Chromatographic Zinc Isotope Separation by Chelating Exchange Resin

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Abstract

Zinc isotope separations were studied by displacement chromatography using the chelating properties of malate, citrate and lactate exchange resin and EDTA as ligands. After each chromatographic operation, the heavier zinc isotopes were found to preferentially fractionated into the carboxylate complex solution phase. The separation coefficients (\( \varepsilon \)) for zinc isotope separation had the largest value and were obtained for the isotopic pairs \(^{68}\text{Zn}/^{64}\text{Zn} (7.16 \times 10^{-4})\) and \(^{66}\text{Zn}/^{64}\text{Zn} (3.08 \times 10^{-4})\), respectively, at 298 ± 1 K. The separation coefficient per unit mass differences (\( \varepsilon/\Delta M \)) for the isotopic pair of \(^{68}\text{Zn}/^{64}\text{Zn} \) was found to range around \( 1.55 \times 10^{-4} \).

Keywords

Zinc isotope separation
Chelating exchange resin
Separation coefficient
Separation coefficient per unit mass differences

Introduction

Zinc is considered as a useful material for cooling water treatment in light water nuclear power plants. A trace amount of soluble zinc injected into the coolant water suppresses \(^{60}\text{Co} \) buildup in water piping systems and reduces the gamma ray dose rate from the piping system carrying the coolant [1–6] and this method has been successfully applied to several nuclear reactors that were used for power generation. However, the use of natural zinc does not provide the maximum benefit due to the adverse effects that are brought about by an increase in radioactive \(^{65}\text{Zn} \) levels caused by the neutron activation of \(^{64}\text{Zn} \) in the reactor core. The use of depleted zinc, i.e., depleted in isotope \(^{64}\text{Zn} \), was found to be more effective to reduce this radiation dose in the cooling water system. The separation of zinc isotopes is therefore of high interest within nuclear science and technology.

Nowadays, zinc isotopes can be obtained using the most advanced centrifuge technology in the world, but this method requires a relatively large amount of energy and therefore has economic restrictions. The present technique could not meet the increasing demands of the \(^{64}\text{Zn} \) depleted zinc isotopes in the world. Compared to this physical method, chemical exchange methods utilizing equilibrium processes have been considered to require less energy consumption and present more economical and environmental clean advantages. The use of chromatographic techniques for the enrichment of isotopes has been regarded as a promising method for large scale production of enriched isotopes. Chromatography operated in the band displacement manner was found to be an efficient process and was characterized by maintaining the self-sharpening band boundaries at both migration band ends. Isotope separation based on ion exchange involves equilibrium formation between the stationary phase resin and the mobile phase solution [7]. The ion exchange displacement chromatography has been successfully applied to the
separation of isotopes of various elements in complex formation systems especially for those using hydroxycarboxylates as ligands [8–12]. Malate, citrate, lactate and EDTA are important ligands that can form a wide variety of complex species with metal ions [13, 14]. Chelating exchange resins have a relatively large zinc adsorption capacity and may be caused by the functional group of the chelating resin [15]. The present study continued the effort to carry out zinc isotope separation by the use of chelating exchange resin. Chromatographic experiments were designed using the four ligands mentioned above in order to elucidate the isotope effect present during zinc complexation. Emphasis was placed on the impact of ligands on the separation coefficient (ε) and the separation coefficient per unit mass differences (ε/ΔM) observed during the separation processes.

### Experimental

#### Chromatographic Process

The chromatographic system used in the present study consisted of five Pyrex columns (100 cm long and 1.0 cm i.d.) and a high pressure pump (Nihon Seimitsu NP-KX-100) that was used for controlling the flow rate of the feeding solution. Columns were connected in series with a Teflon tube. In order to monitor the column pressure, a pressure gauge with a safety device was placed between the first column and the pump. The temperature was set at 298 ± 1 K throughout the experiments by circulating thermostatted water through an extension water jacket out of the columns. An automatic fraction collector (FRC-2100, Iwaki Asahi Techno Glass, Tokyo, Japan) was used for collecting the effluents into small fractions (ca. 1.0 mL). The columns were packed with the commercially available chelating exchange resin D850 (Hangzhou Zhengguang Resin Co., Hangzhou, China). The functional group of this macroporous exchange resin was based on –CH2–N(CH2COOH)2 and the particle size of the resin was 0.4–0.6 mm. At the beginning of the chromatographic operation packed column was conditioned by elution with 2.0 mol L−1 HCl solution in order to remove impurities followed by redistilled water. A 0.1 mol L−1 ZnCl2 with 0.01 mol L−1 HCl solution were used as feeding solution until the length of the zinc adsorption band became ca. 50 cm long. The zinc band was then eluted at a flow rate of 6.0 ± 0.1 mL h−1 using 0.1 mol L−1 ligand solutions. These were prepared from malic acid, citric acid, lactic acid, and EDTA, with ammonia solution, respectively. The eluting solutions were adjusted to each corresponding pH value and transferred to the top of the column. After collection of the zinc fractions the resin was regenerated to the H+ form using 2.0 mol L−1 HCl solution in order to continuously run the five packed columns. The operating conditions for the zinc isotope separation is summarized in Table 1.

### Table 1. Chromatographic experimental conditions of zinc isotope fractionation using different ligands

| Complex system | Zn–EDTA | Zn–malate | Zn–citrate | Zn–lactate |
|----------------|---------|-----------|------------|------------|
| Eluting agent  | (NH4)2–EDTA | (NH4)2–malate | (NH4)2–citrate | NH4–lactate |
| (0.1 mol L−1, pH = 7.5 ± 0.2) | (0.1 mol L−1, pH = 7.4 ± 0.2) | (0.1 mol L−1, pH = 7.2 ± 0.2) | (0.1 mol L−1, pH = 7.3 ± 0.2) |
| Resin          | Macroporous, particle size 0.4–0.6 mm, total exchange capacity ≥ 1.8 mmol mL−1, water content 48–56%, –CH2–N(CH2COOH)2 functional group | | | |
| Temperature (K) | 298 ± 1 | | | |
| Feed solution  | 0.1 mol L−1 ZnCl2 + 0.01 mol L−1 HCl solution | | | |
| Migration length (m) | 9.5 | | | |
| Flow rate (mL h−1) | 6.0 ± 0.1 | | | |

#### Chemical and Isotopic Analysis

The zinc concentrations obtained from each fraction was measured by an atomic adsorption spectrophotometer (ANAX-182F type) using the emission line at 213.9 nm and sensitivity calibration was performed before measurements. The mass determination of the feed solution and fraction samples taken from the front band region were performed by inductively coupled plasma mass spectrometry (ICP-MS, X7 Series) and fraction samples taken from the front band region were performed by inductively coupled plasma mass spectrometry (ICP-MS, X7 Series) with the zinc concentration of 30 ppb diluted with 0.5% HNO3. The dwell time for 64Zn, 66Zn, 68Zn, 69Zn, and 70Zn was 10 ms and for 68Zn and 70Zn was 20 ms. For each isotope measurement, the number of scans were 200 and the measurement error was within 0.2%.

### Results and Discussion

The mechanism of zinc isotope separation by chelating exchange chromatography is illustrated in Fig. 1. Chromatography was conducted in a band displacement manner. At the rear band boundary of zinc ions, the adsorbed zinc ions were released from the resin, as indicated in Eq. 1, and formed complexes with the ligand supplied by the eluent where superscript bars denote the resin phase:

\[
\text{Zn}^{2+} + (\text{NH}_4)_2\text{L} \rightarrow 2\text{NH}_4^+ + \text{ZnL} \quad (1)
\]

During the elution through the zinc adsorption band, the isotope exchange reaction (Eq. 2) took place repeatedly between zinc carboxylate complexes in the external solution phase and the zinc ions on the resin phase:

\[
\text{H}^+\text{Zn}^{2+} + \text{L} \rightarrow \text{ZnL} + \text{H}^+ \quad (2)
\]

where the superscripts H and L denote the heavier and lighter zinc isotopes, respectively.
When the zinc carboxylate complex species reached the front boundary of the zinc adsorption band, the zinc complexes dissociated and the free zinc ions were re-adsorbed onto the resin phase (Eq. 3):

\[ \text{Zn}L + 2\text{H}^+ \rightarrow \text{H}_2L + \text{Zn}^{2+} \] (3)

The results obtained by the above mentioned chemical reactions and isotopic analysis are plotted in Fig. 2.

The zinc concentration found in the effluent fraction corresponded to the front band of zinc adsorption in the columns. The same analyses were performed for the other three ligands that were found to give similar results (not shown). Since the stability constant of zinc complex species is larger than that of ammonium complex species [16, 17] and the total selectivity of the chelating exchange resin for zinc ions is higher than that for protons, the Eq. 1 and Eq. 3 at rear and front boundaries tended to proceed to the right side and resulted in the formation of sharp boundaries at both edges of the band.

It could be obviously found from the profile of isotopic ratios (Fig. 2) that the heavier zinc isotopes were enriched at the front edge and the lighter zinc isotopes enriched at the rear boundary by the present chelating exchange resin. The isotope exchange reaction represented by Eq. 2 concentrated the heavier zinc isotopes in aqueous solution. The results were interpreted by the mechanism that the heavier zinc isotopes were preferentially fractionated into the zinc complex species present in aqueous solution and moved down at a relatively faster velocity to the front edge while the lighter zinc isotopes were retained in the form of zinc ions on the resin and eluted down at a lower velocity, resulting in the accumulation of sharp boundaries at both edges of the band.

The isotope separation coefficient \( \varepsilon \) is used to evaluate chromatographic performance and can be calculated based on Eq. 4 which was derived by Spedding and Kakihana [18, 19]:

\[ \varepsilon = x - 1 = \frac{\sum q_i |R_i - R_o|}{QR_o} \] (4)

where \( x \) refers to the isotope fractionation factor, or the single stage separation factor; \( q \) is the amount of zinc in fraction sample, \( Q \) the total amount of adsorbed zinc on the resin, \( R \) the isotopic percentage of \( ^{64}\text{Zn} \), and the subscripts \( i \) and \( o \) are the fraction number and the original sample, respectively.

The separation coefficients were calculated in each operation from the isotopic enrichment curve of the front boundary and are listed in Table 2.

Each ligand gave distinct \( \varepsilon \) values and EDTA was found to result in the strongest coordination bond with zinc ions as judged by the relatively large \( \varepsilon \) value [20]. When compared to the lactate, citrate and malate ligand systems, the different \( \varepsilon \) value might relate to the complex chemical species in solution. It was reported that
the existence of binuclear complexes of metal ions was probably the reason why the separation coefficient had a large value in the ligand system [21]. In an uranyl carboxylate complex system the existence of binuclear $\text{UO}_2^{2+}$ complexes might create the condition for large isotope fractionation and the separation coefficients followed the order malate > citrate > lactate > tartrate > lactate [22, 23]. On the contrary, the separation coefficients of vanadyl carboxylate system followed the inverse order lactate > citrate > malate [9]. Table 3 presents the separation coefficient $\varepsilon$ and the related data of the isotopic pair of $^{68}\text{Zn}/^{64}\text{Zn}$ obtained in this work along with the results reported from the VO$^{2+}$, Cu$^{2+}$, Gd$^{3+}$ and UO$_2^{2+}$ in malate and EDTA$^{4-}$ ligand exchange systems [9, 10, 24–27].

Regarding the isotopic pair $^{68}\text{Zn}/^{64}\text{Zn}$ (Table 3) the separation coefficients found for lactate and citrate were slightly larger than those observed in a malate system and the $\varepsilon$ value with corresponding ligands obtained in the present work was much larger than that in vanadium system. Apart from the complex structure of zinc and vanadium with those ligands, the large mass difference between $^{68}\text{Zn}$ and $^{64}\text{Zn}$ could also contribute to the formation of a relatively large separation coefficient. Zinc was found to display a much larger separation coefficient when compared with separation coefficients determined for Cu$^{2+}$, Gd$^{3+}$, UO$_2^{2+}$ (Table 3). For a zinc/EDTA ligand a larger value was observed in comparison with the other three ligands and as far as Cu$^{2+}$ and UO$_2^{2+}$ were concerned, the separation coefficients in EDTA were found smaller than those observed in a malate system. The different metal complex structure in ligands and the pH value of the solution used in the present work might have significant impact on results. Further experiments were needed in order to elucidate the mechanism involved in the formation of zinc isotope separation coefficient using different ligand systems.

In general, the isotope effect or the isotope separation coefficient depended on the temperature. Theoretically, $\varepsilon$ is proportional to $\Delta M$ and inversely proportional to the square of both the atomic weight $M$ and the absolute temperature $T$ [26]:

$$\varepsilon \propto \Delta M / M^2 T^2$$  (5)

Table 4 provides an overview of zinc isotope separation coefficient per unit mass differences using a number of different exchange resins [28–31].

It can be seen that each type of resin had different $\varepsilon/\Delta M$ values. The average $\varepsilon/\Delta M$ value for the isotopic pair of $^{68}\text{Zn}/^{64}\text{Zn}$ found in the present study was $1.55 \times 10^{-4}$ which was in agreement with our previous publications using benzo crown resins [31]. When this type of resin is used the cavity size can affect the $\varepsilon/\Delta M$ value significantly, especially when using the benzo-12-crown-4 resin. Benzo-15-crown-5 resin with 0.5 mol L$^{-1}$ zinc chloride was determined to have the largest value (Table 4). It was shown that with the same benzo crown resin and feed solution [30], the $\varepsilon/\Delta M$ value was affected by operation temperature where a high temperature had a small value in alignment with Eq. 5. Ban et al. [12] performed the zinc isotope separation by using a strongly acidic exchange resin and obtained a value for $\varepsilon/\Delta M$ where the isotope effect occurred during cation exchange between the acidic resin and liquid phase. The present study indicated that the determined $\varepsilon/\Delta M$ value was much larger than that of Ban et al. [12] which meant that the present chelating exchange resin might form a more stable binuclear complex with zinc ions and could successfully applied to the zinc isotope separation.

**Conclusion**

Zinc isotope separations were studied by displacement chromatography in malate, citrate, lactate and EDTA systems. The zinc isotope fractionation was confirmed and the lighter isotope was found to be preferentially located on the resin stationary phase. Separation coefficients were calculated for each tested system where the EDTA ligand was found to make a more stable coordination bond with zinc ions that resulted in large $\varepsilon$ values. Polymerization of zinc in ligand complex systems might affect the isotope fractionation. The average $\varepsilon/\Delta M$ value for the isotopic pair of $^{68}\text{Zn}/^{64}\text{Zn}$ was determined to be $1.55 \times 10^{-4}$ which was in agreement with that obtained from benzo crown resins and much larger than

| Table 2. Separation coefficients of zinc in ligand complex systems by chelating exchange resin |
|---------------------------------------------|
| **Zn–EDTA** | **Zn–malate** | **Zn–citrate** | **Zn–lactate** |
|---------------------------------------------|
| Separation coefficient | $^{68}\text{Zn}/^{64}\text{Zn}$ | $^{68}\text{Zn}/^{64}\text{Zn}$ | $^{68}\text{Zn}/^{64}\text{Zn}$ | $^{68}\text{Zn}/^{64}\text{Zn}$ |
| $\varepsilon \times 10^{-4}$ | 7.16 | 5.52 | 6.17 | 5.96 |

| Table 3. Chromatographic separation coefficients of metal ions in ligand complex systems |
|---------------------------------------------|
| **Metal ion** | **Measured isotopic pair** | **Complex formation system** | **T (K)** | **Separation coefficient, $\varepsilon \times 10^{-4}$** | **Reference** |
|---------------------------------------------|
| $\text{Zn}^{2+}$ | 68/64 | Malate | 298 | 5.52 | Present work |
| $\text{VO}^{2+}$ | 50/51 | Malate | 298 | 1.0 | [10] |
| $\text{Cu}^{2+}$ | 63/65 | EDTA$^{4-}$ | 312 | 2.8 | [23] |
| $\text{Gd}^{3+}$ | 158/160 | EDTA$^{4-}$ | 323 | 0.13 | [24] |
| $\text{UO}_2^{2+}$ | 235/238 | EDTA$^{4-}$ | 298 | 0.6 | [27] |
that determined in strongly acidic cation exchange resins.

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**Table 4.** Separation coefficient per unit mass differences (ε/DM) for the isotopic pair of 68Zn/64Zn using different exchange resins

| Resin type | Feed solution | Temperature (K) | 68Zn/64Zn ε/DM (×10^-4) | Reference |
|------------|---------------|----------------|--------------------------|-----------|
| Benzo-12-crown-4 resin | 0.5 mol L⁻¹ ZnCl₂ | 298 | 1.12 | [28] |
| Benzo-15-crown-5 resin | 0.5 mol L⁻¹ ZnCl₂ | 298 | 2.02 | [28, 31] |
| Benzo-18-crown-6 resin | 0.5 mol L⁻¹ ZnCl₂ | 298 | 1.98 | [28] |
| Dibenzo-18-crown-6 resin | 0.5 mol L⁻¹ ZnCl₂ | 298 | 1.60 | [28, 31] |
| Strongly acidic cation exchange resin | 0.05 mol L⁻¹ Zn(NO₃)₂ | 333 | 0.80 | [12] |
| Chelating exchange resin | 0.1 mol L⁻¹ ZnCl₂ with EDTA solution | 298 | 1.79 | Present work |
| | 0.1 mol L⁻¹ ZnCl₂ with malate solution | | 1.38 | |
| | 0.1 mol L⁻¹ ZnCl₂ with citrate solution | | 1.54 | |
| | 0.1 mol L⁻¹ ZnCl₂ with lactate solution | | 1.49 | |