sample 4A. Uncertainties on the measurements are displayed at 1 σ.

nonthermal site and why the noble gases are less contaminated by atmosphere. The greater mantle volatile contribution in Brimstone Basin has also been further substantiated by a recent innovative study that used the rare ¹⁵N¹⁵N isotopologue of N₂ to identify and correct for air contamination in volcanic gas emissions (39). This study revealed that not only did the Brimstone Basin samples contain the greatest contribution of mantlederived N₂ of all of the Yellowstone samples measured but also that the $\delta^{15}N$ of the Yellowstone mantle source is distinct from MORB and in line with previous determinations of plumederived samples (39).

The Brimstone Basin samples analyzed in this study were found to contain excesses in ¹²⁹Xe/¹³⁰Xe (Fig. 4) relative to atmosphere. Excesses in ¹²⁹Xe/¹³⁰Xe relative to atmosphere typically signify the addition of mantle-derived ¹²⁹Xe, produced from the decay of short-lived radiogenic ¹²⁹I, therefore further confirming the lower degree of atmospheric contamination in this sample. Assuming the Yellowstone magmatic source originated with a similar $^{129}\text{Xe}/^{130}\text{Xe}$ to the Icelandic plume source (6.98 \pm 0.07) (20), we calculate that Brimstone Basin sample 4B, which has the highest measured 129 Xe/ 130 Xe (6.541 \pm 0.002), contains between 7 and 12% mantle-derived Xe, with the remainder being from atmospheric contamination occurring at the near surface. These results reveal an unambiguous mantle-derived Xe anomaly within the Yellowstone mantle source.

Brimstone Basin sample 4B shows resolvable excesses in the heavy fissiogenic, and light nonradiogenic, Xe isotopes (Fig. 4). Heavy Xe isotopes ($^{131-136}$ Xe) are produced in different proportions from the spontaneous fission of extinct 244 Pu ($T_{1/2} = 80$ My) and extant 238 U ($T_{1/2} = 4.469$ Gy) (40). Undegassed mantle reservoirs are typically enriched in Pu-derived Xe relative to U, with the ratio of Pu/U-derived Xe utilized to infer the degassing

state of different mantle reservoirs (20). However, measured excesses in fissiogenic Xe isotopes in Brimstone Basin sample 4B are higher than would be expected for a mantle-derived sample, containing up to 12% mantle-derived Xe (Fig. 4). The fissiogenic Xe signature in Brimstone Basin therefore contains an additional Xe component produced from the decay of ²³⁸U in the surrounding Archean crust (SI Appendix, Fig. S9), which precludes using the fissiogenic Xe isotopes to gain further insights into the Yellowstone mantle source.

No significant production pathways exist for primordial 124 Xe, 126 Xe, 128 Xe, and 130 Xe within the mantle. These isotopes are considered to be a remnant of accretionary volatiles preserved since Earth's formation. The primordial Xe isotopes measured within the Brimstone Basin samples can therefore be used to make further inferences on the nature and composition of the Yellowstone mantle source. In Fig. 5, we show that the Yellowstone mantle source exhibits different ¹²⁹Xe/^{124,126,128}Xe ratios from that measured in MORB-derived samples (16-18). Although the difference is slight, the Brimstone Basin sample 4B has lower ¹²⁹Xe/^{124,126,128}Xe ratios than any previously measured MORB-derived sample. The lower ¹²⁹Xe/^{124,126,128}Xe measured in Yellowstone compared with MORB provides further evidence that the Yellowstone gas is unlikely to be sourced from a mantle reservoir with a MORB-like Xe isotopic signature.

Extrapolating the air-Yellowstone mixing line in Fig. 5 to the Extrapolating the all–1 chowstonic mixing line in Fig. 5 to the $^{129}\text{Xe}/^{130}\text{Xe}$ plume mantle endmember, as defined by the Iceland mantle source (6.98 ± 0.07) (20), enables the $^{124}\text{Xe}/^{130}\text{Xe}$, $^{126}\text{Xe}/^{130}$ Xe, and $^{128}\text{Xe}/^{130}\text{Xe}$ of the Yellowstone mantle source to be estimated. The extrapolated $^{124}\text{Xe}/^{130}\text{Xe}$, and $^{129}\text{Xe}/^{130}$ Xe, and ¹²⁸Xe/¹³⁰Xe ratios of the Yellowstone mantle source are shown to be generally higher than the extrapolated values determined for the MORB mantle source. The extrapolated 124 Xe/ 130 Xe, 126 Xe/ 130 Xe, and 128 Xe/ 130 Xe ratios suggest that the Yellowstone

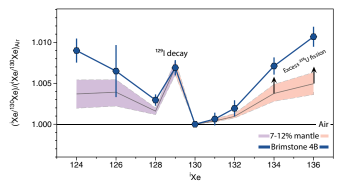


Fig. 4. Xenon isotopic spectrum of Yellowstone volcanic gas (sample 4B). Data are normalized to the isotopic composition of atmosphere and ¹³⁰Xe. Sample 4B is the least air-contaminated sample from Brimstone Basin. It shows resolvable ¹²⁹Xe/¹³⁰Xe excesses, confirming the contribution from mantle-derived xenon. Comparing the ¹²⁹Xe/¹³⁰Xe with that of the Icelandic plume source, we calculate the amount of mantle-derived Xe present within the sample to be between 7 and 12%. The Xe isotopic spectrum corresponding to this estimated mantle contribution is shown by the shaded area (SI Appendix). The estimated heavy isotope composition (orange shaded area) is estimated relative to the Icelandic plume mantle source (20), while the expected light xenon isotope signature (purple shaded area) is calculated based on the amount of primordial Xe present within the MORB source, as no previous estimates for the plume source mantle exist for light Xe isotopes. The excesses in heavy xenon isotopes in the sample are greater than expected for mantle-derived gases, indicating the sample contains an excess fissiogenic Xe component inherited from the surrounding Archean crust. The ¹²⁴Xe/¹³⁰Xe, ¹²⁶Xe/¹³⁰Xe, and ¹²⁸Xe/¹³⁰Xe ratios measured in Yellowstone are higher than would be expected from a MORB source given the excess in ¹²⁹Xe/¹³⁰Xe, indicating that Yellowstone volcanism originates from mantle reservoir with a lower ¹²⁹Xe/^{Primordial}Xe than MORB. Uncertainties represent 1 SE.

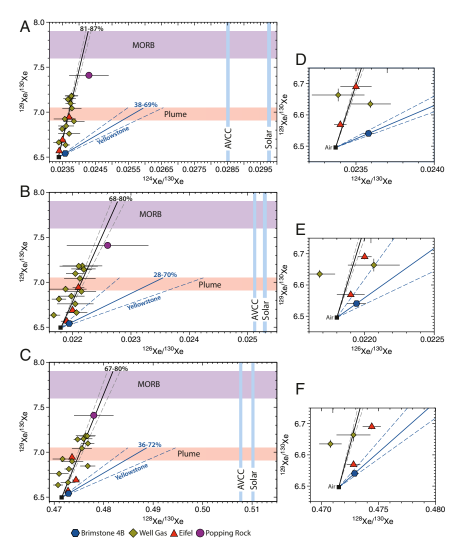


Fig. 5. Difference in Xe isotopic composition between the Yellowstone mantle source and MORB. (*A*–*C*) The ¹²⁹Xe/¹³⁰Xe versus ¹²⁴Xe/¹³⁰Xe (*A*), ¹²⁶Xe/¹³⁰Xe (*B*), and ¹²⁸Xe/¹³⁰Xe (*C*) of Brimstone Basin 4B is plotted together with MORB-derived samples (16–18). *D*–*F* represent zoomed in sections of figures *A*–*C*, respectively. The solid lines in these plots represent the mixing line between atmosphere and the mantle endmember composition. The Brimstone Basin sample has consistently lower ¹²⁹Xe/¹²⁴, ¹²⁶, ¹²⁸Xe ratios (defined from the slope of the mixing lines) when compared with MORB, suggesting that it originated from a mantle source with a lower I/Xe. The endmember ¹²⁹Xe/¹³⁰Xe composition for the MORB (16–18) and plume (20) mantle sources are shown for comparison. Extrapolated ¹²⁴Xe/¹³⁰Xe, and ¹²⁸Xe/¹³⁰Xe values for Yellowstone are higher and closer to primordial AVCC (41) and solar values (2) than that calculated for the MORB source. The percentage of recycled atmosphere, relative to an AVCC starting composition, in the Yellowstone and MORB mantle sources are stated. The solid black line through the MORB data is an error-weighted fit forced through the atmospheric composition, with the dashed lines representing the 1 σ confidence interval. The Yellowstone correlation (solid blue line) is fitted through Brimstone Basin sample 4B, with the dashed lines representing the maximum extent of the uncertainty of the sample. Uncertainties for sample 4B represent 1 SE and are smaller than symbol size.

mantle source has a slightly higher ratio of primordial (Average Carbonaceous Chondrites [AVCC]) (41) to recycled atmospheric Xe than the MORB mantle source (Fig. 5). However, with the exception of ¹²⁴Xe/¹³⁰Xe, the estimated amount of recycled atmospheric Xe isotopic in the Yellowstone plume and MORB mantle sources are within uncertainty of each other (Fig. 5), as has been previously suggested (42).

(Fig. 5), as has been previously suggested (42). These estimates are dependent on the ¹²⁹Xe/¹³⁰Xe ratios chosen as the mantle endmember, so if, for instance, the ¹²⁹Xe/¹³⁰Xe endmember for Yellowstone was greater than that determined for the Iceland mantle source, and more similar to MORB, then the ratio of primordial to recycled Xe within the Yellowstone mantle source would be even higher (Fig. 5). Despite the large uncertainties, the Yellowstone mantle source appears to contain similar, if not slightly lower, proportions of recycled atmospheric Xe compared with the MORB mantle

source. Comparable estimates for the amount of recycled Xe in the plume and MORB mantle sources (42), despite the plume source being less degassed of its volatiles, suggest that volatiles may be more efficiently recycled to the deep plume source mantle.

Although it is difficult to reach definitive conclusions based on a single determination of light isotopes in plume-influenced samples, we suggest that to account for both the high proportion of recycled Xe and the lower ¹²⁹Xe/^{124,126,128}Xe of the Yellowstone samples requires that the Yellowstone mantle source have a lower ¹²⁹Xe/¹³⁰Xe, similar to that measured in Iceland (Fig. 5 and ref. 20). These observations confirm that the plume and MORB mantle reservoirs evolved with different ratios of ¹²⁹Xe to primordial Xe isotopes (20). Since primordial Xe isotopes did not change during ¹²⁹Xe production from the decay of extinct ¹²⁹I over the first 100 My of Earth's history, the differences between the two mantle reservoirs are a function of

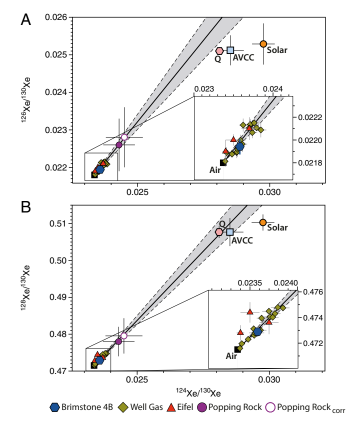


Fig. 6. Primordial light xenon isotopes in the Yellowstone mantle source. The ¹²⁶Xe/¹³⁰Xe (A) and ¹²⁸Xe/¹³⁰Xe (B) versus ¹²⁴Xe/¹³⁰Xe of Brimstone Basin gases plot along the same trend fitted through volcanic gas from Eifel (17), upper magmatic CO2 well gases (16), the measured value and the endmember composition, corrected for atmospheric contamination, of MORB popping rock (18), therefore confirming the presence of primordial xenon in the Yellowstone mantle source. Phase Q (1), AVCC (41), and Solar Wind (2) compositions are shown for reference. Uncertainties for sample 4B represent 1 SE.

their different initial I/Xe ratios (20). The Yellowstone mantle source, as well as that feeding other mantle plumes (20, 43), must therefore have been separated from the MORB source mantle early in Earth's history (before 4.5 Ga), with limited mixing occurring between the two reservoirs since.

The plume origin of the noble gases from Brimstone Basin (Figs. 2, 3, and 5), coupled with their resolvable light Xe isotope excesses from atmosphere, presents an opportunity to determine the accretionary origin of the heavy noble gases in a plume-influenced mantle source. The $^{124}\mathrm{Xe/^{130}Xe},~^{126}\mathrm{Xe/^{130}Xe},$ and ¹²⁸Xe/¹³⁰Xe values plot along the same trend defined previously from the analysis of MORB-derived samples, which is considered to represent a mixing line between air and chondritic Xe: Phase Q, main carrier of chondritic noble gases (1), and AVCC (41). This is, in itself, not sufficient to conclude whether the Yellowstone mantle source is dominated by chondritic or solar noble gases, given the small measured isotope excess and the relatively minor difference between chondritic and solar Xe isotopic signatures (Fig. 6).

Unlike Xe, the Kr isotopic signatures of chondritic and solar endmembers are isotopically heavier and lighter than atmosphere, respectively, therefore constituting a powerful tool for distinguishing between chondritic and solar volatile sources in the mantle (6). However, resolving any mantle Kr contributions has proven difficult, given the relatively small isotopic difference, in comparison with Xe, between the atmosphere and potential accretionary sources (solar and chondritic). Despite this, recent high-precision studies of CO₂ well gas and MORB popping rock have shown the upper mantle to contain a primordial chondritic Kr signature (6, 18).

After correcting for ⁸⁴Kr and ⁸⁶Kr produced by U fission in the crust (SI Appendix), we find that Brimstone Basin sample 4B maintains a ⁸⁶Kr/⁸⁴Kr excess relative to atmosphere, indicating a resolvable primordial mantle Kr component (Fig. 7). The nonatmospheric Kr signature measured within Yellowstone magmatic gas provides a unique insight into the origin of volatiles in the deep plume mantle source. Fig. 7 demonstrates that Kr and Xe isotopes from Brimstone Basin fall along the air-mantle mixing line defined by magmatic CO₂ well gases containing a chondritic inherited noble gas component (6). The identification of chondritic Kr is primarily based on the ⁸⁶Kr/⁸⁴Kr ratio, as the difference between the 82Kr/84Kr and 83Kr/84Kr composition of chondrites and the atmosphere is smaller and therefore more difficult to resolve from the atmosphere (SI Appendix, Fig. S4). Critically, the heavy noble gas composition of the Yellowstone and MORB mantle sources appears to be dominated by chondritic sources, with no evidence for the preservation of solar Kr and Xe isotopes in the mantle.

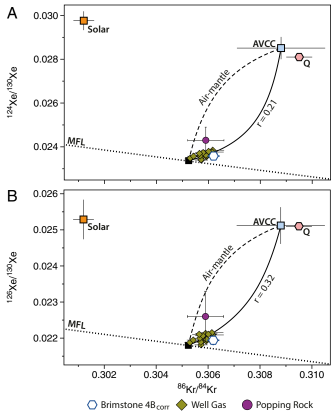


Fig. 7. Chondritic krypton and xenon isotopes in the deep mantle. Brimstone Basin falls along the same trend as magmatic CO2 well gases (6) in 124 Xe/ 130 Xe (A) and 126 Xe/ 130 Xe (B) versus 86 Kr/ 84 Kr space. Magmatic CO₂ well gas and Yellowstone samples have been corrected for crustal U fission production of 86Kr and 84Kr. The curvature of the mixing hyperbola between air and the original mantle composition is defined by [130Xe/84Kr]_{mantle}/[130Xe/ ⁸⁴Kr]_{Air} and is calculated performing a total least-square hyperbolic fit (*SI* Appendix). The trajectory of the mantle-air mixing line for both the magmatic CO₂ well gas and Yellowstone samples indicates that both the plume and MORB mantle sources are dominated by chondritic—and not solar—Kr and Xe. The mass-dependent fractionation line and values for Phase Q (1), AVCC (41), Solar Wind (2), and MORB Popping Rock (18) are shown for reference. Uncertainties of Brimstone 4B represent 1 SE.

The identification of chondritic Kr and Xe in the deep plume mantle source is therefore in line with the chondritic origin of noble gases in the MORB mantle but in contrast with the convincing evidence for solar neon in plume-influenced samples (4). To account for the Ne isotopic dichotomy between the plume and MORB mantle reservoirs, it has been suggested that the MORB mantle originated from the plume mantle but was subsequently modified by additions of chondritic material following the dispersion of the solar nebula (4). This scenario is based on the fact that, in ²⁰Ne/²²Ne versus ³⁶Ar/²²Ne or ¹³⁰Xe/²²Ne space, plume-influenced basalts and MORBs lie along a mixing line between the solar nebula composition and a chondritic component similar to CI chondrites and/or ocean water values. Both mantle reservoirs would hence have originated with a solar noble gas composition, before being modified to variable extents by the latter addition of chondritic material (4).

To account for the chondritic Kr and Xe signature in the deep plume mantle, while preserving the solar Ne signature, therefore requires a mechanism that could overprint solar heavy noble gases without affecting solar light noble gases. Two main scenarios have been proposed to date to account for the origin of chondritic noble gases in Earth's mantle: 1) introduction of chondritic noble gases into the mantle through time via subduction of surface material and/or 2) direct injection of chondritic material to the growing Earth following the dissipation of the solar nebula, with impacting chondritic bodies preferentially populating the MORB mantle relative to the plume mantle (4).

The subduction of chondritic volatiles from Earth's surface could potentially introduce chondritic Kr and Xe to the plume source, considering the efficient recycling of Ar, Kr, and Xe (relative to Ne) to both the deep plume and MORB mantle sources (16, 42). However, subduction of atmospheric Xe to the mantle could not account for the chondritic Xe signature in the lower mantle. This is because the ancient atmospheric Xe component (U–Xe) was not solely chondritic but also contained up to 27% cometary Xe (3). The addition of cometary noble gases appears to be limited to the atmosphere (3), with no evidence of a U–Xe signature in the present-day mantle (18, 21). Furthermore, the efficient recycling of atmospheric Xe is not thought to commence until the atmosphere reached a modern isotopic composition, some 2.5 Ga (18, 44).

The second scenario, concerning the direct addition of chondritic material to the mantle, could be related to the purported late veneer, whereby the addition of $\sim 0.5\%$ of the Earth's mass was added following core formation and the moon-forming impact (45). A late veneer in the form of carbonaceous chondrite-like material could potentially have supplied the majority of Earth's volatiles (22). However, differences in ¹²⁹Xe/¹²⁴,¹²⁶,¹²⁸,¹³⁰Xe between plume (Fig. 5) and MORB mantle reservoirs (20) require that the two reservoirs have experienced limited mixing over the last 4.45 Gy, including during the Moon-forming impact. Furthermore, Archean-aged komatiites that sample the ancient deep mantle are depleted in siderophile elements relative to the modern mantle (46), indicating that siderophile elements added to the Earth during the late veneer were not directly injected into the deep mantle but were progressively mixed in through time via mantle convection (47). The chondritic heavy noble gas component in the deep plume mantle source is therefore unlikely to have been introduced via top-to-bottom mixing of chondritic material arriving to Earth during the late veneer.

Further constraints on the timing of volatile accretion can be provided by examining differences in ${}^{3}\text{He}/{}^{22}\text{Ne}$ between the plume and MORB mantle reservoirs (Fig. 3). During ingassing of a gravitationally captured nebular atmosphere, He and Ne are expected to elementally fractionate in accordance with their different solubilities (34, 48). This results in the ${}^{3}\text{He}/{}^{22}\text{Ne}$ of the mantle rising from the nebular value of 1.5 to \sim 2 to 3 (similar to values observed for primitive plumes; Fig. 3 and ref. 34). It has

been suggested that, in order to increase the ³He/²²Ne from the primitive plume reservoir value to that of the MORB mantle (6.6 ± 2.0) would require at least two further episodes of giant impact-driven atmospheric loss and magma ocean outgassing (34). The preservation of low ³He/²²Ne within the deep mantle reservoir (as sampled here by the Yellowstone plume, with ³He/ ²²Ne of 1.4 to 2.5) therefore indicates that it has remained at least partially isolated following the ingassing of the protosolar nebula and that subsequent giant impacts did not significantly affect its volatile element budget. This is consistent with the observed correlation between ¹⁸²W anomalies and ³He/⁴He in mantle plumes, which suggests that the plume mantle reservoir and its noble gas budget were generated no later than 60 My after the formation of the Solar System (49). We therefore conclude from the complementary constraints brought from ³He/ ²²Ne and Xe isotope systematics that the chondritic heavy noble gases in the Yellowstone plume mantle, at the very least, predate the Moon-forming impact and are unrelated to the chondritic component in the MORB mantle.

The existence of solar Ne in the plume mantle requires that the Earth grew to be Mars-sized within the lifetime of the solar nebula (4). The Mars-sized proto-Earth was likely to have retained chondritic volatiles, as has been shown to be the case for other early accreting planetary embryos (50), prior to the ingassing of a nebular atmosphere. Ingassing of nebular volatiles to Earth's mantle would then preferentially alter the isotopic signature of Ne, relative to Kr and Xe (10, 11), due to the much higher ²⁰Ne/¹³⁰Xe ratio of the solar nebula compared with CI chondrites (8, 9) and greater solubility of Ne in basaltic melts in comparison with Xe (48). We calculate that to raise the ²⁰Ne/ ²²Ne of the mantle from CI chondrites-like values (9.03 \pm 2.46) to its estimated primordial value of 13.23 ± 0.22 (4) would result in a negligible contribution of solar Kr (2%) and Xe (<1%) to the mantle inventory (*SI Appendix*, Fig. S10). These estimates would be even lower if the starting ²⁰Ne/²²Ne composition of the mantle was taken to be the chondritic Ne-B endmember (~ 12.7) (7).

The preservation of primordial chondritic Kr and Xe in the deep Yellowstone mantle source has broad implications regarding the origin and timing of volatile element delivery to Earth. It notably indicates that the Earth has been retentive to chondritic volatiles throughout accretion and that, contrary to the commonly held view, terrestrial volatiles were not solely inherited from Earth's latest stages of accretion (Moon-forming impact and late veneer). Our results imply that the dichotomy between the main sources of heavy (chondritic) and light (solar) noble gases in the deep mantle was already established within the proto-Earth's mantle during the lifetime of the protosolar nebula.

Materials and Methods

Samples were collected in August 2018 from three localities within Yellowstone National Park: Mud Volcano, Turbid Lake, and Brimstone Basin (Fig. 1 and SI Appendix, Table S1). The sites consisted of both hot (Mud Volcano [85 °C], Turbid Lake [80 °C]) and cold (Brimstone Basin [21 °C]) degassing features, which were specifically chosen as they previously showed 40Ar/36Ar values in excess of air (38), thus providing the highest probability of finding mantle-derived Kr and Xe signatures. Gas samples were collected in both refrigeration-grade copper tubes and Giggenbach bottles (i.e., preevacuated 200-cm³ glass flasks filled with 50 to 80 cm³ of 7N NaOH solution) (51). Prior to the samples being sealed, volcanic gas was flushed through the Tygon tubing and copper tubes for a minimum of 30 min to limit air contamination. Sample 4B from Brimstone Basin was flushed with gas overnight for a minimum of 12 h. Giggenbach samples were analyzed for He, Ne, Ar, and high-precision Kr and Xe (SI Appendix, Tables S3 and S4) isotopic analysis at the Centre de Recherches Pétrographiques et Géochimiques (CRPG) noble gas analytical facility, using the ThermoFisher Scientific Helix SFT for helium and Helix MC+ for neon, argon, krypton, and xenon (13). Noble gas concentrations and He, Ne, and Ar isotopes were measured in the copper tube samples in the Noble Laboratory at the University of Oxford, using a dual mass-spectrometer setup (52) (S/ Appendix).

Data Availability. All data pertaining to this study can be found in SI Appendix, Tables S1-S4.

ACKNOWLEDGMENTS. Samples were collected as part of Study YELL-08056: Xenon Anomalies in the Yellowstone Hotspot. We thank Annie Carlson and all of the rangers at the Yellowstone National Park for providing invaluable

- 1. H. Busemann, H. Baur, R. Wieler, Primordial noble gases in "phase Q" in carbonaceous and ordinary chondrites studied by closed-system stepped etching. Meteorit. Planet. Sci. 35, 949-973 (2000).
- 2. A. Meshik, C. Hohenberg, O. Pravdivtseva, D. Burnett, Heavy noble gases in solar wind delivered by Genesis mission. Geochim. Cosmochim. Acta 127, 326-347 (2014).
- 3. B. Marty et al., Xenon isotopes in 67P/Churyumov-Gerasimenko show that comets contributed to Earth's atmosphere. Science 356, 1069-1072 (2017).
- 4. C. D. Williams, S. Mukhopadhyay, Capture of nebular gases during Earth's accretion is preserved in deep-mantle neon. Nature 565, 78-81 (2019).
- 5. C. J. Ballentine, B. Marty, B. Sherwood Lollar, M. Cassidy, Neon isotopes constrain convection and volatile origin in the Earth's mantle. Nature 433, 33–38 (2005)
- 6. G. Holland, M. Cassidy, C. J. Ballentine, Meteorite Kr in Earth's mantle suggests a late accretionary source for the atmosphere. Science 326, 1522-1525 (2009).
- 7. M. Moreira, S. Charnoz, The origin of the neon isotopes in chondrites and on Earth. Earth Planet. Sci. Lett. 433, 249-256 (2016).
- 8. E. Mazor, D. Heymann, E. Anders, Noble gases in carbonaceous chondrites. Geochim. Cosmochim. Acta 34, 781-824 (1970).
- V. S. Heber et al., Noble gas composition of the solar wind as collected by the Genesis mission. Geochim. Cosmochim. Acta 73, 7414-7432 (2009).
- 10. H. Mizuno, K. Nakazawa, C. Hayashi, Dissolution of the primordial rare gases into the molten Earth's material. Earth Planet. Sci. Lett. 50, 202-210 (1980).
- 11. C. L. Harper, S. B. Jacobsen, Noble gases and Earth's accretion. Science 273, 1814–1818
- 12. H. Wang et al., Lifetime of the solar nebula constrained by meteorite paleomagnetism. Science 355, 623-627 (2017).
- 13. R. Yokochi, B. Marty, A determination of the neon isotopic composition of the deep mantle. Earth Planet. Sci. Lett. 225, 77-88 (2004).
- 14. S. Péron et al., Neon isotopic composition of the mantle constrained by single vesicle analyses, Earth Planet, Sci. Lett. 449, 145-154 (2016).
- 15. A. Raquin, M. Moreira, Atmospheric ³⁸Ar/³⁶Ar in the mantle: Implications for the nature of the terrestrial parent bodies. Earth Planet. Sci. Lett. 287, 551-558 (2009).
- 16. G. Holland, C. J. Ballentine, Seawater subduction controls the heavy noble gas composition of the mantle. Nature 441, 186-191 (2006).
- 17. D. V. Bekaert, M. W. Broadley, A. Caracausi, B. Marty, Novel insights into the degassing history of Earth's mantle from high precision noble gas analysis of magmatic gas. Earth Planet. Sci. Lett. 525, 115766 (2019).
- 18. S. Péron, M. Moreira, Onset of volatile recycling into the mantle determined by xenon anomalies. Geochem. Perspect. Lett. 9, 21-25 (2018).
- 19. M. Trieloff, J. Kunz, D. A. Clague, D. Harrison, C. J. Allegre, The nature of pristine noble gases in mantle plumes. Science 288, 1036-1039 (2000).
- 20. S. Mukhopadhyay, Early differentiation and volatile accretion recorded in deepmantle neon and xenon. Nature 486, 101-104 (2012).
- 21. A. Caracausi, G. Avice, P. G. Burnard, E. Füri, B. Marty, Chondritic xenon in the Earth's mantle. Nature 533, 82-85 (2016).
- 22. F. Albarède, Volatile accretion history of the terrestrial planets and dynamic implications, Nature 461, 1227-1233 (2009).
- 23. H. Craig, J. E. Lupton, J. A. Welhan, R. Poreda, Helium isotope ratios in Yellowstone and Lassen Park volcanic gases. Geophys. Res. Lett. 5, 897-900 (1978).
- 24. P. L. Nelson, S. P. Grand, Lower-mantle plume beneath the Yellowstone hotspot revealed by core waves. Nat. Geosci. 11, 280-284 (2018).
- 25. J. B. Lowenstern, W. C. Evans, D. Bergfeld, A. G. Hunt, Prodigious degassing of a billion years of accumulated radiogenic helium at Yellowstone. Nature 506, 355–358 (2014).
- 26. P. J. Valbracht, T. Staudacher, A. Malahoff, C. J. Allègre, Noble gas systematics of deep rift zone glasses from Loihi Seamount, Hawaii. Earth Planet. Sci. Lett. 150, 399-411
- 27. P. Sarda, T. Staudacher, C. J. Allègre, Neon isotopes in submarine basalts. Earth Planet. Sci. Lett. 91, 73-88 (1988).
- 28. A. Buikin et al., Noble gas isotopes suggest deep mantle plume source of late Cenozoic mafic alkaline volcanism in Europe. Earth Planet. Sci. Lett. 230, 143-162 (2005).

advice and help when collecting the samples. M.W.B., D.V.B., D.J.B., and B.M. were supported by the European Research Council (PHOTONIS Project Grant 695618). This work was partially supported by Grants G-2016-7206 and G-2017-9696 from the Alfred P. Sloan Foundation and the Deep Carbon Observatory (to P.H.B.) and UK National Environment Research Council Deep Volatile Grant NE/M000427/1 (to C.J.B.). We also thank Laurent Zimmerman for providing help with the analysis. Finally, we thank the editor for efficient handling of our manuscript and the two anonymous reviewers for their insightful comments. This is CRPG contribution 2998.

- 29. J. Hopp, M. Trieloff, R. Altherr, Neon isotopes in mantle rocks from the Red Sea region reveal large-scale plume-lithosphere interaction. Earth Planet. Sci. Lett. 219, 61-76 (2004).
- 30. D. W. Graham, Noble gas isotope geochemistry of mid-ocean ridge and ocean island basalts: Characterization of mantle source reservoirs. Rev. Mineral. Geochem. 47, 247-317 (2002).
- 31. C. Gautheron, M. Moreira, Helium signature of the subcontinental lithospheric mantle. Earth Planet. Sci. Lett. 199, 39-47 (2002).
- 32. J. M. Day et al., The helium flux from the continents and ubiquity of low 3He/4He recycled crust and lithosphere. Geochim. Cosmochim. Acta 153, 116-133 (2015).
- 33. M. M. Jean, B. B. Hanan, J. W. Shervais, Yellowstone hotspot-continental lithosphere interaction. Earth Planet. Sci. Lett. 389, 119-131 (2014).
- 34. J. M. Tucker, S. Mukhopadhyay, Evidence for multiple magma ocean outgassing and atmospheric loss episodes from mantle noble gases. Earth Planet. Sci. Lett. 393, 254-265 (2014).
- 35. J. M. Tucker, S. Mukhopadhyay, H. M. Gonnermann, Reconstructing mantle carbon and noble gas contents from degassed mid-ocean ridge basalts. Earth Planet. Sci. Lett. 496, 108-119 (2018).
- 36. N. Dygert et al., Plate tectonic cycling modulates Earth's ³He/²²Ne ratio. Earth Planet. Sci. Lett. 498, 309-321 (2018).
- 37. J. B. Bergfeld, A. G. Hunt, W. Shanks, W. Evans, "Gas and isotope chemistry of thermal features in Yellowstone National Park, Wyoming" (Scientific Investigations Report 2011-5012, US Geological Survey, 2011).
- 38. D. Bergfeld, W. C. Evans, J. B. Lowenstern, S. Hurwitz, Carbon dioxide and hydrogen sulfide degassing and cryptic thermal input to Brimstone Basin, Yellowstone National Park, Wyoming. Chem. Geol. 330, 233-243 (2012).
- 39. J. Labidi et al., Hydrothermal ¹⁵N¹⁵N abundances constrain the origins of mantle nitrogen. Nature 580, 367-371 (2020).
- 40. C. J. Ballentine, P. G. Burnard, Production, release and transport of noble gases in the continental crust. Rev. Mineral. Geochem. 47, 481-538 (2002)
- 41. R. O. Pepin, On noble gas processing in the solar accretion disk. Space Sci. Rev. 106, 211-230 (2003).
- 42. R. Parai, S. Mukhopadhyay, The evolution of MORB and plume mantle volatile budgets: Constraints from fission Xe isotopes in southwest Indian ridge basalts. Geochem. Geophys. Geosyst. 16, 719-735 (2015).
- 43. M. K. Pető, S. Mukhopadhyay, K. A. Kelley, Heterogeneities from the first 100 million years recorded in deep mantle noble gases from the Northern Lau Back-arc Basin. Earth Planet. Sci. Lett. 369, 13-23 (2013).
- 44. R. Parai, S. Mukhopadhyay, Xenon isotopic constraints on the history of volatile recycling into the mantle. Nature 560, 223-227 (2018).
- 45. T. S. Kruijer, T. Kleine, M. Fischer-Gödde, P. Sprung, Lunar tungsten isotopic evidence for the late veneer, Nature 520, 534-537 (2015).
- 46. S. Marchi, R. M. Canup, R. J. Walker, Heterogeneous delivery of silicate and metal to the Earth by large planetesimals. Nat. Geosci. 11, 77-81 (2018).
- 47. W. D. Maier, Progressive mixing of meteoritic veneer into the early Earth's deep mantle. Nature 460, 620-623 (2009).
- 48. G. Lux, The behavior of noble gases in silicate liquids: Solution, diffusion, bubbles and surface effects, with applications to natural samples. Geochim. Cosmochim. Acta 51, 1549-1560 (1987).
- 49. A. Mundl et al., Tungsten-182 heterogeneity in modern ocean island basalts. Science 356, 66-69 (2017).
- 50. M. W. Broadley, D. V. Bekaert, B. Marty, A. Yamaguchi, J. A. Barrat, Noble gas variations in ureilites and their implications for ureilite parent body formation. Geochim. Cosmochim. Acta 270, 325-337 (2020).
- 51. W. F. Giggenbach, R. L. Goguel, Methods for the Collection and Analysis of Geothermal and Volcanic Water and Gas Samples, (Chemistry Division, Department of Scientific and Industrial Research, 1988).
- 52. P. H. Barry et al., Noble gases solubility models of hydrocarbon charge mechanism in the Sleipner Vest gas field. Geochim. Cosmochim. Acta 194, 291-309 (2016).