Leaching Behavior of Novel Multi-element Hollandite Ceramic for Immobilizing Radioactive Cesium

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1. Introduction

The structural formula $A_x(B^{3+},B^{14+})_8O_{16}$, defining hollandite, exhibits a distinctive BO₆ octahedral framework. This unique structure is characterized by its ability to form square-shaped tunnels and cubic-like cavities, known as A sites. These sites are adept at housing large ions, for instance, Cs⁺ and Ba²⁺. In the realm of nuclear waste management, titanium-based hollandite ceramics have emerged as a focal point for the encapsulation of the radioactive isotope ¹³⁷Cs, thanks to their inherent structural robustness, thermodynamic consistency, resilience to chemical decay in various environments, and capacity to manage the significant heat output from cesium isotopes. Furthermore, Ti4+ ions are essential for capturing electrons during the β-decay transition of Cs (¹³⁷Cs \rightarrow ¹³⁷Ba + β(e-) and Ti⁴⁺ + β(e-) \rightarrow Ti³⁺) [1 - 4].

¹³⁷Ba + β (e-) and Ti⁴⁺ + β (e-) \rightarrow Ti³⁺) [1 - 4]. Considering the β -decay of ¹³⁷Cs to ¹³⁷Ba over a 30year period, the capability of hollandite structures to securely encase both Cs⁺ and Ba²⁺ for extended timeframes is a crucial research topic. Studies have thus been directed towards the (Ba2+,Cs+)x(B3+,Ti4+)8O16 variant of hollandite, investigating its creation, structural properties, resilience against radiationinduced damage, chemical robustness, and thermodynamic reliability [4 - 7]. However, there is a lack of research on $Cs_x(B^{3+},Ti^{4+})_8O_{16}$ hollandite, in terms of its thermodynamic stability when altering the cesium ratio and incorporating diverse dopants at the Bsite such as Al³⁺, Cr³⁺, Fe³⁺, Ga³⁺, Mn³⁺, and Ti⁴⁺.

We intend to enhance phase stability through the incorporation of five or more elements in roughly equal amounts inspired by the concept of high-entropy alloys. This method is known for generating a high level of configurational entropy, inducing lattice distortion, and producing a synergistic "cocktail effect," which allows certain high-entropy alloys to display outstanding properties such as enhanced mechanical strength, corrosion resistance, stability at elevated temperatures, and resistance to radiation, unlike conventional alloys. Although the B-site sublattice of hollandite does not fully satisfy the criteria for a high-entropy alloy (requiring an entropy above 1.61R), applying highentropy concepts to the design of hollandite could offer significant benefits, including improved thermal stability, resistance to corrosion, and tolerance to radiation. Up until now, there has been a lack of research into hollandite materials that incorporate multiple elements, underscoring the innovative aspect of our study in the development of multi-element hollandite materials.

During long-term disposal, radioactive nuclides may dissolve from the waste form into groundwater. Accordingly, analysis of the leaching behavior of radioactive nuclides in waste form is necessary. The Product Conductivity Test (PCT) method is a standard test method of ASTM for determining chemical durability of glass waste form but it is also widely used for evaluating the leaching property of the ceramic waste form [8]. Therefore, in the present study, leaching behavior of designed multi-element hollandites were measured by the PCT method.

2. Methods and Results

2.1 Synthesis of Multi-element Hollandite

Four multi-element hollandite samples with the compositions Cs_{1.2}Al_{0.2}Fe_{0.3}Ga_{0.4}Mn_{0.3}Ti_{6.8}O₁₆, Cs_{1.4}Al_{0.1} Fe_{0.5}Ga_{0.4}Mn_{0.4}Ti_{6.6}O₁₆, Cs1.5Fe0.8Ga0.4Mn0.3Ti6.5O16, $Cs_{1.6}Fe_{0.9}Ga_{0.3}Mn_{0.4}Ti_{6.4}O_{16}$ were synthesized using a sol-gel route. CsNO₃, Al(NO₃)₃, Fe(NO₃)₃, Ga(NO₃)₃, Mn(NO₃)₂, and titanium ethoxide were used as starting chemicals. The Ti reagents, in the desired stoichiometric proportions, were initially dissolved in anhydrous ethanol within a dry beaker, while the Cs, Al, Fe, Ga, and Mn reagents were separately dissolved in pure water in another beaker. Once the solid powders were completely dissolved, the aqueous solution was gradually introduced into the ethanol-based solution. The resulting mixture was continuously stirred and heated at approximately 120°C for six hours. During this period, the solvent evaporated, leading to the formation of a gel. The gel was subsequently subjected to calcination at 1273 K for two hours to eliminate any remaining organic components [Fig. 3]. The final samples were found to be crystalline, as confirmed by powder X-ray diffraction (XRD) and scanning electron microscopy (SEM) analyses as shown in Fig. 4.

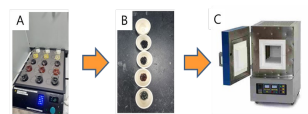


Fig. 3. Synthesis process of multi-element hollandite

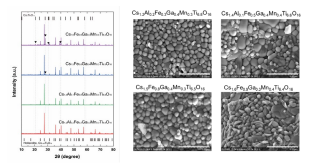


Fig. 4. XRD data and SEM images of multi-element hollandites

2.2 Leaching Test of Multi-element Hollandite

For the PCT procedure, the test specimens were initially ground using an agate mortar, and the resulting powders were passed through 100 to 200 mesh sieves to obtain particles of the desired size. The selected particles were then transferred into the test vessel, which was subsequently filled with fresh ultrapure water. The vessel was securely capped and sealed before being placed in a convection oven preheated to 90°C. After a period of 7 days, the leachate was filtered using a 0.45 μ m syringe filter and analyzed with an inductively coupled plasma mass spectrometer (ICP-MS).

The normalized elemental mass loss (g/m2) based on the relase of element i, NL_i can be calculated using Eq 1, where NC_i is normalized concentration (g/L) of element i, SA is surface area (m2) of specimen and V is the leachate volume (L).

$$\mathrm{NL}_{i} = \frac{\mathrm{NC}_{i}}{SA / V} \tag{1}$$

3. Conclusions

The designed multi-element hollandites were successfully synthesized experimentally using the solgel method. Additionally, the leaching behavior of the produced samples was evaluated through the PCT method.

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REFERENCES

[1] G.C.C. Costa, H. Xu, A. Navrotsky, Thermochemistry of barium hollandites, J. Am. Ceram. Soc. 96 (2013) 1554–1561. [2] V. Aubin-Chevaldonnet, D. Caurant, A. Dannoux, D. Gourier, T. Charpentier, L. Mazerolles, T. Advocat, Preparation and characterization of (Ba,Cs)(M,Ti)8016 (M = Al3+, Fe3+, Ga3+, Cr3+, Sc3+, Mg2+) hollandite ceramics developed for radioactive cesium immobilization, J. Nucl. Mater. 366 (2007) 137–160.

[3] M. Carter, E. Vance, H. Li, Hollandite-rich Ceramic Melts for the Immobilisation of Cs, MRS Online Proc. Libr. 807 (2003). https://doi.org/doi:10.1557/PROC-807-249.

[4] R.W. Cheary, An analysis of the structural characteristics of hollandite compounds, Acta Crystallogr. Sect. B 42 (1986) 229–236.

[5] H. Xu, G.C.C. Costa, C.R. Stanek, A. Navrotsky, Structural behavior of Ba1.24Al2.48Ti5.52O16 hollandite at high temperature: An in situ neutron diffraction study, J. Am. Ceram. Soc. 98 (2015) 255–262.

[6] M. Zhao, Y. Xu, L. Shuller-Nickles, J. Amoroso, A.I. Frenkel, Y. Li, W. Gong, K. Lilova, A. Navrotsky, K.S. Brinkman, Compositional control of radionuclide retention in hollandite-based ceramic waste forms for Cs-immobilization, J. Am. Ceram. Soc. 102 (2019) 4314–4324.

[7] R. Grote, M. Zhao, L. Shuller-Nickles, J. Amoroso, W. Gong, K. Lilova, A. Navrotsky, M. Tang, K.S. Brinkman, Compositional control of tunnel features in hollandite-based ceramics: structure and stability of (Ba,Cs)1.33(Zn,Ti)8O16, J. Mater. Sci. 54 (2019) 1112–1125.

[8] ASTM, C1285-14 Standard Test Mehods for Determining Chemical Durabililty of Nuclear, Hazardus, and Mixed Waste Glasses and Multiphase Glass Ceramics: The Product Consistency Test (PCT), (2014).