Radiochemical Studies of the Transactinide Element, Rutherfordium (Rf) at JAERI

Y. Nagame*
Advanced Science Research Center, Japan Atomic Energy Research Institute, Tokai, Ibaraki 319-1195, Japan

Received: June 30, 2005; In Final Form: August 27, 2005

Radiochemical studies of element 104, rutherfordium (Rf), at JAERI (Japan Atomic Energy Research Institute) are reviewed. The transactinide nuclide $^{261}$Rf has been successfully produced in the reaction $^{248}$Cm$(^{18}$O, 5n) at the JAERI tandem accelerator. The excitation function of the reaction has been measured with a rotating wheel catcher apparatus and the maximum production cross section has been determined to be about 13 nb. On-line ion-exchange experiments on Rf together with the group-4 elements Zr and Hf in acidic solutions have been conducted with a rapid ion-exchange separation apparatus. From the systematic study of the anion-exchange behavior of Rf, it has been found that the properties of Rf in HCl and HNO$_3$ solutions are quite similar to those of Zr and Hf, respectively, confirming that Rf is a member of the group-4 elements. However, we have observed an unexpected chemical behavior of Rf in HF solutions; the fluoride complex formation of Rf is significantly different from those of Zr and Hf. The current progress in the Rf aqueous chemistry is briefly summarized and prospects of extending chemical studies of transactinide elements at JAERI are considered.

1. Introduction

Presently, we know more than 20 man-made transuranium elements. According to the actinide concept,$^1$ the 5f electron series ends with element 103, lawrencium (Lr), and a new 6d electron transition series is predicted to begin with element 104, rutherfordium (Rf). The elements with atomic numbers $Z \geq 104$ are called transactinide elements or recently called superheavy elements.$^2$ The currently known transactinide elements, elements 104 through 112, are placed in the periodic table under their lighter homologues in the 5d electron series, Hf to Hg. Elements from 113 to 118 except for 117 have been recently synthesized$^3$ and would be in the successive 7p electron series (see Figure 1), although the discoveries of elements with $Z \geq 112$ are waiting to be confirmed.$^5$

Studies on the chemical properties of the elements at the uppermost end of the periodic table are extremely interesting and challenging subjects in the fields of nuclear and radiochemistry. It is also of special interest to assess the magnitude of the influence of relativistic effects on chemical properties. From the calculations of electron configurations of heavier elements, it is predicted that sudden changes in the structure of electron shells may appear due to relativistic effects which originate from the increasingly strong Coulomb field of a highly charged atomic nucleus.$^7$ Therefore, it is expected that heavier elements show a drastic rearrangement of electrons in their atomic ground states, and as electron configurations are responsible for chemical behavior of elements, such relativistic effects can lead to surprising chemical properties. Increasing deviations from the periodicity of chemical properties based on extrapolation from lighter homologues in the periodic table are consequently predicted.$^5$ It would be no longer possible to deduce detailed chemical properties of the transactinide elements simply from the position in the periodic table.

The transactinide elements must be produced at accelerators using reactions of heavy-ion beams with heavy target materials and must be identified by measurement of their decay or that of their known daughter nuclei with unambiguous detection techniques. Chemical experiments with the transactinide elements are generally divided into the following 4 basic steps: i) synthesis of transactinide nuclides, ii) rapid transport of the synthesized nuclides to chemical separation apparatuses, iii) fast chemical isolation of a desired nuclide and preparation of a sample suitable for nuclear spectroscopy, and iv) detection of nuclides through their characteristic decay properties. Because of the short half-lives and the low production rates of the transactinide nuclides, each atom produced decays before a new atom is synthesized. This means that any chemistry to be performed must be done on an "atom-at-a-time" basis. Thus, rapid and very efficient radiochemical procedures must be devised. Recent comprehensive reviews on the chemistry of the transactinide elements are seen in References 2, 11, and 12.

In order to carry out chemical experiments of the transactinide elements with single atoms, we developed some experimental apparatuses: a beam-line safety system for the usage of a gas-jet coupled with a radioactive target and recoil chamber, a rotating wheel catcher apparatus for the measurement of $\alpha$ and spontaneous fission (SF) decay of transactinide nuclei, and an automated rapid ion-exchange separation apparatus based on high performance liquid chromatography connected with an on-line $\alpha$-particle detection system.$^7$

The transactinide nuclide, $^{78+}$Rf, which is commonly used for recent chemical studies of element 104, was successfully produced in the reaction $^{246}$Cm$(^{18}$O, 5n) at the JAERI tandem accelerator for the first time in Japan. The excitation function of the reaction was measured with the rotating wheel catcher apparatus MANON (Measurement system for $\alpha$ particle and spontaneous fission events ON-line), and the maximum

---

*Corresponding author. E-mail: nagame.yuichiro@jaea.go.jp.
FAX: +81-29-282-5927.

---

© 2005 The Japan Society of Nuclear and Radiochemical Sciences
Published on Web 11/07/2005
production cross section was determined to be 13 nb at around 94-MeV ¹⁸O energy.⁴ On-line anion-exchange experiments of Rf together with the group-4 elements Zr and Hf in acidic solutions, HCl, HNO₃, and HF, were conducted with the rapid ion-exchange separation apparatus, AIDA (Automated Ion-exchange separation apparatus coupled with the Detection system for Alpha-spectroscopy).¹⁵,¹⁶ AIDA enables us to perform cyclic discontinuous column chromatographic separations of short-lived nuclides in aqueous solutions and automated detection of α-particles within a typical cycle of 1–2 min. The adsorption behavior of Rf as a function of the acid concentration has been systematically studied with AIDA.¹⁵,¹⁶

In this paper, the progress in the radiochemical studies of Rf and future research plans for extending chemical studies of the transactinide elements at JAERI are summarized.

2. Production of the Transactinide Nuclide ²⁶¹Rf

A schematic representation of the experimental set-up including a target and recoil chamber arrangement and the rotating wheel catcher apparatus MANON is shown in Figure 2. A ²⁴⁸Cm target of 590-μg/cm² thickness prepared by electrolysis of Cm(NO₃)₃ from isopropyl alcohol onto a 2.2-mg/cm²-thick beryllium backing foil was bombarded by an ¹⁸O beam delivered from the JAERI tandem accelerator. The ¹⁸O⁺ beam passed through a 2.0-mg/cm² HAVAR entrance window foil, 0.09-mg/cm² He cooling gas and the beryllium target backing, and finally entering the ²⁴⁸Cm target material. The beam intensity was 250–300 particle n.a. The reaction products recoiling out of the target were stopped in a volume of He gas (~1 bar) that had been loaded with potassium chloride (KCl) aerosols generated by sublimation from the surface of KCl powder at 640 °C. The products adsorbed onto the surface of the aerosols were swept out of the recoil chamber with the He gas flow (2.0 L/min) and were transported through a Teflon-capillary (2.0 mm i.d., 20 m long) by a He/KCl gas-jet system to MANON. The transported nuclei were deposited on polyethylene terephthalate foils of 120 μg/cm² thickness and 20 mm diameter mounted on the circumference of a stepwise rotating stainless steel wheel that has 80-position collection sites with an 80-cm diameter. The wheel was rotated every 110 s to position the foils between six pairs of Si PIN photodiodes for α-particle detection. Each detector had an active area of 18 × 18 mm². A detection efficiency of approximately 40% for α-particles was achieved. The energy resolution (FWHM) was about 30 keV for the top detectors and 100 keV for the bottom ones.

To determine the gas-jet transport yield, we first collected the recoil of ²⁵⁵Md (T½ = 2.3 min) produced in the reaction ²⁴⁸Cm(¹⁸O, 5n) behind the target in an aluminum catcher foil to get a reference value (100%), while ²⁵⁵Md transported by the gas-jet was collected on a glass filter. ²⁵⁵Fm (T½ = 25 h), the EC decay daughter of ²⁵⁵Md, in both samples was chemically separated and subjected to α-spectroscopy. By comparing the production rate measured after transport through the gas-jet with the absolute production rate from the catcher foil, the transport efficiency was determined to be approximately 35%. The production cross sections were evaluated from the measurement of the mother-daughter correlations of α-particle energies between ²⁵⁵Rf (Eα = 8.28 MeV) and ²⁵⁷No (Eα = 8.22, 8.27, and 8.32 MeV).¹⁷ A photograph of the MANON apparatus is shown in Figure 3.

The sum of α-particle spectra measured in the six top detectors for the production of ²⁶¹Rf at 94-MeV ¹⁸O beam energy is shown in Figure 4. The total beam dose was 2.35 × 10¹⁵. In the α-energy range of 8.10–8.40 MeV, α lines from ²⁶⁰Rf and its daughter ²⁵⁷No are clearly seen. No contributions from other nuclides in this energy window are observed, although there exist several α lines originating from the transfer reaction products from the Pb impurity in the ²⁴⁸Cm target. A total of 166 events were registered both in the top and bottom detectors in the singles measurement and 57 α–α correlation events were detected at this beam energy. Assuming a 100% α-decay branch for both ²⁶⁰Rf and ²⁵⁷No,¹⁷ the production cross sections of ²⁶⁰Rf in this reaction were evaluated to be 8 ± 2, 13 ± 3, and 8 ± 2 nb at the ¹⁸O beam energies of 91, 94, and 99 MeV, respectively. The contribution of the direct production of ²⁵⁷No via the ²⁴⁸Cm(¹⁸O, α5n) reaction is assumed to be negligible in the studied energy region. In Figure 5, the cross sections are plotted as a function of the ¹⁸O bombarding energy together with the literature data,¹⁸,¹⁹ where the relative cross section values in Reference 19 are normalized to the present results. The maximum cross section of about 13 nb at the ¹⁸O energy of 94 MeV was obtained.¹⁴ This results in a production rate of about 2 atoms per min under the given conditions.

Figure 3. View of the MANON (left) and the part of the detectors inside the apparatus (right).

Figure 4. Sum of α-particle spectra measured in the bombardment of the ²⁴⁸Cm target with 94-MeV ¹⁸O ions.

Figure 2. Schematic representation of a target and recoil chamber arrangement with a He/KCl gas-jet system and the apparatus MANON (Measurement system for Alpha-particle and spontaneous fission events ON-line).
3. Aqueous Chemistry of Element 104 Rf

The pioneering works, the first generation experiments, so far performed have concentrated on the question how well the periodic table accommodates the transactinide elements as transition metals in the seventh period. The results with a few atoms have justified placing the elements Rf through element 108, hassium (Hs), into groups 4 to 8 of the periodic table.2 The results with a few atoms and for macro amounts of atoms; however, the first generation experiments still show conflicting results and some of them are criticized due to unsatisfactory experimental conditions.20 There are few convincing data for examining relativistic effects and for considering thermodynamic chemical properties of the transactinide elements even in those of Rf and element 105, dubnium (Db). Therefore, it is of great importance to study detailed chemical properties of the transactinide elements with high statistics and to compare these with properties deduced from extrapolations of periodicity and from modern relativistic molecular orbital calculations: the second generation experiments. In the following, the second generation studies on Rf at JAERI are outlined.

3.1. On-line Production of $^{249}$Rf, $^{169}$Hf, and $^{85}$Zr. As mentioned above, one of our experimental approaches is a detailed comparison of the chemical properties of Rf with those of the expected lighter homologues Hf and Zr. Thus, the chemical experiments on Rf should be conducted together with the homologues under strictly identical conditions. For the on-line production of Rf and the homologues, we used two different targets; one was the mixed target of $^{248}$Cm (610 $\mu$g/cm$^2$ in thickness) and Gd (39.3%-enriched $^{152}$Gd of 36 $\mu$g/cm$^2$ in thickness) prepared by electrodeposition onto a 2.4-mg/cm$^2$ beryllium backing and then, on the resulting natGd (660-$\mu$g/cm$^2$ thickness) mixed target to produce $^{169}$Hf and $^{248}$Cm (18O, 5n) and natGe (18O, x) reactions, respectively. The other was the natGd (370-$\mu$g/cm$^2$ thickness) and Gd (660-$\mu$g/cm$^2$ thickness) mixed target to produce $^{169}$Hf and $^{7.86}$-min $^{85}$Zr through the $^{168}$Gd(18O, x) and $^{169}$Ge(18O, x) reactions, respectively. The $^{168}$Ge was deposited by vacuum evaporation onto a 2.4-mg/cm$^2$ beryllium foil for the simultaneous production of $^{261}$Rf and the homologues $^{85}$Zr. 

3.2. Atom-at-a-time Chemistry of Rf. The chemical study needs to be carried out on phenomena that give the same results for only a few atoms and for macro amounts of atoms; the question arises as to whether a meaningful chemistry with single atoms is possible. For single atoms, the classical law of mass action is no longer valid, because the atom cannot exist in the different chemical forms taking part in the chemical equilibrium at the same time. Guillaumont et al. suggested for single-atom chemistry the introduction of a specific thermodynamic function, the single-particle free enthalpy,21,22 An expression equivalent to the law of mass action is derived, in which activities are replaced by probabilities of finding the species in the appropriate phases. According to this law, an equilibrium constant (distribution coefficient) of the atom between two phases is correctly defined in terms of the probabilities of finding the atom in one phase or the other. If a static partition method is used, this coefficient must be measured in repetitive experiments. Since dynamical partition methods can be considered as repetitive static partitions, the displacement of the atom along the chromatographic column gives a statistical result.2,21 The concept for single atom chemistry is depicted in Figure 6.

For short-lived atoms, the partition equilibrium must be reached during the lifetime of the nuclides, which requires fast reaction kinetics. Based on the kinetics of a single-step exchange reaction, Borg and Dienes suggested that at certain conditions a measurement of the partition of the atom between the phases with very few atoms will already yield an equilibrium constant close to the “true” value provided that both states are rapidly sampled.23 The above interpretation indicates that chromatographic systems with fast kinetics are ideally suited for single atom separation as there is rapid, multiple sampling of the adsorbed or mobile chemical species.2,11

To perform rapid and repetitive chemical experiments with Rf, we employed the apparatus AIDA15,24 that consists of a modified ARCA (Automated Rapid Chemistry Apparatus)25 which is a miniaturized computer controlled liquid chromatography system, and an automated on-line $\alpha$-particle detection system. The schematic drawing of AIDA is given in Figure 7. There are two column magazines, each containing 20 microcolumns. In the modified ARCA as shown in Figure 8, two different paths to supply solutions are available; the first eluent goes through the collection site to the microcolumn, while the second is directed to the column after one-step forward movement of the column magazine to avoid cross-contamination at the collection site.

3.3. Anion-exchange Behavior of Rf in HCl and HNO$_3$. The experimental procedures and the result on the anion-exchange behavior of Rf in HCl solution are summarized here. The reaction procedures recoiling out of the target were transported by the He/KCl gas-jet system to the collection site of AIDA (see Figure 8). After collection for 125 s, the site was mechanically moved to the top of one of the microcolumns, where the products were dissolved with 170 $\mu$L of hot (= 80°C) 11.5 M HCl and were fed onto the 1.6 mm i.d. x 7.0 mm chromatographic column filled with the anion-exchange resin.

Figure 5. Excitation function for the $^{248}$Cm(18O, 5n) reaction. The data taken from References 18 and 19 are also shown.

Figure 6. Concept for single atom chemistry. Courtesy of M. Schädel.
MCI GEL CA08Y (particle size of about 20 µm) at a flow rate of 1.0 mL/min. Then, the products were eluted with 290 µL of HCl through the second path with concentrations between 4.0 M and 9.5 M at a flow rate of 1.1 mL/min. The effluent was collected on a Ta disk as fraction 1 and evaporated to dryness using hot He gas and a halogen heat lamp for α-particle measurement (see Figure 7). The remaining products in the column were eluted with 250 µL of 4.0 M HCl at a flow rate of 1.1 mL/min. The effluent was collected on another Ta disk and evaporated to dryness as fraction 2. The pair of disks were automatically transported to the α-spectroscopy station equipped with eight 600-mm² PIPS (passivated ion-implanted planar silicon) detectors. Counting efficiencies of each detector ranged from 30% to 40% depending on geometric differences of the dried α sources. Alpha-particle energy resolution was 80–200 keV FWHM. After the α-particle measurement, the 493 keV γ-radiation of 169Hf was monitored with Ge detectors to determine the elution behavior of Hf and its chemical yield. The anion-exchange experiments with 85Zr and 169Hf produced from the Ge/Gd mixed target were conducted under the same conditions as those with 261Rf and 169Hf. The effluents were collected in polyethylene tubes and were assayed by γ-ray spectroscopy.

Each separation was accomplished within 20 s and the α-particle measurement was started within 80 s after the collection of the products at the AIDA collection site. The chemical yield of 169Hf including deposition and dissolution efficiencies of the aerosols was approximately 60%. A photograph of the effluent collection part of AIDA is shown in Figure 9.

Figures 10(a) and 10(b) show the typical α spectra observed in the elution with 11.5 M HCl and with 4.0 M HCl, respectively. There are only a few α events from the α decay of 261Rf– 257No at 11.5 M HCl, which means Rf is strongly adsorbed on the anion-exchange resin, while at 4.0 M HCl, Rf is not retained in the column.

In order to evaluate the contribution to the adsorption distribution of 261Rf on the anion-exchange resin of the α events from 257No formed from the α decay of 261Rf during the collection at AIDA before the chemical separation, the adsorption behavior of No on the same anion-exchange resin was examined. On-line experiments with 255No produced in the reaction 248Cm(12C, 5n) were separately conducted according to those with 261Rf in HCl.
and HNO₃ solutions, and it was found that this contribution was negligible. As shown in Figure 10(a), the Cm isotopes, recoil products of the target, the isotopic composition of the ²⁴⁴Cm target is as follows: ²⁴⁴Cm (1.12 at.%) and ²⁴⁶Cm (1.31 at.%), and the Fm isotopes, transfer reaction products from the Cm target, are mostly eluted with 11.5 M HCl. The present result is consistent with the fact that the chloride complex formation of element 104 in 12 M HCl was stronger than that of the trivalent actinides as reported by Hulet et al. The trivalent rare earth element isotopes in the figure, ¹⁵₀,¹⁵₁Dy, the recoil products of the target, the isotopic composition of the trivalent actinides as reported by Hulet et al. Another interesting feature is the adsorption sequence on the anion-exchange resin among Rf, Zr, and Hf. The adsorption order that reflects the strength of the chloride complex formation is Rf ≥ Zr > Hf. The present outcome, however, contradicts the prediction with the relativistic molecular density-functional calculations by Pershina et al. where the sequence of the chloride complex formation is expected to be Zr > Hf > Rf.

In order to obtain information on the chemical species and structure of Rf complexes in HCl, we performed measurements of the extended X-ray absorption fine structure (EXAFS) spectra of Zr and Hf chloride complexes at the KEK (High Energy Accelerator Research Organization) Photon Factory. It has been found that the Zr and Hf complex structure in HCl solution changes from presumably [M(H₂O)₈]⁴⁺ (M = Zr and Hf) to the anionic chloride complex [MCl₆]²⁻ with the increase in the HCl concentration from 9 to 12 M, which is consistent with the results of the anion-exchange experiments. Therefore, we assume the structure of the Rf complex, in analogy to those of Zr and Hf, is the octahedral [RfCl₆]²⁻ in conc. HCl. The difference in the strength of the chloride complex formation of Zr and Hf was also demonstrated; the affinity of the Cl⁻ ion for Zr is higher than that for Hf: [ZrCl₆]²⁻ > [HfCl₆]²⁻, which agrees with the adsorption sequence in Figures 11(a) and 11(b) and the result by Huffman et al. This confirms that the sequence in the chloride complex strength among these elements is Rf ≥ Zr > Hf.

Non Th⁴⁺-like behavior of Rf was also probed with anion-exchange experiments in 8 M HNO₃. From 217 experiments, a total of 20 α events from ²⁶⁰Rf and ²⁵³No were observed including 5 time-correlated α pairs. Although Th⁴⁺ formed anionic complexes and was strongly adsorbed on the anion-exchange resin, Rf was eluted from the column with 8 M HNO₃ as expected for a typical group-4 element as shown in Figure 12. The adsorption value for Th was obtained by batch experiments using radiotracer ²³⁴Th. The above results definitively confirm that Rf is a member of the group-4 elements, but does not resemble the pseudo-homologue Th.

### 3.4. Fluoride Complex Formation of Rf

The experimental procedures with AIDA are basically the same as those described in Section 3.3 and the details are given in Reference 15.
In this study, we used two different microcolumns, 1.6 mm i.d. × 7.0 mm and 1.0 mm i.d. × 3.5 mm, to measure a wide range of $K_d$ values. From the 4226 cycles of the anion-exchange experiments, a total of 266 $\alpha$ events from $^{261}$Rf and its daughter $^{257}$No were registered including 25 time-correlated $\alpha$ pairs of $^{261}$Rf and $^{257}$No. Figures 13(a) and 13(b) show the variation of the adsorption probability of Rf, Zr, and Hf as a function of the initial concentration of HF, $[\text{HF}]_{\text{ini}}$ for the two columns. The adsorption values of Zr and Hf were also measured in separate experiments using $^{89m}$Zr and $^{167}$Hf produced in the $^{89}$Y(p, n) and $^{154}$Eu(19F, n) reactions, respectively. The contribution of the $\alpha$ events from the daughter nucleus $^{257}$No formed from the $\alpha$ decay of $^{261}$Rf before the chemical separation procedures was also evaluated to be less than 5% in HF solution. As shown in Figure 13, the adsorption values of Zr and Hf obtained in the different experiments agree well, and the adsorption of Zr and Hf is absolutely equal over a wide range of HF concentrations and steeply decreases agree well, and the adsorption of Zr and Hf is absolutely equal over a wide range of HF concentrations and steeply decreases as, $\log [\text{HF}]_{\text{ini}} / \text{M}$.

The present result is consistent with those reported in References 34–37 where a different behavior of Rf as compared to that of the homologues in the fluoride complex formation was observed. It is well known that the dissociation of HF is as follows,

$$\begin{align*}
H^+ + F^- &\rightarrow HF \\
HF + F^- &\rightarrow HF_2^-
\end{align*}$$

According to the dissociation constants $K_1 = 935 \text{ M}^{-1}$ and $K_2 = 3.12 \text{ M}^{-1}$ for eqs (1) and (2), respectively, above 1 M $[\text{HF}]_{\text{ini}}$, the concentration of the anionic $HF_2^-$ species is more than one order of magnitude higher than that of the free $F^-$. Thus, the $K_d$ values for Rf, Zr, and Hf in Figure 14 are plotted as a function of the concentration of $HF_2^-$. The $K_d$ values for Rf were deduced from the adsorption probability in Figure 13 by assuming that the kinetics in the fluoride complex formation and the anion-exchange processes of Rf are as fast as those of the homologues Zr and Hf as described in Reference 16, while those for Zr and Hf were from the on-line column chromatographic experiments with AIDA. A decrease of the $K_d$ values for these elements with $[\text{HF}_2^-]$ is explained as the displacement of the metal complex from the binding sites of the resin by the counter anion $HF_2^-$ as,

$$R^-\text{MF}_{4+} + nHF_2^- \rightleftharpoons nR^-HF_2^- + \text{MF}_{4+}$$

where R denotes the resin. As shown in Figure 14, the $K_d$ values of these elements decrease linearly with $[\text{HF}_2^-]$ in the log $K_d$ vs. $\log [\text{HF}_2^-]$ plot. It should be noted that the slopes for Zr and Hf are clearly $-3$ (dashed line), while that for Rf is significantly different, i.e. $-2$ (solid line). Equation 3 implies then that Rf is likely to be present as the hexafluoro complex, $[\text{RF}_{4+}]$ similar to the well known $[\text{ZrF}_{4+}]$ and $[\text{HF}_{4+}]$ at lower $[\text{HF}]_{\text{ini}}$, while Zr and Hf are likely to be present in the form of the heptafluoro complexes, $[\text{ZrF}_{7}]^3-$ and $[\text{HF}_{7}]^3-$ as suggested in References 39 and 40. The activity coefficients of the involved chemical species, however, are not available in the case of the higher concentration of HF, thus the definite identification of the anionic fluoride complexes is presently an open question.

Recently, we have studied the anion-exchange behavior of Rf in the mixed HF/HNO$_3$ solutions to further the understanding of...
the fluoride complex formation of Rf. The $K_d$ values for Rf and the homologues Zr and Hf have been measured as a function of the fluoride ion concentration $[F^-]$. The results indicate that at constant concentration of the nitrate ion $[NO_3^-] = 0.01 \, M$, the formation of the hexafluoro anionic complexes of Zr and Hf occurs at a fluoride ion concentration of about $10^{-5} \, M$, while that of Rf started at around $10^{-3} \, M$. There was about two-orders of magnitude difference in the fluoride ion concentration of Rf and the homologues for the formation of the hexafluoro complexes. This also clearly demonstrates that the formation of the fluoride complexes of Rf is much weaker than those of the homologues Zr and Hf.

4. Summary and Perspectives

Anion-exchange behavior of rutherfordium $^{260}$Rf produced in the $^{248}$Cm/$^{18}$O, $^{9}$Sn reaction was systematically studied based on an atom-at-a-time scale. The adsorption probabilities of Rf in anion-exchange chromatography were measured with high statistics, which has made it possible to perform a detailed comparison of the properties of Rf with those of the homologues Zr and Hf. It was found that the anion-exchange behavior of Rf is quite similar to those of Zr and Hf in HNO$_3$ and HCl solutions, clearly confirming that Rf is a member of the group-4 elements, and Rf does not behave like the tetravalent pseudo-homologue Th. On the other hand, a large difference in the adsorption behavior of Rf and the homologues on the anion-exchange resin was observed in HF solution; the fluoride complex formation of Rf is much weaker than those of the homologues. According to the HSAB (Hard and Soft Acids and Bases) concept, the fluoride anion is a hard anion and interacts stronger with (hard) small cations. Thus, a weaker fluoride complex formation of Rf as compared to Zr and Hf would be reasonable if the size of the RF$^+$ ion is larger than those of Zr$^{4+}$ and Hf$^{4+}$ (0.08 nm vs. 0.07 nm) as predicted by Johnson and Frickie. The fact that the chloride complex formation of Rf is slightly stronger than those of the homologues would also be explained by this concept, because the chloride anion is a relatively soft anion. (The different adsorption behavior of Zr and Hf in HCl is not interpreted with this concept, since the radii of Zr$^{4+}$ and Hf$^{4+}$ are known to be very similar.)

In further studies of Rf aqueous chemistry, experiments on reversed-phase extraction chromatography with AIDA are under way to study complex formation ability of Rf with various kinds of organic extractants. Also experiments are being performed to obtain some thermodynamic equilibrium constants of Rf in fluoride complex formation. On the basis of these quantitative data, the influence of relativistic effects on chemical properties of Rf will be discussed.

Anion-exchange experiments with $^{34}$s-$^{36}$Db produced in the $^{248}$Cm/$^{18}$F, $^{9}$Sn reaction with the cross section of 1.3 nb are also being conducted with AIDA. To obtain more accurate data, however, we need to develop a new device to shorten the time for the preparation of the $\alpha$ sources. An improved AIDA based on a continuous ion-exchange system is under construction.

Acknowledgment. The research work reported was carried out by many colleagues that make up the Heavy Element Nuclear Chemistry Group at JAERI. The list includes K. Akiyama, M. Asai, S. Goto, H. Haba, M. Hirata, S. Ichikawa, Y. Ishii, I. Nishinaka, M. Sakama, T. K. Sato, A. Toyoshima, and K. Tsukada. The present study is supported in part by the JAERI-University Collaboration Research Program and by the JAERI on the Scientific Cooperation between JAERI and GSI in Research and Development in the Field of Ion Beam Application. We would like to thank all colleagues participated in the experiment and contributed to this study from Wako, Niigata, Osaka, Kanazawa, Tokyo, Tsukuba, Shizuoka, Tokushima, Darmstadt, Mainz, Bern/Villigen, and Munich. We are grateful to M. Schädel for his continuous help and valuable suggestions from the beginning of this work. We also acknowledge W. Brichte, H. W. Gügeler, H. Kudo, J. V. Kratz, Y. Oura, A. Shinohara, K. Sueki, A. Türler, and A. Yokoyama for their kind and constant support. Our thanks go to the staff of the JAERI tandem accelerator, Advanced Science Research Center, and Nuclear Physics Group at JAERI, and other all colleagues for their continuous help. The author appreciates kindness of P. J. Karol for reading and correcting the manuscript. Last, he is sincerely indebted to H. Nakahara for his thoughtful and constant advice and encouragement throughout the course of this work.

References

(12) M. Schädel, J. Nucl. Radiochem. Sci. 3, 113 (2002); M.


(35) M. Hirata, private communication.


