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Recovery of No-Carrier-Added ⁴¹Ca, ⁴⁴Ti, and ²⁶Al from High-Energy Proton-Irradiated Vanadium Targets

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Abstract.

Many useful and needed radionuclides for medicinal, astrophysical, and environmental research are produced naturally in inefficient quantities or not-at-all. In the method described here, rare cosmogenic isotopes were produced via spallation reactions in metallic vanadium and separated without adding any carriers. In the SINQ facility at the Paul Scherrer Institut, the vanadium targets were irradiated for two years with high-energy protons (\leq 590 MeV). After a cooling period of eight years, only relatively long-lived radionuclides such as 32 Si, 44 Ti, 41 Ca, and 26 Al remain present. After target dissolution, 32 Si was first separated for a prospective half-life redetermination. The remaining 32 Si-free solution was used for extracting 44 Ti, 41 Ca, and 26 Al, three key isotopes which are scientifically interesting for nuclear astrophysics research as well as medical applications. Each separation scheme employed ion-exchange and extraction chromatography; developed and optimized using inactive model solutions analyzed with Inductively Coupled Plasma – Optical Emission Spectrometry (ICP-OES). The irradiated samples were tracked with γ -ray spectroscopy for γ -ray emitting impurities. As a result, radiochemically pure sample solutions of 44 Ti, 41 Ca, and 26 Al were obtained as "ready for use" in different application fields.

Introduction

In a previous paper [1], we reported on the separation of ³²Si from vanadium disks, which had been irradiated with high-energetic protons in the Swiss Spallation Neutron Source (SINQ) target at PSI for two years. The work was aimed at providing enough sample material for a reliable redetermination of the ³²Si half-life. The remaining solution contained, among others, three exotic carrier-free nuclides: ⁴⁴Ti, ⁴¹Ca, and ²⁶Al, which are of particular interest in several fields of nuclear science:

i) Typically, ⁴⁴Ti is formed via nucleosynthesis in supernovae [2]. It decays via electron capture (EC) and has an associated dual γ -ray emission used for young supernovae identification [2, 3]. Its half-life, $t_{1/2} = 59.1(3)$ yr [4], prevents the nuclide from being directly seen in meteorites since its travel time to Earth is much longer than the life of ⁴⁴Ti. Instead, the large excesses of its stable

ii) ⁴¹Ca is mainly formed within supernovae [6] and is an ideal candidate for nuclear dating because its

granddaughter, ⁴⁴Ca, indirectly quantifies ⁴⁴Ti [5, 6]. Medically, ⁴⁴Ti is sought after as the parent for the 44Ti/44Sc generator [7]. 44Sc decays via EC and β + emission ($t_{1/2} = 3.97(4)$ hr [4]) and is an ideal candidate for Positron Emission Tomography (PET). Most PET isotopes require on-site production via cyclotron, subsequent separation, and preparation for medical applications. A nuclide generator widens the number of facilities able to use PET without a cyclotron [7]. Fortunately, ⁴⁴Ti only decays to ⁴⁴Sc, rather than the longer-lived ^{44m}Sc ($t_{1/2} = 58.61(10)$ hr [4]). The benefits of 44Sc as a PET nuclide are: a short halflife, a biologically friendly stable daughter ⁴⁴Ca, and, necessary for a generator, a mother with a half-life longer than the daughter nuclide.

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half-life $(9.94(15) \times 10^4 \text{ yr } [8])$ bridges the age gap between two commonly used dating radionuclides, ^{14}C and ^{36}Cl ($t_{1/2} = 5.700(30) \times 10^3 \text{ yr}$ and $3.013(15) \times 10^5 \text{ yr}$ respectively) [9]. In addition, ^{41}Ca can be used as a long-term medical tracer for sorption within bones [10]. For accurate analysis, large quantities of ^{41}Ca for standards are necessary to date meteorites [6], bones [11], and limestone [12].

iii) 26 Al ($t_{1/2} = 7.17(24) \times 10^5$ yr) decays via EC and β + with an associated γ -ray emission [13]. Like the preceding nuclides, it plays a role in nuclear astrophysical research where the information gained from 26 Al γ -ray analysis ranges from nucleosynthesis [14] to the supernova explosion of massive stars [6]. It is also cosmogenically formed through neutron-induced spallation of argon in the atmosphere [15] or through muon interactions in quartz [16]. Its constant surface production allows dating with 26 Al/ 10 Be isotope ratios within ice [17] and caves [18].

In order to study the nuclear astrophysics processes described above as well as for establishing these nuclides as environmental dating tools, sufficient sample amounts in the requested quality fitting to the specific envisaged experiment - for instance cross section measurements or determination of nuclear decay properties like half-lives, emission probabilities or branching ratios – are urgently needed. Complementary production routes exist for all three isotopes. 41Ca can easily be produced via neutron activation of ⁴⁰Ca in a nuclear reactor, but the reaction product contains large amounts of stable Ca, preventing from its use in many applications. Non-carrier added ⁴⁴Ti can be obtained by proton irradiation of natural scandium as explored by [7]. The challenge with this approach is that the daughter nuclide of the generator is the same element as the target material, so that extremely high separation of the titanium from the scandium is required. 41Ca and 26Al can also be produced in (p,n) reactions, but the natural abundances of the needed target materials, 41K and ²⁶Mg, are only 6.7% and 11%, respectively. To some extent, this reduces the advantage of the higher cross sections. Moreover, highly enriched material has to be used to suppress the production of unwanted byproducts. The advantage of isotope production using the SINQ target at PSI is the vast range of interesting isotopes and the beam time for free. This outweighs the effort to develop and perform the more complicated chemical separations.

The excitation functions for all three isotopes from proton-irradiated V have been determined earlier [19,20]. For ⁴⁴Ti, no chemical separation was needed because of the good measurability of the radiation [19], however ⁴¹Ca and ²⁶Al both required chemical separation prior to detection [20]. A key difference between the separation procedures presented here and by the other authors [20] is the irradiated vanadium targets here were approximately three times larger and irradiated longer. As a result, separations occurred from a

larger vanadium matrix with more undesired radionuclides within the irradiated samples. Presented here is the separation of ⁴¹Ca, ⁴⁴Ti, and ²⁶Al from a vanadium matrix solution via chromatographic separation methods after the previous removal of ³²Si, reported earlier [1].

Experimental Studies

Materials and Instruments

All separations and purifications utilized ionexchange and extraction column chromatography. Titanium and calcium separations exclusively used extraction resins, whereas aluminum separations required a series of ion-exchange and extraction resins. For the ion-exchange experiments, Dowex® resins were used for both cation- (Dowex® 50WX8 resin, 200-400 mesh, Sigma-Aldrich, Germany) and anion-exchanges (Dowex® 1X8 resin, 200-400 mesh, Sigma-Aldrich, Germany). The extraction resins (TrisKem SAS, Bruz, France) were normal DGA (50-100 µm), LN1 (20-50 μm), and LN3 (20-50 μm). Each experiment was optimized first with non-irradiated metallic vanadium disks identical to the irradiated targets ($\emptyset = 9 \text{ mm}$, 1 mm thick, $\approx 417 \pm 4$ mg, chemical purity: 99.8% V, Goodfellow Cambridge Limited, U.K.). Single element standards (1000 mg/L in 2% HNO₃, Sigma-Aldrich, Germany) were used to develop the separation methods. Preconditioning and eluents were diluted HNO₃ (69 wt.-%, Honeywell, U.S.A.) or HF (40 wt.-%, Merck, Germany) prepared with ultra-pure water (upt- H_2O , 18.2 $M\Omega \times cm$, Veolia S.A.).

Each setup utilized an in-house made pressurized column apparatus comprised of: the chosen resin, polymethyl methacrylate columns (inner- \emptyset = 10 mm, and 300 or 100 mm length), squeeze tubes (Ismaprene PharMed®, Cole-Parmer Instrument Company, LLC., U.S.A.), PE frits (\emptyset = 10 mm, 30 μ m pore-size, Kinesis Group from Cole-Parmer Instrument Company, LLC., U.S.A.), and a peristaltic pump (REGLO Digital MS-2/8, Cole-Parmer Instrument Company, LLC., U.S.A.).

For the analysis during method development, ICP-OES (Agilent 5110, Agilent Technologies, U.S.A) was employed using inactive solutions. The associated breakthrough curves can be viewed in the Supplementary Information (SI). The irradiated samples were tracked via γ-ray spectrometry using a high-purity germanium (HPGe) detector (Mirion Technologies, California, U.S.A.) as well as via liquid scintillation counting (LSC) using a Packard Tri-Carb® 2250CA.

Irradiation and First Separation

At the SINQ facility of PSI, 150 vanadium targets were irradiated for two years under a high-energy proton beam (≤ 590 MeV) [21]. The Gaussian shaped

beam irradiated the stacked targets perpendicular to the stack lengthwise, thus each target was subjected to a different power intensity resulting in a different composition of radionuclides per target [22]. After irradiation, the targets were placed in intermediate storage for eight years without previous measurement. The initial target dissolution and first chromatographic separation of ³²Si via cation-exchange chromatography are described elsewhere [1], thus not a subject in this study. After the ³²Si removal, ⁴¹Ca, ⁴⁴Ti, ²⁶Al, and the vanadium bulk remained in 50 mL 3 M HNO₃, the starting solution for the proceeding separations.

General Separation Procedure

Unless otherwise stated, all chromatographic separations were carried out according to the following outline. Before assembly, each resin was saturated in upt-H₂O for at least 24 hr, then added into the column, held between two PE frits. Solutions were introduced through a feeder connected to a peristaltic pump with squeeze tubing maintained at 1.50 mL/min. Prior to the experiments, the resins were pre-conditioned with the solvent of the loading solution; stated for each separation specifically. The experiments then followed these general steps: (i) the loading solution containing all matrix elements (Load), (ii) a wash of the same solvent (Wash), (iii) a chosen eluent solution to wash-off desired elements (Elute), and eventually (iv) one or more consecutive solutions preparing the resin for reuse (post-conditioning). Unless otherwise stated, the Load and Wash were collected together in preparation for the subsequent element separation. However, the Elute was further utilized for purification procedures.

Calcium Separation

For the calcium separation from vanadium, a 30 mm high resin bed of DGA resin was preconditioned with 10 mL 3 M HNO₃ solution. The initial vanadium solution after ³²Si separation [1] was used as the Load without further manipulation. The Load, 50 mL 3 M HNO₃, and Wash, 20 mL 3 M HNO₃, were collected for further separations. Calcium was eluted with 20 mL upt-H₂O, followed by 10 mL 3 M HNO₃ post-conditioning.

The calcium fractions then underwent two purification steps, where the second step additionally served to concentrate the calcium fraction (see Fig. 1). The collected Elute from the bulk vanadium separation was acidified with 0.66 mL conc. HNO₃ to reach a 20.66 mL 0.5 M HNO₃ Load. The LN3 resin (bed height: 30 mm) was preconditioned with 10 mL 0.5 M HNO₃, followed by the Load and 5 mL 0.5 M HNO₃ Wash. The Load and Wash were collected together for the next purification step. In preparation for resin reuse, 15 mL upt-H₂O then 10 mL 0.5 M HNO₃ were added as post-conditioning.

Finally, the solution was purified and concentrated using DGA resin (bed height: 30 mm). Here, the Load

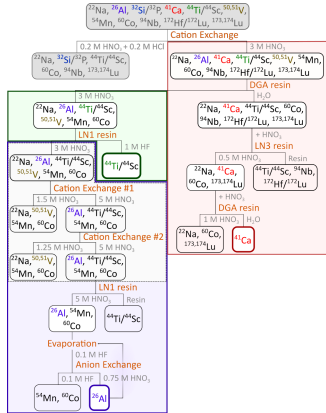


Fig. 1. Full separation schematic starting from the initial vanadium target solution and ³²Si separation (gray) [1]. Highlighted in red are the ⁴¹Ca separations: vanadium removal with DGA resin, purification #1 with LN3 resin, and purification #2 and concentration with DGA resin. In green is the ⁴⁴Ti separation: vanadium removal with LN1 resin. Finally, in purple are the ²⁶Al separations: two-step vanadium removal with cation exchange resin, purification #1 with LN1 resin, concentration via evaporation, and purification #2 with anion exchange resin. The evaporation and 2nd purification are repeated. Fig. 2 is a more detailed display of the two-step vanadium removal (dashed gray within the purple). The final fractions are bolded in their respective colors.

was the collected Load and Wash from the previous purification on LN3 resin, 25.66 mL 0.5 M HNO₃, which was then acidified with 0.876 mL conc. HNO₃ to reach 1 M HNO₃. The calcium solutions from ten vanadium targets were used as the Load (approx. 265 mL 1 M HNO₃). The resin was preconditioned with 15 mL 1 M HNO₃ before adding the \approx 265 mL Load, 20 mL 1 M HNO₃ Wash, 20 mL utp-H₂O Elute (final ⁴¹Ca solution), then 15 mL 1 M HNO₃ post-conditioning.

Titanium Separation

The 70 mL 3 M HNO₃ Load and Wash obtained from the calcium separation from vanadium was used directly as the Load solution here. First, the LN1 extraction resin (bed height: 20 mm) was preconditioned with 15 mL 3 M HNO₃. Sequentially, the 70 mL Load, 20 mL 3 M HNO₃ Wash, and 20 mL 1 M HF Elute were fed through the column (see Fig. 1). The Load and Wash were collected for the separation of alumi-

num. Following the elution, 10 mL upt- H_2O and 15 mL 3 M HNO₃ were used to prepare the resin for a subsequent vanadium solution.

Aluminum Separation

Here, the aluminum separation from the vanadium matrix was optimized utilizing a two-step cation exchange. For Step #1, the 90 mL 3 M HNO₃ Load and Wash from the titanium separation from vanadium was diluted to 1.5 M HNO₃ with 90 mL upt-H₂O as the first Load. A 210 mm resin bed height column was preconditioned with 20 mL 1.5 M HNO₃ followed by the 180 mL Load and 30 mL 1.5 M HNO₃ Wash. The Elute, 40 mL 5 M HNO₃, was collected in two separate 20 mL fractions; the Load and Wash were discarded (see Fig. 2, Step #1).

In preparation for Step #2, each collected Elute fraction from Step #1 was diluted to 1.25 M with 60 mL upt- H_2O . These solutions were then used as the Load for Step #2 in the same order as collection in Step #1 (see Fig. 2). For Step #2, a 130 mm resin bed height column was preconditioned with 20 mL 1.25 M HNO₃ followed by: both Loads, 2x 80 mL 1.25 M HNO₃; Wash, 30 mL 1.25 M HNO₃; then Elute, 35 mL 5 M HNO₃. The Elute was collected as the separated 26 Alfraction and used for further purification; the Load and Wash were discarded. For both cation exchange separations, upt- H_2O was the post-conditioning until pH \approx 4. This was followed by 20 mL 1.5 M or 1.25 M HNO₃ for Step #1 and #2, respectively.

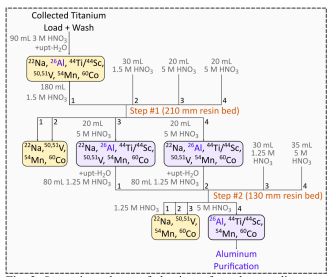


Fig. 2. Separation scheme of aluminum from the vanadium matrix using two cation-exchange resin bed heights. Step #1 shows how the Elute was collected in two fractions, then diluted and used as the Load for Step #2. Shown is the expanded dashed gray section of Fig. 1.

Following the separation from vanadium, the ²⁶Alfraction was subjected to two purifications and one concentration step (see Fig. 1). The collected Elute, 35 mL 5 M HNO₃, was used directly as the Load for a first purification using the LN1 extraction resin (bed height: 30 mm). The resin was preconditioned with

 $10~\text{mL}~5~\text{M}~\text{HNO}_3$, proceeded by the 35~mL~Load and $10~\text{mL}~5~\text{M}~\text{HNO}_3$ Wash, collected together for the next step. Post-conditioning consisted of $10~\text{mL}~\text{upt-H}_2\text{O}$, $20~\text{mL}~1~\text{M}~\text{HF},~10~\text{mL}~\text{upt-H}_2\text{O}$, and $10~\text{mL}~5~\text{M}~\text{HNO}_3$.

The second purification required a matrix change from HNO₃ to HF. This was achieved via evaporation which additionally served as a concentration step. On a PTFE evaporation dish, the collected Load and Wash from the first purification (45 mL 5 M HNO₃) was evaporated to dryness. The equivalent of three vanadium target solutions were concentrated to one solution, i.e., 135 mL 5 M HNO₃. The solid residue was then dissolved in 5 mL 1 M HF and upt-H₂O was added to reach a final concentration of 0.1 M HF that was used as Load for the second purification step.

A 30 mm bed height anion-exchange resin was preconditioned with 15 mL 0.1 M HF. This was followed by: 50 mL Load, 10 mL 0.1 M HF Wash, and then 20 mL 0.75 M HNO₃ Elute. For post-conditioning, 10 mL 1 M HF, 20 mL upt-H₂O, and 15 mL 0.1 M HF were used. For the final purification, the Elute was once more evaporated to dryness before repeating the anion-exchange separation.

Results & Discussion

The qualitative γ -ray spectra of 41 Ca, 44 Ti, and 26 Al after successful separation from the irradiated vanadium targets are shown in the following section. Unless otherwise stated, the obtained fractions were measured with the varying full volumes and normalized against respective acquisition time. Stated recovery yields are calculated from the nonradioactive method development with ICP-OES. For detailed elution profiles, see the SI.

Calcium

DGA resin is well-known to retain calcium in nitric acidic environments [20, 23] and elute out in water [24]. Vanadium was removed from calcium by a factor of 19.91(0.05) x 10⁴ while calcium was recovered at 99.8% \pm 2%. As previously mentioned, ⁴¹Ca is a non- γ ray emitting nuclide, thus, γ-ray spectrometry was used to trace contaminants. The targeted impurities were ²²Na, ⁴⁴Ti/⁴⁴Sc, ⁶⁰Co, ⁹⁴Nb, ¹⁷²Hf/¹⁷²Lu, and ^{173,174}Lu, shown in Fig. 3 (Spectrum 1). The only predicted contaminants after separation through DGA resin were ^{173,174}Lu (SI Fig. 2), ¹⁷²Hf, and ⁹⁴Nb [23]. Nonetheless, multiple other radionuclides were additionally seen, e.g., ²²Na and ⁶⁰Co. For titanium, some retention is possible on DGA resin at 3 M HNO₃ [23], but this was not seen in the model elution profiles (SI Fig. 2). The results of the model break-through curves can be attributed to the detection limit of the ICP-OES, which is much higher as opposed to γ-spectrometric measurements.

The first purification, using the LN3 extraction resin, removed Hf(IV), Lu(III), and Ti(IV) entirely but Nb(V) only partially. Between the different available LN extract resins (LN1, LN2, and LN3), Ca(II) retains the least for any given nitric acid concentration in LN3, whereas Hf(IV), Ti(IV), and Nb(V) retain in all concentrations of HNO3 for all LN resins. Lu(III) has moderate retention at 0.1 M HNO3, and Ca(II) has no retention at the same concentration [25]. Thus, it was chosen as the Load concentration. The successful removal of $^{172}{\rm Hf}$ (t_{1/2} = 1.87(3) yr) was tracked by the daughter's γ -ray emission ($^{172}{\rm Lu}$, t_{1/2} = 6.70(3) days) multiple weeks after separation [26]. The purification with LN3 resin successfully removed $^{44}{\rm Ti}/^{44}{\rm Sc}$, $^{173,174}{\rm Lu}$, and $^{172}{\rm Hf}/^{172}{\rm Lu}$, see Fig. 3 (Spectrum 2).

For the final purification and concentration step, the reported maximum retention for calcium on DGA resin, 1 M HNO₃, was the chosen condition for the Load [23]. All remaining contaminants, 22 Na, 60 Co, and 94 Nb, were successfully removed, and the final spectra matched a measured background spectrum (see Fig. 3, spectrum 3). For this purification, calcium was recovered at $100\% \pm 0.7\%$ with an average of 5 kBq/g. Finally, the successful separation and average activity concentration of 41 Ca was verified by LSC (see SI Fig. 4).

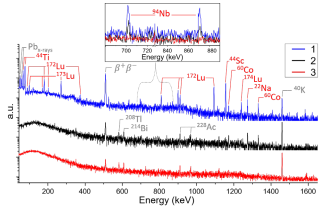


Fig. 3. The γ-ray spectra of 41 Ca solutions after: (1) separation from vanadium on DGA resin, 22 Na, 44 Ti/ 44 Sc, 60 Co, 94 Nb, 172 Lu/ 172 Hf, and 173 Lu remain; (2) separation from 44 Ti/ 44 Sc, 172 Lu/ 172 Hf, and 173 Lu on LN3 resin, where 22 Na, 60 Co, and 94 Nb remaining; (3) separation of 22 Na, 60 Co, and 94 Nb on DGA resin. The sub-spectra shows the energy range of 94 Nb, notably its absence in the final solution.

Titanium

The LN1 extraction resin was found to be the most reliable for separating titanium from the vanadium matrix in 3 M HNO₃ [25]. Titanium was recovered at 98.8% \pm 0.2% and vanadium was removed by a decontamination factor of 6.99(0.1) x 10⁴. Although traces of $^{44}\mathrm{Ti}$ were found in the $^{41}\mathrm{Ca}$ -fraction and in the Load for the aluminum separation, the majority was collected in the $^{44}\mathrm{Ti}$ Elute. Because of its high activity, a 5 mL aliquot of the 20 mL Elute was used for γ -ray analysis (see Fig. 4). In addition to vanadium, $^{22}\mathrm{Na}$ and $^{60}\mathrm{Co}$

were successfully removed with LN1 resin and no other γ -ray emitting nuclides were found. The average activity concentration of 44 Ti being 100-200 kBq/g.

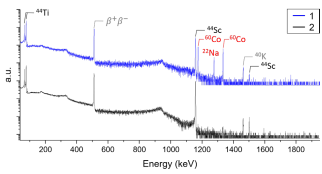


Fig. 4. The γ -ray spectra of 3 M HNO₃ Load (1) and 1 M HF Elute (2) of the titanium separation on LN1 extraction resin. Note the daughter's emission (44 Sc) at 1157 and 1499 keV [4].

Aluminum

Due to the large concentration ratio between the vanadium and aluminum in solution and their similar retention behavior on cation-exchange resins [27], a two-step approach was chosen to remove the vanadium matrix. The applied separation scheme, with a split collected Elute, proved successful for maximum aluminum separation from vanadium while minimizing the radioactive waste (see SI Fig. 6 and 7). Aluminum was recovered at $99.6\% \pm 0.3\%$ and vanadium removed by a factor of $5.88 \pm 0.01 \times 10^{-6}$ following the second step. After both cation-exchange steps, $^{44}\text{Ti}/^{44}\text{Sc}$, ^{54}Mn , and ^{60}Co remain as contaminants (see Fig. 5).

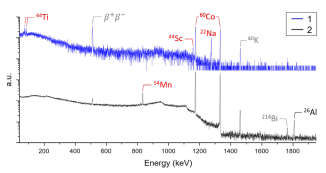


Fig. 5. The γ -ray spectra of 26 Al before (1) and after (2) the vanadium separation with both cation exchange resin columns.

Regarding the distribution coefficients (K_d) for Ti(IV), Mn(II), and Co(II), it is expected that small quantities are still present after cation-exchange since their K_d values are between V(IV) and Al(III) [27]. As seen previously, LN1 extraction resin removed ⁴⁴Ti, thus it was used as the first purification step for ²⁶Al, before addressing ⁵⁴Mn and ⁶⁰Co. However, since the LN1 extraction resin does not retain Mn(II), Co(II), nor Al(III) under the chosen conditions [25], a different approach was necessary to separate these nuclides.

The separation of aluminum was achieved by converting aluminum into an anionic complex ([AIF₄]⁻). At low HF concentrations, [AIF₄]⁻ is retained on anion-exchange resins, while Mn(II) and Co(II) are not [28, 29]. Contrary to the model experiments where aluminum was recovered at $100\% \pm 2\%$ and all contaminants removed (SI Fig. 9), traces of ⁵⁴Mn and ⁶⁰Co were still visible after the separation. As a result, approximately 80% of both contaminants were removed, while ²⁶Al was quantitatively recovered. Consequently, a repetition of the evaporation and subsequent separation on the anion-exchange resin removed the remaining ⁵⁴Mn and ⁶⁰Co (see Fig. 6). The average activity concentration was 5-10 Bq/g.

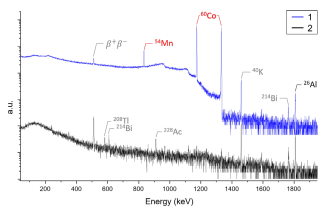


Fig. 6. The γ-ray spectra of 26 Al after concentration via evaporation (1) and the final 26 Al fraction after the second anion-exchange separation (2).

Conclusion

Rare and exotic isotopes such as 41Ca, 44Ti, and 26Al, are urgently needed for applications in nuclear medicine or nuclear astrophysics. Usually, their production is very challenging, but these nuclides have been identified in exceptionally high quantities as by-products due to proton-induced spallation of metallic vanadium targets. Here, the development of a selective and robust chromatographic separation method has proven to successfully isolate ⁴¹Ca, ⁴⁴Ti, and ²⁶Al from the vanadium matrix. In the process of developing the optimal separation conditions, inactive model solutions, precisely matching the expected matrix composition, were analyzed using ICP-OES. For the radioactive solutions, the separation procedure was carried out without the addition of stable carriers. Consequently, the ⁴¹Ca, ⁴⁴Ti, and ²⁶Al were obtained as no-carrier-added. The radiochemical purity of the final fractions was confirmed by γ-ray spectrometry and LSC. Thus, vanadium targets should be considered as candidates for long-term irradiations to provide a variety of no-carrier-added rare isotopes. In particular, a sufficiently high amount of "ready-for-use" 44Ti can be provided for the development of a future radionuclide generator system as well as other exotic radioisotopes, i.e., ⁴¹Ca and ²⁶Al, for

instrument calibration or dedicated studies on their own.

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