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The Xenon Record of Extinct Radioactivities in the Earth

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Abstract. Analyses of xenon from well gas rich in carbon dioxide reveal a large excess of radiogenic xenon-129 from the decay of extinct iodine-129. Smaller excesses observed in the heavy xenon isotopes are from fission. These results place narrow limits on any age difference between the earth and the oldest meteorites. The occurrence of excess radiogenic xenon-129 in well gas also suggests that any quantitative degassing of existing solid materials to form the atmosphere must have been limited to a very early period of the earth's history, approximately the first 10⁸ years. Alternatively, this observation is consistent with a model of the earth's continuous, but still incomplete, degassing since its time of formation.

There is unambiguous evidence from several radioactive dating methods that most meteorites formed about 4.6×10^9 years ago (1, 2). This age is obtained by comparing present concentrations of certain long-lived radioactive nuclides, ²³⁸U, ²³⁵U, ⁸⁷Rb, and ⁴⁰K, with the concentrations of their respective stable decay products, ²⁰⁶Pb, ²⁰⁷Pb, ⁸⁷Sr, and ⁴⁰Ar. It has recently been established that two radioactive nuclides, ¹²⁹I and ²⁴⁴Pu, were incorporated into meteorites at the time of their formation (3), but essentially the entire inventory of these two nuclides has decayed over the life-span of 4.6×10^9 years of meteorites. These two nuclides are now referred to as extinct radioactivities.

The half-lives of ¹²⁹I and ²⁴⁴Pu are 17 × 10⁶ (4) and 82 × 10⁶ (5) years, respectively, and their decay products identified in meteorites are radiogenic ¹²⁹Xe (¹²⁹Xe) from ¹²⁹I and fissiogenic ¹³¹⁻¹³⁶Xe (¹³¹⁻¹³⁶Xe) from ²⁴⁴Pu. Because it is not possible to calculate the age of a body solely from the concentration of a stable decay product, total meteorite ages cannot be obtained by either ¹²⁹I-¹²⁹TXe or ²⁴⁴Pu-¹³¹⁻¹³⁶Xe dating. However, early ¹²⁹I : ¹²⁷I and ²⁴⁴Pu : ²³⁸U ratios in meteorites decreased rapidly because of the shorter half-lives of the nuclides in the numerator, and since the nuclides in each ratio display similar geochemical behavior, measurements of the ¹²⁹TXe: ¹²⁷I and ²⁴⁴Pu : ²³⁸U ratios present in meteorites formed (6, 7). For example, it has been found that chondrite meteorites were formed when the ¹²⁹I : ¹²⁷I ratio was about 1.1 × 10⁻⁴ (8) and the ²⁴⁴Pu : ²³⁸U ratio was found to vary from 10⁻⁸ to 10⁻⁵, and the initial ²⁴⁴Pu : ²³⁸U ratio varied from 0.002 to 0.006 (9), which suggests that the achondrites formed later than the chondrites. Meteorite age differences of a few million years have been detected by measurements of ¹²⁹TXe : ¹²⁷I and ¹³¹⁻¹³⁶fXe : ²³⁸U ratio varied from 0.002 to 0.006 (9), which suggests that the achondrites formed later than the chondrites. Meteorite age differences of a few million years have been detected by measurements of ¹²⁹TXe : ¹²⁷I and ¹³¹⁻¹³⁶fXe : ²³⁸U ratio (9, 10).

The chronology of early events for the earth is not so well established. The earth's interior is inaccessible, and the rocks and minerals of the earth's crust show a wide range of ages with an upper limit of about 3.5×10^9 years. Further, the continuous formation of new crustal material along oceanic ridges suggests that even today the earth's crust is not completely isolated from its mantle. Thus, the fundamental assumption of all radioactive dating, that the system has remained isolated or closed over the time period being measured, may not be valid for the accessible portions of the earth. Age estimates of the

earth based on ${}^{207}\text{Pb}$: ${}^{206}\text{Pb}$ ratios are undoubtedly among the more reliable since chemical fractionation would alter neither the ratio of the two parents, ${}^{235}\text{U}$: ${}^{238}\text{U}$, nor the ratio of the decay products, ${}^{207}\text{Pb}$: ${}^{206}\text{Pb}$. The ${}^{207}\text{Pb}$: ${}^{206}\text{Pb}$ ratios in oceanic sediments, lead ores, and old shield-area rocks suggest that the earth is approximately the same age as meteorites (2, 11). Even so, no clear evidence has previously been presented for the occurrence on earth of the decay products of the extinct radioactivities, ${}^{129}\text{I}$ or ${}^{244}\text{Pu}$.

The search for decay products of extinct elements in the earth has included several studies of xenon from gas wells (12-15). Boulos and Manuel (14) suggested that the xenon from deep gas wells up to a depth of 54.9 km was enriched in the heavy isotopes because of isotopic fractionation but noted that ¹²⁹Xe seemed to be enriched because of the decay of 129 I within the earth. Butler *et al.* (15) reported a small component of excess 129 Xe and an enrichment of the heavy xenon isotopes in CO₂ well gas from Harding County, New Mexico. They noted that the ¹²⁹Xe anomaly, if real, was due to the decay of extinct ¹²⁹I. In view of the uncertainty in the identification of the decay product of ¹²⁹I in terrestrial gas samples, a reinvestigation of CO₂ well gas from Harding County was undertaken. The samples were taken from a well drilled in 1969 in a field known as the Bueyeros Pool. The well has a depth of 6432 m and a wellhead pressure of 480 pounds per square inch (3312 N/m²). Methods of gas analysis and data reduction were described by Bennett and Manuel (12). The results of mass spectrometric analyses on duplicate samples are shown in Table 1 together with the isotopic composition of atmospheric xenon (16). The xenon concentration per unit volume of gas is about 1/10,000 of that in air.

Within the limits of statistical error, the isotope ratios from these two analyses are identical and correspond to atmospheric xenon at all mass numbers except 129, 132, 134, and 136. The abundance pattern of the xenon isotopes relative to that in the atmosphere (16) does not fit the pattern expected from isotopic fractionation (17). The ¹²⁹Xe isotope is enriched by about 10 percent over its atmospheric abundance, and the ¹³⁶Xe isotope is enriched by about 12 percent. The amount of excess ¹²⁹Xe we attribute to the decay of now extinct ¹²⁹I and show this as ¹²⁹Xe near the bottom of Table 1. In a similar manner the amount of excess ¹³⁶Xe we attribute to fission decay and designate this as ^{136f}Xe. We are certain that both of these anomalies are real and that terrestrial xenon therefore contains a record of extinct elements that were present in the very early history of the earth.

The presence of ^{129r}Xe in the CO₂ well gas is evidence that part of the xenon was derived from material that has not mixed with atmospheric xenon since a very early period when the earth's original inventory of ¹²⁹I had not completely decayed. The CO₂ may have resulted from volcanic emanations or from decomposition of carbonate rock through contact with intrusive magmas (15). Thus, the well may contain a mixture of gases from magmas and carbonates. The great majority of the New Mexico pools rich in CO₂ gas lie close to regions of recent igneous activity (18), and it seems likely that the very old xenon component, indicated by radiogenic ¹²⁹Xe, came from magmas, while the bulk of the CO₂, and probably other constituents, were derived from crustal carbonates.

Xenon isotope	Relative abundance of isotopes		
	Sample 1 (126 cm ³)	Sample 2 (78 cm ³)	Air (16)
¹²⁴ Xe	2.35 ± 0.10	2.41 ± 0.06	2.35
¹²⁶ Xe	2.25 ± 0.10	2.30 ± 0.06	2.21
¹²⁸ Xe	46.9 ± 0.5	47.5 ± 0.5	47.0
¹²⁹ Xe	711 ± 6	707 ± 4	648
¹³⁰ Xe	$\equiv 100$	≡ 100 ⁻	≡ 100
¹³¹ Xe	521 ± 5	524 ± 3	519
¹³² Xe	677 ± 6	676 ± 5	659
¹³⁴ Xe	278 ± 2	279 ± 3	256
¹³⁰ Xe	241 ± 2	245 ± 3	217
	Gas concentrations (10 ⁻¹	⁵ cm ³ per 1 cm ³ of gas)	
¹³⁰ Xe (total)	300 ± 60	400 ± 80	3,550,000
129r Xe	189	236	
¹³⁶ fXe	72	112	

Table 1. Xenon in samples of CO₂ well gas and in air. Volumes are given at standard temperature and pressure.

Enrichments due to radiogenic and fissiogenic xenon were calculated by subtracting atmospheric xenon (16) from the average xenon observed in the two samples of CO_2 well gas,

$$\delta_{i} = ({}^{i}Xe : {}^{130}Xe)_{CO2 well} - ({}^{i}Xe : {}^{130}Xe)_{air}$$
(1)

where i is the mass number and 130 Xe is the reference isotope because it is shielded from any contribution from fission. The results are shown in Fig. 1 with the yields of xenon isotopes from the spontaneous fission of 238 U (19) and 244 Pu (20).

The large excess of ¹²⁹Xe relative to excess ¹³¹⁻¹³⁶Xe is unlike any known fission yields (21). Butler *et al.* (15) have considered alternative possible origins for excess ¹²⁹Xe in CO₂ gas and concluded that the decay of now extinct ¹²⁹I is the most likely source. Srinivasan *et al.* (22) have shown that natural iodine, as incorporated in the mineral iodyrite (AgI), contained a small amount of ¹²⁹I from nuclear reactions in nature. They showed that nuclear reactions in the earth yielded a maximum equilibrium ratio for the two iodine isotopes of ¹²⁹I : ¹²⁷I $\leq 3.3 \times 10^{-15}$. This ratio and the average iodine content of limestone, 14 parts per million (23), can be used to show that only about 4×10^{-20} cm³ of excess ¹²⁹Xe observed in our analyses. Since neither the present levels of ¹²⁹I (22) nor any known nuclear reactions in nature (15) are capable of producing the excess ¹²⁹Xe shown in Fig. 1, we conclude that this is the decay product of part of the earth's original inventory of ¹²⁹I.

From our measurements, it is not possible to unambiguously determine the origin of the fission products, ^{131-136f}Xe. As can be seen from Fig. 1, the average yields at ¹³¹Xe, ¹³²Xe,

and ¹³⁴Xe relative to the yield of ¹³⁶Xe are slightly higher than the yields reported for the spontaneous fission of ²³⁸U (19). Butler *et al.* (15) also reported high yields of ¹³¹Xe and ¹³²Xe in CO₂ gas but ruled out any appreciable contribution from neutron-induced fission of ²³⁵U by the high ¹³⁴Xe : ¹³⁶Xe ratio in the latter. A mixture of fission products from ²⁴⁴Pu and ²³⁸U, about 31 percent of the excess ¹³⁶Xe from ²⁴⁴Pu and about 69 percent of the excess ¹³⁶Xe from ²³⁸U, fits the average yields from our two analyses. However, within the statistical error of our measurements the total enrichment of heavy xenon isotopes, ^{131-136f}Xe, could be from either ²³⁸U (19) or ²⁴⁴Pu (20). Even in the absence of definite information on the origin of ^{131-136f}Xe, the data from Table 1 place narrow limits on the age of the earth relative to that of meteorites.



Fig. 1. The excess xenon isotopes found in CO_2 gas wells relative to atmospheric xenon (16). The excess for each isotope of mass number i is expressed as δ_i , which is

defined by Eq. 1 (see text). The enrichment at each mass number is compared with that expected from the fission of 244 Pu (20) and 238 U (19).

It is impossible to make a quantitative calculation of the formation time of the earth from the concentration of ^{129r}Xe in the CO₂ wells without information on the iodine content and the quantity of magma that released this decay product. However, even if all the observed ^{136f}Xe in the CO₂ well is from ²⁴⁴Pu and if both ^{129r}Xe and ^{136f}Xe were derived exclusively from magma with the highest iodine : uranium ratio observed in deep-seated rocks (24), it can be shown by the method of Sabu and Kuroda (9) that the earth formed and began to accumulate radiogenic ^{129r}Xe no later than the achondrites. This means that any period of complete degassing, as might have accompanied the differentiation of the earth into a core, mantle, crust, and atmosphere, must have been limited to approximately the first 10⁸ years of the earth's history. It should be emphasized that this age estimate is based on an assumption that all of the ^{136f}Xe in the CO₂ wells is from ²⁴⁴Pu. If part of the ^{136f}Xe is from uranium or if part of the decay products were derived from material with a lower iodine : uranium ratio than assumed above, then an even earlier time of retention of these decay products in the earth is indicated.



Fig. 2. The ratio of xenon in air (16) to xenon released from lunar soil at 800°C (25). The line shows the effects of isotopic fractionation. Deviations from isotopic

fractionation at ¹²⁹Xe and ¹³⁶Xe are due to radiogenic ¹²⁹Xe in air and fissiogenic ¹³⁶Xe in the moon, respectively.

The occurrence of ^{129r}Xe in well gas suggests the possibility of radiogenic ¹²⁹Xe in the terrestrial atmosphere. A comparison (Fig. 2) of atmospheric xenon (16) with xenon implanted in lunar soil (25) from the solar wind shows that these two types of xenon are related by isotopic fractionation (17) except for a clear excess of ^{129r}Xe in air and an apparent excess of ^{136f}Xe in the moon. The fractional abundance of ¹²⁴Xe and ¹²⁶Xe in natural xenon is only about 10⁻³, and deviations from the fractionation line for these two isotopes are probably the result of nuclear reactions (26).

In conclusion, the presence of radiogenic ¹²⁹Xe in CO₂ wells shows that the earth started to accumulate this decay product no later than did some achondrites. Any quantitative degassing of the earth must have been limited to a relatively short time period. It seems more likely to us that the degassing process was never quantitative but has decreased over geologic time as suggested by Turekian (27). The presence of ¹²⁹Xe in the atmosphere with no clear evidence of ^{136f}Xe suggests that the initial retention of these decay products in terrestrial material may predate the formation of achondrites. The decay products of extinct elements in the earth are excellent tracers for studies of the events in the early history of the earth (28).

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