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Improved Limit on the Electron Capture Decay Branch of ¹⁷⁶Lu

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Abstract

We have performed searches for the electron-capture decay branches of ¹⁷⁶Lu to the ground state and first excited state of ¹⁷⁶Yb. No evidence of either decay mode was observed. From these measurements we have established upper limits on both of these possible branches that are each >20 times more stringent than the single previously published limit.

Keywords: Natural radioactivity, geochronology, electron-capture, Ge detector

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1. Introduction

The long-lived naturally occurring nuclide 176 Lu (J^{π} = 7⁻) β ⁻ decays to levels in ¹⁷⁶Hf. This decay is the basis for a dating method increasingly applied to rocks and meteorites. The utility of this dating method has been hampered by uncertainty in the half-life, as recently reviewed by Begemann et al. (2001). To some extent the half-life estimates vary systematically according to the approach used for its determination, with the lowest values, $(3.50 \text{ to } 3.57) \times 10^{10}$ years, implied from mass spectrometric measurements of Lu- correlated ¹⁷⁶Hf excesses in meteorites whose ages were independently known (Patchett and Tatsumoto, 1980; Blichert-Toft et al., 2002; Bizzarro et al., 2003). A half-life value of $(3.73 + 0.02) \times 10^{10}$ years obtained from the direct counting experiment of Nir-El and Lavi (1998) agrees with the $(3.72 \pm 0.03) \times 10^{10}$ years) value determined by Scherer et al. (2001) through age comparisons for terrestrial rocks. However, Browne and Junde (1998) evaluated the results of a large number of direct counting measurements and recommended a higher value of $(4.00 + 0.22) \times 10^{10}$ years. Moreover, the recent and precise γ - γ coincidence measurement of the ¹⁷⁶Lu half-life of $(4.08 + 0.03) \times 10^{10}$ years by Grinyer et al. (2003), which agrees very well with the earlier direct γ -ray counting results of $(4.08 + 0.24) \times 10^{10}$ years by Norman (1980) and (4.05 + $(0.09) \times 10^{10}$ years by Gerke et al. (1990), reaffirms a systematic difference of order 15% in the half-life deduced from these two approaches, which must be resolved before the ¹⁷⁶Lu-¹⁷⁶Hf method can be established as a reliable geo- and cosmo-chronometer.

It has been suggested that the discrepancies involving age comparisons could be reconciled if ¹⁷⁶Lu also underwent significant electron capture decay. ¹⁷⁶Lu is unstable with respect to electron-capture decay to ¹⁷⁶Yb. The Q_{EC} for decay to the ¹⁷⁶Yb ground state is 106.2 keV (Firestone, 1996). Thus, EC decays to both the $J^{\pi} = 0^+$ ground state and $J^{\pi} = 2^+$ 82-keV first excited state of ¹⁷⁶Yb are possible. These EC decay branches would be 7th and 5th forbidden transitions, respectively, and thus are expected to be negligibly small. In fact, there are no observed cases of such highly-forbidden transitions (Singh et al., 1998). The decay scheme of ¹⁷⁶Lu is illustrated in Figure 1. The published

limit on the EC decay branch of ¹⁷⁶Lu of < 10% was reported by Arnold (1954) in his study of the decay of ¹⁷⁶Lu. This upper limit was deduced from a search for Yb K x-rays that would be produced by the EC decay of ¹⁷⁶Lu. Because of the recent inconsistencies encountered in using the ¹⁷⁶Lu/¹⁷⁶Hf chronometer, it was felt that a new search for the EC decay of ¹⁷⁶Lu was warranted.

2. Experimental

Two plastic bottles, each containing 5 grams of 99.9% pure LuCl₃ ${}^{\circ}$ 6H₂O purchased from Alfa-Aesar were placed against the plastic endcap of a 110 cm³ high-purity germanium detector. X- and γ -ray data from approximately 20 – 800 keV were acquired in 4096 channels for a period of 65 hours using an ORTEC PC-based acquisition system. The energy resolution of this system was measured to be 1.4 keV full width at half maximum at 88 keV. Portions of the spectrum obtained from the Lu sample are shown in figures 2 and 3. The Hf K x rays and the 88-, 202-, 307-, and 401-keV γ rays produced by the β^{-} decay of 176 Lu were clearly observed. However, no evidence of Yb K x rays or 82-keV γ rays was seen. In order to have reference spectra of Yb and Hf K x rays, we also fluoresced samples of Hf metal powder and Yb₂O₃ with 122-kev γ rays from a 57 Co source.

3. Results

As can be seen from the decay scheme shown in Figure 1, all of the beta decays of ¹⁷⁶Lu eventually produce γ -ray transitions through the 88-keV level in ¹⁷⁶Hf. Thus, in order to establish a limit on the EC decay branch to the 82-keV level in ¹⁷⁶Yb, we determined both the net area of the 88-keV peak and the gross area of an equal width energy interval centered on 82 keV in the spectrum obtained from the Lu₂O₃ sample. The net area of the 88-keV peak was determined to be N₈₈ = 79111 ± 534 counts, whereas the gross area of the 82-keV region was found to be 46623 ± 216 counts. Since no peak was observed at 82 keV, we multiplied the gross area of this interval by 2 and then took the square root to obtain a 1 σ upper limit, (N₈₂ < 305) on the net area of 82-keV peak. Both

the 88-keV level in ¹⁷⁶Hf and the 82-keV level in ¹⁷⁶Yb decay mainly by internal conversion electrons rather than by photon emission. The total internal conversion coefficient, α_T for the 88-keV transition is 5.914, and that for the 82-keV transition is 7.125 (Hager and Seltzer, 1968). Because of the small energy difference between these two transitions, we assumed the same photopeak detection efficiencies for them. Thus, the upper limit we can establish on the ¹⁷⁶Lu EC branch to the 82-keV level in ¹⁷⁶Yb is

$$B_{82} < \frac{(1 + \alpha_T(82)) N_{82}}{(1 + \alpha_T(88)) N_{88}} = \frac{(8.125)(305)}{(6.914)(79111)} = 0.0045 \text{ (or } 0.45\%).$$

The beta decay of ¹⁷⁶Lu produces Hf x rays through the atomic vacancies created by internal conversion electrons emitted from the decays of the 998-, 597-, 290-, and 88keV levels in ¹⁷⁶Hf. Hf K_{α} x rays are emitted in 25.4% of all ¹⁷⁶Lu beta decays (Browne and Firestone, 1986). Yb K_{α} x rays would be produced in 38.4% of all ground state to ground state EC decays of 176 Lu (Browne and Firestone, 1986; Schonfeld, 1998). Yb K_a x rays are 3.3 keV lower in energy than those of Hf (Browne and Firestone, 1986). Thus in order to establish an upper limit on the EC decay of ¹⁷⁶Lu to the ground state of ¹⁷⁶Yb, we determined the net area of the Hf K_{α} x-ray doublet, N_{Hf} = 331382 ± 576, and established an upper limit on the net area (N_{Yb}) of a possible Yb K_{α} x-ray peak in the spectrum obtained from the Lu₂O₃ sample. To establish this upper limit, we added a scaled version of the spectrum obtained from the fluorescence of the Yb₂O₃ sample to the spectrum obtained from counting the Lu₂O₃ sample. We then compared the resulting sum to the original Lu spectrum to see if we could observe "shoulders" on the low-energy sides of the Hf x-ray peaks that would be caused by the presence of the Yb x rays. In the Yb_2O_3 fluorescence spectrum $N_{Yb} = 18065 + 247$. Our upper limit on the scale factor at which we could still detect the Yb x rays in the summed spectrum was 0.10. Again we assumed that because of the small energy difference between these two transitions, the photopeak detection efficiencies for the Yb and Hf K_{α} x rays were equal. Thus, our 1 σ upper limit on the EC decay of ¹⁷⁶Lu to the ¹⁷⁶Yb ground state is:

$$B_{gs} < [0.10 \text{ x } 18065 \text{ x } 0.254] / [331382 \text{ x } 0.384] = 0.0036 \text{ (or } 0.36\%).$$

4. Conclusion

As expected, no evidence for the EC decay of ¹⁷⁶Lu to either the ground state or first excited state of ¹⁷⁶Yb was observed. The upper limits we have been able to establish on these possible EC transitions are each >20 more stringent than the single previously published limit of <10% for the EC decay of ¹⁷⁶Lu. Our results are also about a factor of two more stringent than that recently obtained from a mass spectrometric search for excess ¹⁷⁶Yb in $(1.0 - 2.7)x10^9$ year-old Lu-bearing samples (Amelin and Davis, 2003). Taken together, these results indicate that an explanation for discrepancies between ¹⁷⁶Lu half-life estimates based on age comparisons and those based on direct counting methods must be sought elsewhere.

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Figure 1. Decay scheme of 176 Lu. The possible electron-capture decay branches to the ground and first-excited states of 176 Yb are illustrated. All energies are given in keV.



Figure 2. Gamma–ray spectrum observed from the 10-g sample of Lu_2O_3 . Peaks attributed to the β^- decay of ^{176}Lu are indicated.



Figure 3. Expanded region of the spectrum observed from the 10-g sample of Lu₂O3. The expected position of the 82-keV γ ray from the possible EC decay of ¹⁷⁶Lu is indicated.