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Efficient immobilization of ionic corrosion products by a silica-hydroxyapatite composite via a cold sintering route

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We have successfully demonstrated a new method of radioactive waste immobilization by hosting a waste-bearing form in another waste matrix. A cold sintering route was used to consolidate a silica-incorporated hydroxyapatite (Si-HAp) composite at 200 °C by applying a uniaxial pressure of 500 MPa for a short holding time of 10 min. The higher relative sintered density of up to $98.0 \pm 1.3\%$ was achieved by 25 wt% Si loaded HAp. Results from high resolution X-ray diffraction, micro-hardness, and high resolution scanning electron microscopy confirmed the densification with good mechanical strength (micro-hardness = 2.9 ± 0.3 GPa). For practical applications, two kinds of wastes (25 wt% ionic corrosion product-sorbed EDTA functionalized mesoporous silica and 75 wt% ionic corrosion product-sorbed HAp) were mixed, consolidated and tested. The chemical stability of the solidified composite matrix was positively assessed for low leaching rates of 5.9×10^{-9} to 1.2×10^{-5} g per m² per day using a standard product consistency test. The consolidated composite can bear compressive stress up to 358 MPa, which is orders of magnitude higher than the waste acceptance criteria of 3.5 MPa. The low process temperature can make this sintering process very powerful for the immobilization of radionuclides with volatility and low boiling point. Such a low temperature solidified matrix hosting various wastes may be a promising path for waste management because of its simplicity, reliability, scalability, cost effectiveness and environmental friendliness.

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1. Introduction

The effective management of radioactive waste has been discussed worldwide as a crucial issue for many years. The smart use of a variety of natural and anthropogenic wastes to capture and immobilize radioactive wastes could be an innovative technology to solve the gigantic environmental problems in an efficient and effective manner.^{1–3}

The nuclear power industry has grown strongly in the past 50 years to meet global energy needs. According to the report from International Atomic Energy Agency (IAEA) 2018, about 448 nuclear power reactors with a total installed capacity of 391.721 GWe are in operation worldwide. The spent ion exchange resin, which is produced from nuclear power facilities installed worldwide, is more than 3500 m³ per year.^{1,4,5} The future of radioactive waste (RW) generated as a result of nuclear reactor operations has not been properly and effectively addressed. RW is generated in various ways, such as mining and fuel processing activities, spent fuel assemblies, ion exchange resins (IER),^{6,7} radioisotope production laboratories,⁸ and other

sources (membrane process, evaporator, and filters, *etc.*).⁹ A large volume of radioactive sludge and solutions is produced by various activities related to the decontamination of laundry and working area.

In addition, the decommissioning of nuclear power plants also adds a significant amount of low and intermediate radioactive wastes. The spent ion exchange resin is one of the most secondary waste sources due to its tedious pre-treatments, conditionings and immobilization procedures. Moreover, the nuclear power plants and nuclear accidents such as the Fukushima Daiichi¹⁰ and Chernobyl nuclear power plants have released an enormous amount of contaminated water (⁹⁰Sr and ¹³⁷Cs, *etc.*) that needs immediate and effective treatment and management.^{11–13} There are many solidification methods to solidify low and intermediate level radioactive wastes such as cementation, bituminization, vitrification, polymer and geopolymer encapsulation, incineration, thermal organic reduction process *etc.*^{9,14–16} Ordinary Portland cement (OPC), bitumen (asphalt) and vitrification techniques are widely used to incorporate the spent IER wastes. A conventional solidification route for spent ion exchange resins by being hosted in ordinary Portland cement is outlined in Fig. 1. It is apparent that the conventional solidification flowsheet involves a variety of activities, such as dewatering, watering, drying, mixing of cement and additives, and solidification. Each step during waste conditioning contributes to increase the secondary waste

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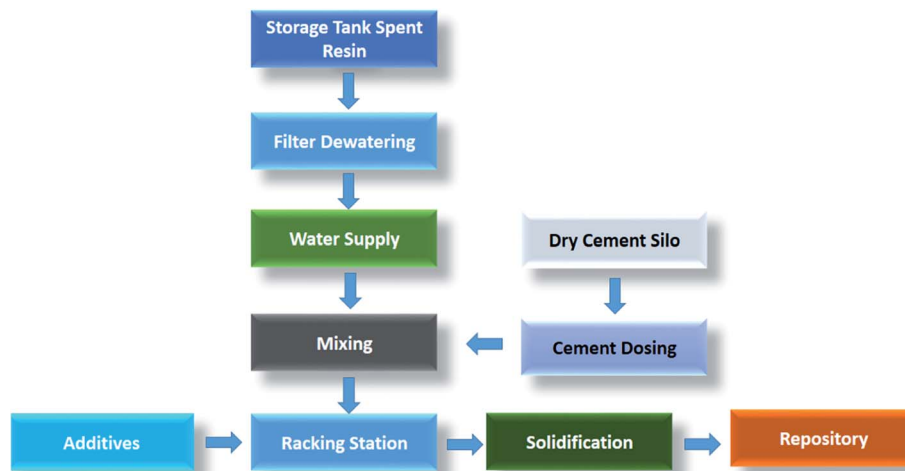


Fig. 1 Flowsheet for the conventional solidification of ion exchange resin with ordinary Portland cement.⁷

volume and extend the process. Therefore, it is imperative to explore a novel route for efficient and rapid waste disposal without adding binders and additives.

All available immobilization techniques have certain advantages and disadvantages associated with them. For instance, OPC has good compressive strength (15–64 MPa) and thermal stability (~ 1000 °C). OPC has been readily available and widely used over the last 40 years. On the other hand, moderate leaching resistance (10^{-5} g per m^2 per day), relatively less waste loading (10–25 wt%), the requirement for binding additives, and humid atmosphere swelling are the problems of the OPC matrix. Bitumen matrix has good leaching resistance (10^{-6} to 10^{-7} g per m^2 per day) and waste loading (25–50 wt%) compared to OPC with low thermal stability (up to 120 °C); however, potential fire risk, restricting its applications in the US and many other countries. The vitrification process has fairly good waste loading (50 wt%), the tendency to chemically bind metal ions into its amorphous network, showing excellent leaching resistance (10^{-8} to 10^{-9} g per m^2 per day) and lowest swelling. Drawbacks of this method, however, are a high processing temperature (>1200 °C) and an exceptional cost. Therefore, novel, efficient, cost-effective and durable materials are required to separate radionuclides and directly immobilize the immense nuclear waste that accumulates in on-site storage facilities.^{2,17}

Here, we present a simplified, single-step radioactive waste management route for the removal of radioactive contaminants by ethylene diamine tetra acetic acid functionalized mesoporous silica (EDTA-Si) and calcium hydroxyapatite (HAP) and sintered to form composite *via* cold sintering for immobilization and storage. The unique property of the proposed matrix is its ability to capture radionuclides and transform itself into a mechanically and chemically durable solid composite matrix without additives. Both constituents of the composite (EDTA-Si and HAP) are readily available and possess ion exchange properties.^{3,18–21} HAP is the most important bioactive material with many natural sources in the form of minerals, phosphate-based rocks, and animal and fish bones.²² It can easily be synthesized

by co-precipitation, sonochemical and hydrothermal methods and is commercially available as well.²³ HAP is widely investigated for its ion exchangeability and its ability to remove a number of ionic species from their aqueous solutions. HAP was also studied and reported for its crystal structure flexibilities to incorporate and hold a number of cationic and anionic species.^{3,24} Silica has been used as a thickening agent to reinforce and improve the mechanical properties of nanomaterials.²⁵ Silica-reinforced calcium hydroxyapatite has been synthesized for dentistry and biomedical applications.²⁶ Silica-incorporated HAP nanoparticles may increase the mechanical strength and chemical stability of nanocomposites due to micro-encapsulation phenomena. In addition, the functionalized silica nanoparticles bear excellent sorption capacities for a number of metal ions (Co, Cu, Ni, Mn, Cr, Zn, Fe, Cs, Sr).^{21,27} Therefore, it may also be possible to consolidate spent silica-based sorbents with spent HAP-based sorbents by mixing them at an optimized weight ratio and converting them into a solid monolith for final waste storage applications. In addition, the use of a spent sorbent (having no significant commercial value) to stabilize/encapsulate another waste at low temperature may have an economic advantage over other conventional solidification techniques.

Many researchers have performed high-temperature sintering (>1000 °C) of silica-substituted HAP to enhance the mechanical properties of composites for medical applications.^{28–31} The immobilization of radioactive wastes by HAP has also been suggested elsewhere.^{3,32} Nevertheless, the sorption, solid-state mixing, and solidification properties of two spent wastes (metal ion captured silica and HAP) are the least investigated. Additionally, high-temperature sintering has certain disadvantages, such as the volatilization of sorbed volatile radionuclides, the dehydroxylation of hydroxyapatite to form unwanted apatite phases (β -TCP), and the high processing cost. Further, the radionuclides captured by organic group functionalized sorbents can be released at high-temperature (>200 °C) due to the degradation of the organic matrix.¹ Recently, Hassan *et al.* reported the cold sintering of pure HAP, and cobalt



sorbed HAP with good density, mechanical and chemical properties.³³

Cold sintering is a densification process of a board set of inorganic powders at extremely low temperatures up to ~ 200 °C using a uniaxial pressing.^{34,35} The initial compaction is achieved by rearrangement of solid particles under mechanical force of the applied pressure. Thereafter, as a result of the combined effect of pressure and temperature, atoms, ions, molecules, and clusters diffuse at the interfaces to reduce the surface free energy and porosity, densely packing the particles together to achieve the maximum density.

The aim of this study is to explore new avenues for radioactive waste disposal by hosting one waste into another to form a low temperature processed waste matrix. There can be certain disadvantages if we immobilize individual sorbents by following conventional methods: such as sintering problems of ionic corrosion products (ICPs) captured EDTA-Si at low temperature, low metal loading capacity of HAP and requirement of the binding matrix (cement, bitumen, glass) to consolidate each individual sorbents. The strategy to incorporate and immobilize spent sorbents by themselves containing ICPs may be promising to enhance the overall waste loadings and reduce the secondary waste volume. Therefore, we designed and investigated the new composite consisting of a combination of spent organic and inorganic type sorbents. Standard mechanical and durability tests were performed to ensure the waste acceptance criteria for the immobilized waste matrix.

2. Experimental

2.1 Chemical procedures

Hydroxyapatite powder was synthesized by a co-precipitation method with a little modification.³⁶ HAP is a well-known sorbent for trapping monovalent and divalent metal ions from aqueous media.^{3,37,38} A detailed sorption study on the optimization of the HAP powder dose to remove a maximum amount of metal ions from a mixture may be found in our previous study.³ Briefly, 40 g L^{-1} HAP powder was found sufficient to remove more than 95% of the metal ions (Co, Cr, Cu, Fe, Mn, Ni, and Zn) with a total concentration of 7 mM at pH 6. In another study, EDTA-Si was synthesized and applied to capture Co, Cu and Ni ions from aqueous solution.²¹ Here, we applied the synthesized EDTA-Si powder ranging from 3 to 9 g L^{-1} to optimize the sorbent dose. More than 95% of the dissolved metal ions were removed as the sorbent dose increased to 9 g L^{-1} at pH 4. The optimized solid-to-liquid ratios for HAP (40 g L^{-1}) and EDTA-Si (9 g L^{-1}) were used for further experiments.

2.2 Sintering methodology for pure and ICPs sorbed Si-HAP powder

We used the synthesized HAP powder as a host matrix for silica-based sorbents. For this purpose, commercial silica nanoparticle (Sigma Aldrich) and synthesized pure HAP powder were initially used to optimize silica loading into the HAP matrix. Fig. 2 shows the schematic diagram of mixing, densifying and

sintering of silica-HAP (Si-HAP). Five samples in triplicate were prepared to take 0, 1, 10, 20 and 25 wt% of silica powder with HAP. Uniform mixing of nano-powders was performed with a 3D mixer at 30 rpm for 12 h using stainless steel balls by mixing the powders with a small amount of ethanol. After mixing, ethanol was evaporated at room temperature, followed by drying the mixed powder under vacuum at 110 °C for 12 h. To make sintered pellets, 1.10 g of powder was put into a specially designed 13 mm stainless steel mold. The mold was wrapped with a heating band and then placed in a Carver press machine at a uniaxial pressure of 500 MPa. The ceramic heating band obtained from Samsung Electronics Co. was used to provide a constant sintering temperature of 200 °C for 10 min. Thereafter, the mold was allowed to cool and the pressure was slowly released before the sample ejection from the mold at room temperature. Optimized cold sintering parameters (applied pressure, sintering temperature, holding time) were applied for sintering of Si-HAP-composite with silica from 1 to 25 wt%.³⁹ A 25 wt% loading of silica powder to HAP was assumed to be the optimum loading among studied conditions because the sintered pellets start breaking during cooling and ejection from the mold at Si loading ≥ 30 wt%. Furthermore, it was not possible to measure the physical and mechanical properties of broken pellets.

After optimizing the pure Si loading into pure HAP matrix, the cold sintering of ICPs sorbed EDTA-Si (ICPs-Si) and ICPs-sorbed HAP composite (25 wt% ICPs-Si and 75 wt% ICPs-HAP) was performed under identical sintering conditions as shown in Fig. 2.

2.3 Characterization

High-resolution XRD patterns were measured with SmartLab, RIGAKU, 2θ scan range from 20 to 60°, the step size of 0.02 at 45 kV and 200 mA at a scan speed of 4° min^{-1} with a Cu K_α incident radiation source of wavelength ($\lambda = 1.54056 \text{ \AA}$) in reflection geometry. The International Center for Diffraction Data Powder Diffraction Files (HAP, ICDD-PDF) 09-0432 were used to identify the crystalline phases.

Attenuated total reflectance-Fourier transform infrared (ATR-FTIR) spectroscopy (Nicolet iS50, Thermo Fisher) was used in the scan range of 4000 to 400 cm^{-1} by a built-in DLATGS ATR detector with a resolution of 4 cm^{-1} . Duplicate measurements were made for each sample using two specimens per sample.

Scanning electron microscopy (SEM) (SU5000 Hitachi) was used to examine the morphology and grain growth. The elemental mapping of the sintered samples was carried out with an Octane Elite EDX detector installed onto an SU5000 SEM device (Hitachi). Prior to SEM analysis, the sintered pellets were broken and mounted on the individual substrate holder using carbon adhesive tape coated with an Osmium Plasma Coater before sputtering.

The hardness of the sintered pellets was measured using a Vickers microindenter (Model: 402MVD, Wolpert Wilson Instruments) under an applied load of 200 gf for 10 s. The upper and lower surfaces of the sintered pellets were polished to $1 \mu\text{m}$ before the hardness measurements. Ten indents were made in



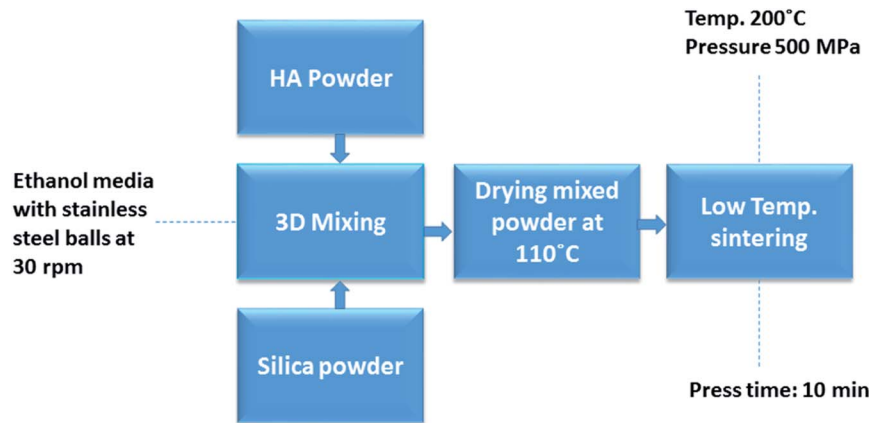


Fig. 2 Schematic of pure Si-HAp and ICPCs-sorbed Si-HAp mixing, compaction, and sintering.

a line passing through a center, and an average hardness value was determined for each pellet.

The bulk density of the samples was measured by Archimedes' method. Sintered pellets were immersed in boiling de-ionized water for 10 min prior to density measurement to remove trapped air. The percentage relative sintered density (% RD) was evaluated using eqn (1).⁴⁰

$$\% \text{RD} = \frac{\rho_b}{\rho_R} \times 100 \quad (1)$$

where ρ_b is the bulk density of the sintered sample measured by Archimedes' method and ρ_R is the theoretical density of the samples. The theoretical density of the samples was normalized and the density values of 2.94 g cm^{-3} and 2.2 g cm^{-3} were used for HAp³ and SiO₂,⁴¹ respectively.

Inductively coupled plasma-optical emission spectroscopy (ICP-OES) was used to determine the concentration of various elements in aqueous solution and sintered samples. The leachate collected during the durability test was also analyzed for the leached quantities of the matrix elements.

Chemical stability is an important parameter to ensure the durability and leach resistance of the developed waste matrix. Different dynamic and static tests developed by the American Society of Testing Materials (ASTM) and Material Characterization Centre (MCC) to investigate the matrix durability in aqueous solutions. In this case, the product consistency test (PCT) devised by ASTM C1285-02 was used under static conditions to measure the normalized leach rates of the constituent elements of the sintered matrix.^{33,42}

3. Results and discussions

XRD patterns were measured to investigate the phase purity and crystal structure of pure HAp sintered samples and silica incorporated HAp sintered samples (Fig. 3). The broad peaks show low crystallinity and amorphous nature of the dried HAp and Si-HAp. It is also known from Fig. 3 that the peaks in pure HAp are identical to those in Si-HAp, suggesting the integrity of the apatite structure even with an increase in the silica dose up to 25 wt%. Moreover, due to the amorphous nature of silica,

XRD peaks only reflect the presence of the pure HAp phase. The phase purity of HAp and HAp containing different silica compositions was confirmed by comparing the XRD peaks with JCPDS PDF No. 09-432.⁴³

FTIR spectra shown in Fig. 4 confirmed the incorporation of silica into the HAp matrix in the range of 1 to 25 wt%. For comparison, two samples were analyzed with 0 wt% silica and 100 wt% silica. The characteristic absorption peaks of the phosphate group (PO_4^{2-}) appear at 566, 603, 963 and 1044 cm^{-1} . A small peak around 875 cm^{-1} (in all samples except

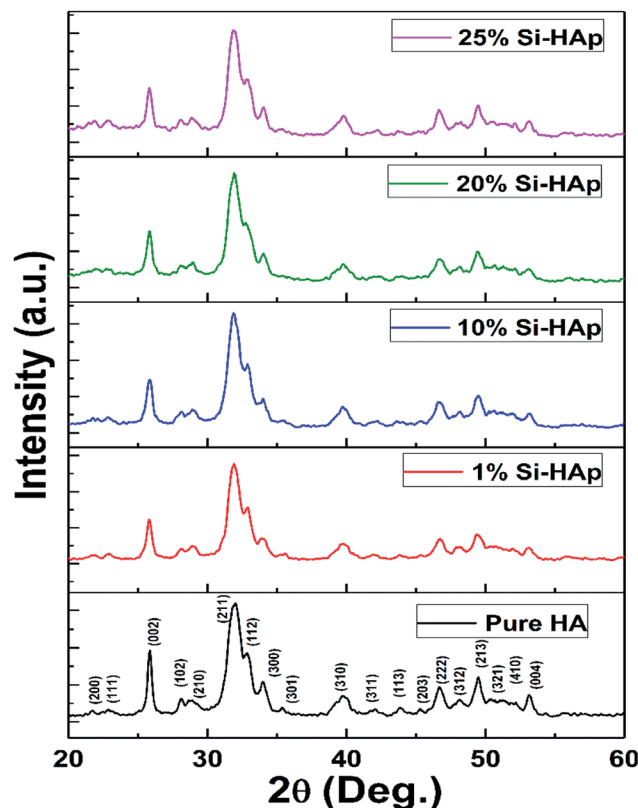


Fig. 3 XRD patterns of the nano-crystalline Si-HAp composite sintered at 200°C , 500 MPa, 10 min in comparison to the pure dried HAp.



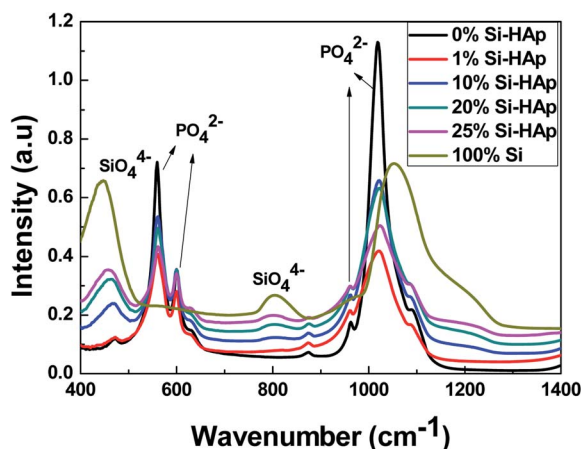


Fig. 4 FTIR spectra of Si-HAp sintered composite in comparison with pure Si and HAp.

pure silica) can be attributed to carbonates. Carbonates can be substituted by phosphates due to the absorption of CO_2 from the environment under alkaline conditions ($\text{pH} = 10.3$) during synthesis.⁴⁴ The appearance of two new peaks at the wavenumbers 444 and 814 cm^{-1} was assigned to the SiO_4^{4-} groups, clearly demonstrating the presence of silica in the HAp matrix,^{43,45} as the peak intensity increased with raising silica incorporation. Gibson *et al.* reported the incorporation of silica at up to 0.4 wt% into apatite synthesized by wet precipitation reaction method.⁴⁶ They proposed a replacement of the phosphate group (PO_4^{2-}) with the silicate (SiO_4^{4-}) group during the synthesis process, which is different from our solid-state mixing of separately synthesized silica and HAp components. Bogya *et al.* reported the existence of non-incorporated amounts of silica in the form of a silica polymer coating on the nano-grains of HAp.²²

During sintering, the additional amount of silica nanoparticles can diffuse at the grain boundaries to form a glassy layer, resulting in densification and good mechanical properties. The silica nanoparticles possess a high surface energy value due to the very small particle size up to 7 nm and the large surface area ($495\text{ m}^2\text{ g}^{-1}$) compared to synthesized HAp ($112\text{ m}^2\text{ g}^{-1}$). These properties also suggest that it is a better candidate for micro-encapsulation when it is in excess. Thus, the good physico-mechanical properties of the consolidated matrix may be due to the silica incorporated HAp structure and micro-encapsulation phenomena.

Fig. 5 shows the fractured surface micrographs of sintered samples of pure HAp and Si-HAp with different wt% of silica. The cold sintering has caused a limited grain growth, and the sintered pure matrix of HAp was composed of nano-grains.⁴⁰ Silica itself is known as a grain growth inhibitor and produces sintered bodies with limited or very small grain growth.⁴⁷ Low temperature sintered Si-HAp composites have shown a homogeneous microstructure in this study due to good 3D mixing technique (Fig. 5).²² The presence of silica might have contributed to achieve good densification. This effect may be attributed to the accumulation and consolidation of SiO_2 along with the

amorphous phases of HAp at the pores under the combined effect of cold sintering parameters (*i.e.* uniaxial pressure and temperature) (Fig. 5a–e).⁴⁸

The EDX mapping of 25 wt% ICPS-Si and 75 wt% ICPS-HAp is shown in Fig. 5f–n. From the images, it is obvious that the sample is highly densified with the homogenized distribution of all the ICPS (Fe, Co, Ni, Cu, Cr, Zn, and Mn).

The effect of silica incorporation on the % RD and micro-hardness of the sintered samples is shown in Fig. 6. The measured physico-mechanical properties of the cold sintered 25 wt% Si-HAp composite confirmed the successful sintering at $200\text{ }^\circ\text{C}$ and 500 MPa within 10 min. Cold sintered samples (25 wt% ICPS-Si and 75 wt% ICPS-HAp) showed a good micro-hardness of $2.9 \pm 0.3\text{ GPa}$. The % RD and micro-hardness results have shown an increasing trend with the increase in wt% of silica. The increase in % RD and hardness may be associated with the condensed packing of porous cavities in HAp by silica diffusion during cold sintering which is a pressure-driven thermo-chemical process. Similar effects of silica incorporation on mechanical properties of concrete have been observed elsewhere at different experimental conditions.^{48,49}

Fig. 7 shows the stress–strain curve for the sintered sample of 25 wt% ICPS-Si and 75 wt% ICPS-HAp composite. From Fig. 7, it is apparent that the consolidated matrix is strong enough to stand the compressive load of up to 358 MPa. The bending and compressive strengths of dense HAp ceramics are in the wide range of 40–250 MPa and 120–900 MPa, respectively.^{50–54} The variation in mechanical properties can be attributed to the statistical nature of the strength distribution, which is affected by the residual microporosity, grain size, impurities, synthesis and sintering route *etc.*⁵⁴

All results discussed in the above-mentioned studies support deformation tendency in HAp, and it behaves differently than the typical ceramics such as aluminum oxide and glass. This load bearing behavior of the HAp matrix can provide the extra mechanical stability of the waste form in the event of sudden mechanical shocks and accidents.

Radioactive waste can be lethal to human and hazardous to the environment and should be stored safely under the multi-layered barrier. The embedded components of the matrix may undergo chemical degradation under extreme environmental conditions, which can cause a leakage of hazardous waste to the environment. Therefore, the standard PCT was performed to evaluate the normalized leach rates (NLRi) for 25 wt% ICPS-Si and 75 wt% ICPS-HAp composite for Ca, Co, Cr, Cu, Fe, Mn, Ni, P, Si and Zn, that are presented in Table 1.

The obtained NLRi values ranged from 5.9×10^{-9} for Fe to $1.2 \times 10^{-5}\text{ g per m}^2\text{ per day}$ for Si revealed good chemical stability of sintered composite for hosted elements. The measured NLRi values for Ca and P were found in close agreement with the literature values.^{55,56} Hassan *et al.* studied the possibility of cobalt immobilization by pure HAp under the identical sintering condition to our study.³³ Their reported NLRi value for cobalt ($2.5 \times 10^{-5}\text{ g per m}^2\text{ per day}$) is very close to our reported value for cobalt ($3.3 \times 10^{-5}\text{ g per m}^2\text{ per day}$). However, the NLRi values for Ca ($1.5 \times 10^{-5}\text{ g per m}^2\text{ per day}$) and P ($3.2 \times 10^{-5}\text{ g per m}^2\text{ per day}$) are higher as compared to our reported



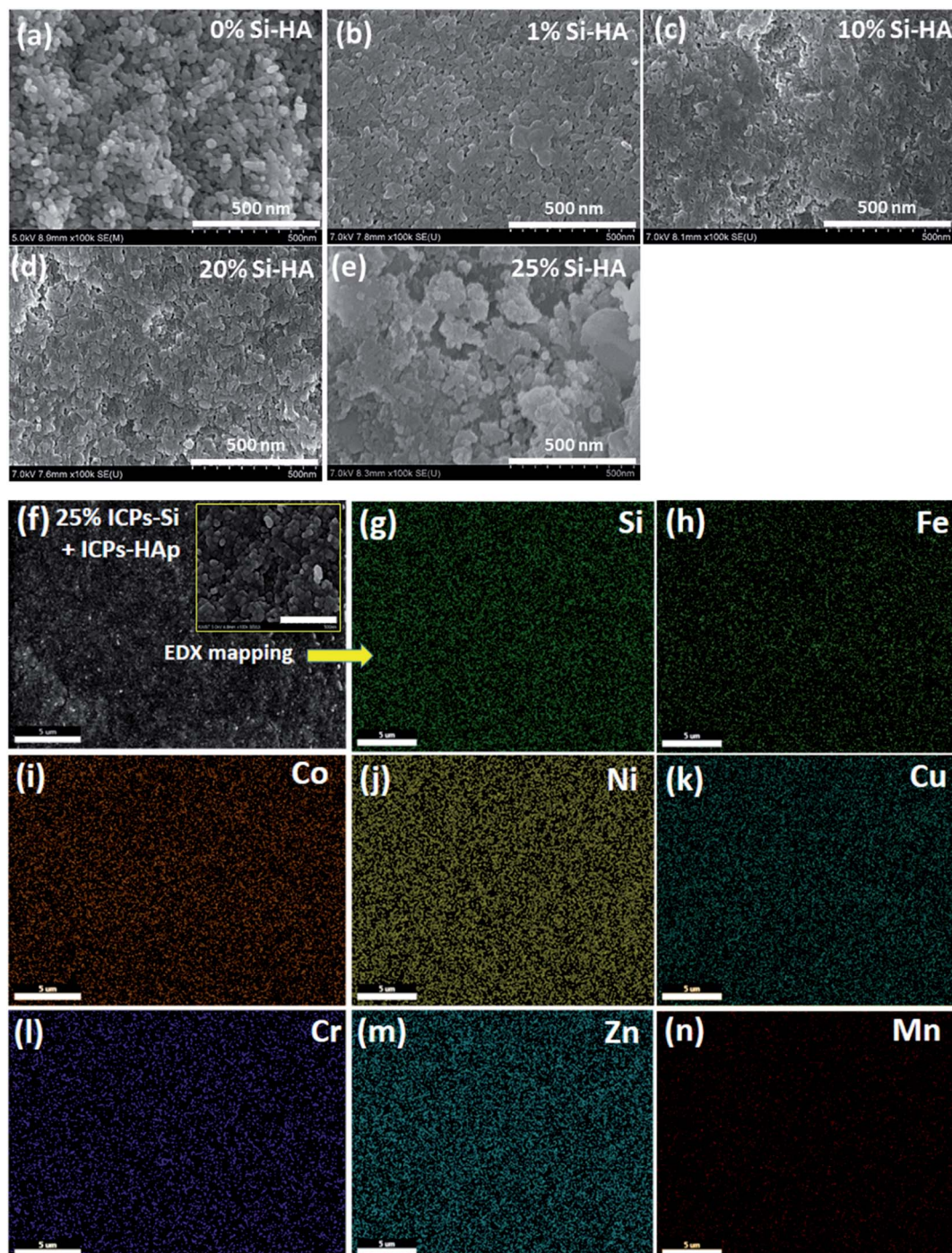


Fig. 5 (a–e) SEM images of pure and silica mixed HAP; (f) EDX elemental mapping showing the distribution and increased amount of silica; images (g–n) show the EDX mapping of 25 wt% ICPs-Si and 75 wt% ICPs-HAP.

composite values. The presence of silica in HAP might have posed an extra barrier by making a silicate-rich layer at a higher sintered density to retain different ions, especially Ca and P.^{45,57} It is hard to make a direct comparison with the limited experimental results in the literature, which were achieved at different compositions, sintering and testing conditions. However, the reported dissolution behavior of iodate substituted HAP, Co-HAP and our measured values are in close agreement to qualify the stability performance.

Based on our previous studies^{33,58} and current results, the reported immobilization route may be used for the solidification of radionuclides by capturing them from the liquid and gaseous wastes. The developed matrix might be used to immobilize gaseous waste (radioiodine) captured by fumed silica, silver functionalized silica or bismuth functionalized silica based gas adsorbents/filters. It is worth mentioning that the reported sintering temperature is very low and is safe for the immobilization of volatile radionuclides such as radioiodine.⁵⁸



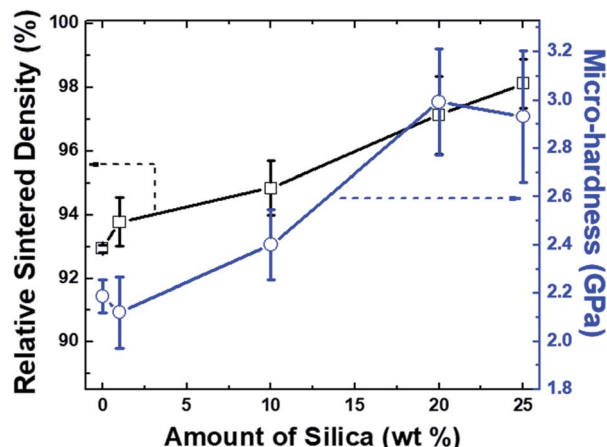


Fig. 6 Effect of silica contents on the relative sintered density (black) and micro-hardness (blue) of HAp; error bars can also be seen at each data point.

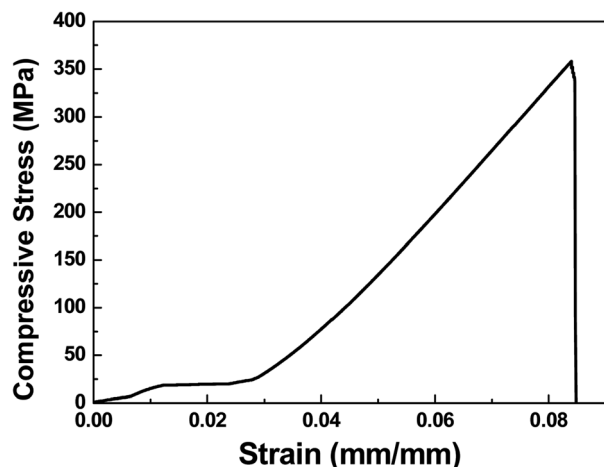


Fig. 7 The stress vs. strain curve of the composite with 25 wt% ICPs-Si and 75 wt% ICPs-HAp. The cross head speed was maintained at 0.1 mm min^{-1} and the load was applied at a cross-section along the z-direction of the cylindrical pellet. The length-to-diameter ratio of the pellets was almost equal to 1.

Table 1 Normalized leach rate (g per m^2 per day) for a waste composite with 25 wt% ICPs-Si and 75 wt% ICPs-HAp

Element	NLRi (g per m^2 per day)
Ca	5.3×10^{-7}
Co	3.3×10^{-5}
Cr	2.0×10^{-6}
Cu	1.9×10^{-5}
Fe	5.9×10^{-9}
Mn	1.4×10^{-8}
Ni	2.3×10^{-5}
P	3.1×10^{-7}
Si	1.2×10^{-5}
Zn	5.1×10^{-7}

4. Conclusions

In this paper, we investigated the possible use of waste to accommodate another waste with a view of reducing secondary waste volume. The sintering conditions for the Si-HAp composite have been successfully optimized to a loading of 25 wt% silica. The composite matrix provided excellent micro-hardness ($2.9 \pm 0.3 \text{ GPa}$) with a relative sintered density of $98.0 \pm 1.3\%$ after cold sintering at $200 \text{ }^\circ\text{C}$. The waste matrix revealed excellent structure integrity by showing the compressive strength of 358 MPa. The cold sintering route for immobilization of waste containing 25 wt% ICPs-Si and 75 wt% ICPs-HAp was successfully demonstrated. The durability of the cold sintered composite matrix showed a promising low leach rates (5.9×10^{-9} to $1.2 \times 10^{-5} \text{ g per m}^2 \text{ per day}$) for all matrix elements. Low temperature, fast process time and sintering methods without artificial binders/additive-free make this immobilization route for radioactive waste an environmentally friendly, energy-efficient and simplified process.

Conflicts of interest

There are no conflicts to declare.

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