Sonochemical recovery of uranium from nanosilica-based sorbent and its biohybrid

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ABSTRACT

Use of nanomaterials to remove uranium by adsorption from nuclear wastewater is widely applied, though not much work is focused on the recovery of uranium from the sorbents. The present work reports the recovery of adsorbed uranium from the microstructures of silica nanoparticles (SiO2-M) and its functionalized biohybrid ([BHM], synthesized with Streptococcus lactis cells and SiO2-M, intensified using ultrasound. Effects of temperature, concentration of leachant (nitric acid), sonic intensity, and operating frequency on the recovery as well as kinetics of recovery were thoroughly studied. A comparison with the silent operation demonstrated five and two fold increase due to the use of ultrasound under optimum conditions in the dissolution from SiO2-M and [BHM] respectively. Results of the subsequent adsorption studies using both the sorbents after sonochemical desorption have also been presented with an aim of checking the efficacy of reusing the adsorbent back in wastewater treatment. The SiO2-M and [BHM] adsorbed 69% and 67% of uranium respectively in the second cycle. The adsorption capacity of [BHM] was found to reduce from 92% in the first cycle to 67% due to loss of adsorption sites in the acid treatment. Recovery and reuse of both the nuclear material and the sorbent (with some make up or activation) would ensure an effective nuclear remediation technique, catering to UN’s Sustainable Development Goals.

1. Introduction

Uranium is the most common radionuclide used as fuel in the nuclear power industry and hence it also constitutes the highest proportion in the nuclear wastewater, posing environmental hazard [1-3]. Due to limited reserves of uranium in India, its recovery from unconventional uranium sources viz. wastewater from uranium mines (containing 480–560 ppm of uranium) [4], fertilizer industry (viz. uranium bearing phosphate as raw material) [5,6] and nuclear reprocessing facilities, mill tailings [4], rock phosphate [6,7] or even seawater (having 3 ppb of uranium) [8] is of particular interest. Efforts are being made to recover the uranium from such lean sources to conserve the natural reserves and also to minimize the spread of contamination of nuclear materials in the environment [9]. Chemical precipitation, solvent extraction, cementation, electrochemical oxidation, evaporation and adsorption are a few of the conventional methods used for the removal of uranium from aqueous solution [10-13]. Most of these methods also have some drawbacks such as uneconomical operation, generation of toxic wastes, low efficiency and high energy consumption [12]. Specifically, adsorption is the most efficient and simplest process for removing contaminants from wastewater [11,13,14] though it also generates secondary waste in the form of used sorbents, which is a very significant aspect in nuclear wastewater treatment. If proper methods are developed for efficient recovery from the sorbents, its subsequent re-use in the fuel cycle is possible and in addition sorbent can also be reused. Recycle of uranium will prevent waste of valuable fissile material resource and also reduce the associated environmental concerns.

Recent developments in nanotechnology have paved a way for the use of nanomaterials such as silica nanoparticles [13], graphene oxide [14], titanium oxide [15], carbon nanotubes [16], nanozeolite composites [17], magnetic nanoparticles [18] and nanoalumina [19,20] as effective sorbents. Silica based matrices has gained a lot of interest as
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In the present work, parametric studies and leaching kinetics of uranium from the ²/BHM bio-hybrid (comprising bacterial cells and silica nanoparticles) as well as from spray dried silica microstructures (SiO₂M) were studied. Biological components like bacterial cells are highly sensitive against various chemicals and are mechanically less stable [29]. Thus, this study helps to understand the changes(transformations) happening in physical as well as chemical properties of ²/BHM and SiO₂M during the process of sonochemical leaching of uranium. The work also opens a new dimension in nuclear remediation technique wherein recovery of nuclear materials of interest from bio-hybrids could be achieved, allowing its recycle in the fuel cycle and preventing the generation of secondary wastes. The study further demonstrates the reusability of both the adsorbents (SiO₂M and ²/BHM) after sonication induced recovery. Though recovery of uranium from activated carbon can be achieved easily by burning off the sorbent, it would cause environmental pollution and sorbent recyclability is not be possible demonstrating the importance of the approach presented in current work.

2. Materials and methods

2.1. Materials

Colloidal silica nanoparticle suspension (40 v/v%) was purchased from Visha Chemicals, India. The colloidal solution contains 40 wt% silica, having a specific surface area of 220 m²/g and density as 1.3 g/ml at 25 °C, as specified by the manufacturer. The particle size varied between 12 nm and 17 nm. A stock solution of uranium (500 ppm) was prepared using UO₂(NO₃)₂·6H₂O (Merck, Germany) in double-distilled water. S. lactis cells were overnight grown in MRS Broth and used for further studies. A high concentration of uranium would ensure that the binding sites on the sorbent are saturated with the metal ion, as also reported by Mishra et al. [29]. The concentration of uranium in the mine wastewater is also in the same range confirming the selection of concentration as 500 ppm. Nitric acid (~16 M) was obtained from M/s Sigma Aldrich. All chemicals were of analytical grade.

2.2. Synthesis of SiO₂ microstructures (SiO₂M) and bio-hybrid microstructures (²/BHM)

For the synthesis of spray dried SiO₂ microstructures, 100 mL of silica NP (2 v/v%) solution was prepared and used as feed through the nozzle of spray dryer (LU222, Labultima, India) maintained under a flow of hot air at inlet temperature (140 °C) and flow rate 2 mL min⁻¹. After spray drying, fine dried powder was recovered at 55 °C as the outlet temperature. The spray dried powder was stored at 4 °C and used for further studies.

In the case of ²/BHM, overnight grown S. lactis cells (6 w/v%) were mixed in silica NP (2 v/v%) solution, stirred for 30 min and used as feed for spray dryer operated as mentioned above. The spray dried powder was stored at 4 °C and used for further studies. The details of the synthesis of both SiO₂M and ²/BHM are reported in details, in our earlier publication [29].

2.3. Synthesis of uranium loaded SiO₂ microstructures (SiO₂M) and bio-hybrid microstructures (²/BHM)

For the synthesis of uranium loaded SiO₂M and ²/BHM, SiO₂M (1 g L⁻¹) and ²/BHM (1 g L⁻¹) was mixed separately in uranium solution (100 mg L⁻¹), and agitated on the shaker (MaxQ 4000, Thermo, USA) working at room temperature (298 K) at 100 rpm for 2 h. Solution was centrifuged at 9000 rpm for 5 min (Eppendorf centrifuge, Model-5810R, Germany). The supernatant was used to analyze residual uranium concentration and pellet (uranium loaded SiO₂M and uranium loaded ²/BHM) was stored till further use.

2.4. Leach solutions and experimental setup for dissolution

The resultant uranium adsorbed sorbents (SiO₂M and ²/BHM) were treated with different concentrations of nitric acid (HNO₃) and ultrasonic baths of different frequencies (20 ± 2, 30 ± 3 and 40 ± 3 kHz) have been used. The in-house built ultrasonic cleaning bath has dimensions of height, width and depth as 240 mm × 140 mm × 60 mm, respectively (Fig. 1). Two transducers (M/s Morgan Matrac make) of size 2” each operating at the same frequency were attached to the base of the tank. Uranium loaded sorbents (SiO₂M and ²/BHM) taken in a beaker with 10 mL of 4 M HNO₃ solution were mounted on a metal basket and dipped in the bath. The acoustic intensity was maintained at 8 W cm⁻² (80 W) in all experiments, except in cases where the effect of ultrasonic intensity was studied. The power dissipation was varied between 20 and 100 W to study the effect of intensity, in order to achieve the requisite acoustic intensity. The ultrasonic bath was filled with distilled water up to a level of 30 mm to provide a medium for sonication. The bath was placed in a fumehood to manage the NOx fumes generated where suitable dilution was provided with air. Dissolution studies were carried out in batches for 30 min each (unless otherwise mentioned). All the experiments were carried out in triplicate, and the mean values have been reported with experimental errors typically between ±3%.

2.5. Analysis of leach liquor samples

The leach liquor samples were withdrawn at equal intervals of time, filtered and analyzed for dissolved uranium concentration. Estimation of uranium was performed using inductively-coupled plasma optical
emission spectroscopy (ICP-OES). In each set, uranium solution (without sorbent) was used as control.

2.6. Characterization of morphology

To study the changes in morphology of the sorbents (SiO$_2$M and BHM) before and after desorption of uranium, Scanning Electron Microscopy (SEM) (Make: M/s Tescan Vega3) analysis was carried out. Energy Dispersive Spectroscopy (EDS) studies were also carried out in parallel using detector from M/s Oxford Instruments. To understand the involvement of functional groups of sorbents, Fourier transform infrared spectroscopy (FTIR) of the sorbents was recorded using Bruker FT-IR Alpha II spectrometer, at a resolution of 4 cm$^{-1}$. Raman spectrometer (M/s W Tec make, Model Alpha 300R) was used to make a comparative study of the uranyl compounds present in the biohybrid before and after sonication.

2.7. Reusability studies after sonication

In order to test the reusability of the nanoparticles, 1 g L$^{-1}$ of sorbent and uranyl nitrate solution of 50 mg L$^{-1}$ concentration (initial pH = 7.0) was agitated on a rotary shaker operating at room temperature (298 K) and 150 rpm for 24 h. The objective was to measure the uranium adsorption capacity on reuse for adsorption.

3. Results and discussion

3.1. Recovery of uranium using only acid (in the absence of ultrasonic field)

The obtained results for recovery of uranium from both SiO$_2$M and BHM are shown in Fig. 2 (a) and Fig. 2 (b) respectively. Nitric acid is widely used for uranium recovery [39]. The extent of dissolution of uranium with 8 M nitric acid at the operating temperature 45 $^\circ$C was less than 6% and 20% in 1 h for both SiO$_2$M and BHM respectively in the absence of ultrasonic field. The concentration of the acid was deliberately taken as 8 M to allow considerable recovery of metal from the microstructures, though results were not very promising in the absence of ultrasound. In addition, Despite the higher adsorption capacity of the BHM compared to the SiO$_2$M, as reported earlier [23], the recovery of uranium from the BHM was faster than SiO$_2$M.

The effect of concentration of the acid on the biohybrid on uranium recovery was studied through Scanning Electron Microscopy (SEM) analysis. In Fig. 3 (a-d), micrographs show the trends for degradation of the biohybrid at different acid concentration at room temperature in the absence of ultrasound. Higher recovery of uranium from the biohybrid compared to the self assembled SiO$_2$M can be explained by the micrographs (Fig. 3 (b)) which show the disintegration of the BHM in acid beyond 4 M acid concentration. At 4 M acid concentration, fissures appear on the surface and beyond 4 M concentration, the biohybrid collapses to a deformed two-dimensional structure (Fig. 3 (c)) from its three-dimensional dough-nut shaped morphology. The biohybrid shows complete disintegration at 16 M acid (Fig. 3 (d)). Similar results were observed in the presence of ultrasound (20 kHz and 40 kHz) thereby establishing acid concentration as the sole reason for the degradation of biohybrid and not ultrasonic cavitation. On sonicating the solution in water, degradation of the cells was not observed. This observation has also been confirmed by earlier reports [40–44] that state that cell lysis is brought about by ultrasound only at very high intensity (>10 W/cm$^2$).

Though the effect of ultrasound on bacterial cells is limited, lysis of tumor cells is reported say using focused ultrasound [45]. However, effect of nitric oxide on cell disintegration is widely reported [46]. Therefore, it may be concluded that at higher nitric acid concentration, the nitric oxide generation leads to the disintegration of the cells more dominantly as compared to ultrasound.

The particle size distribution of the biohybrid in various acid concentrations corroborates the initiation of degradation of the microstructures beyond 4 M acid (Fig. 3 (e)). The particle size of the biohybrid remains intact till 4 M, beyond which it gets skewed towards size <1 μm. Around 7% of the total particle volume is less than 1 μm when treated with 8 M acid and 55% becomes lower in the case of acid of 16 M concentration as depicted schematically in Fig. 4.

Thermal effect on biohybrid was also studied in the absence of ultrasound to confirm degradation of biohybrid microstructure beyond 70 $^\circ$C. The degradation of biohybrid was observed to some extent at 70 $^\circ$C (Fig. 5 (a)), but complete disintegration is seen beyond 100 $^\circ$C (Fig. 5 (b)) which is attributed to the combined effect of acid and temperature on the cells. Therefore, high acid concentration and high temperature intensification of uranium recovery is ruled out for the biohybrid.

3.2. Intensification of recovery using ultrasound

The poor recovery rates of uranium from SiO$_2$M and BHM microstructures necessitated the use of sonochemical intensification. Ultrasound is an interesting alternative for low temperature (<70 $^\circ$C) process intensification. Since the piezoelectric transducers used to generate
Ultrasonic field has a Curie temperature of 120°C, the bath temperature is also kept sufficiently low to prevent damage to the piezoelectric crystal (Lead Zirconate Titanante, PZT, in our case) and the glue bonding the transducer to the reactor bottom. Ultrasound has been demonstrated to successfully increase mass transfer rates, improve yields, and initiate reactions and even change the reaction pathways [34,36,37]. In order to establish the intensification effects of ultrasound in the dissolution and recovery process of uranium from the microstructures, the extent of dissolution at 30 kHz ultrasonic field under similar operating conditions was studied. It is clearly established from Fig. 2 (a) for SiO$_2$M and Fig. 2 (b) for BHM that the ultrasonic dissolution yields five- and two-fold enhancement in the extent of dissolution, respectively. The intensification is caused by the cavitating bubbles that act as microreactors of high temperature and pressure, resulting in faster dissolution of uranium in the acid before their collapse on the surface of the silica substrates. The impact of collapse of bubbles further leads to dislodging of the contaminants from the surface facilitating enhanced recovery.

The micrographs of the SiO$_2$M and BHM before adsorption of the metal, after adsorption and after recovery of metal by sonication have been given in Fig. 6. Fig. 6 (a) shows the microstructure of the silica (SiO$_2$M) before adsorption. The SiO$_2$M after adsorption of uranium is represented in Fig. 6 (b). Fig. 6 (c) shows SiO$_2$M after desorption using 30 kHz ultrasonic field applied at 16 M acid concentration for 30 min. As evident from the micrographs, the SiO$_2$M shows no difference in morphology before and after recovery of uranium. The results clearly confirm that the morphology remains unaffected by the temperature, high acid molarity and the effect of the cavitating bubbles leaching out the adsorbed U from its surface.

On the other hand, the BHM presents a contrasting picture based on the comparison of the native material, after adsorption and finally after desorption based on ultrasound. The morphology of the doughnut shaped self-assembled spray dried structure shows the prominence of cells near the surface in the case of native adsorbent (Fig. 6 (d)). The doughnut shaped structure can be explained by the repulsion of the negatively charged silanol groups on silica and the negatively charged cell wall of the cells, as discussed elsewhere [29]. These cells on the surface gets occupied with uranium due to the adsorption and the prominent imprints of the cells are less visible (Fig. 6 (e)). After sonication treatment using 16 M acid leading to uranium recovery, the biohybrid doughnut structure collapse to a deformed structure (Fig. 6 (f)). This was mainly due to the high acid concentration and not due to ultrasound effects as also discussed earlier and confirmed by dissolving the biohybrid in 16 M acid in the absence of any ultrasonic field. The acid degraded the cells and the three dimensional doughnut structure collapsed, subsequently fragmenting into smaller particles, even in the absence of ultrasound. Therefore, detrimental effect of high acid concentration on the biohybrid is conclusively proved, also confirming that use of ultrasound is not contributing to any negative effects.

Raman spectra of the uranium adsorbed biohybrid before sonication and after sonication using 4 M acid for 30 min are shown in Fig. 7 (a). The uranyl (UO$_2^{2-}$) peaks in the spectra (at 177 and 845 cm$^{-1}$) before sonication reduce in intensity after sonication, indicating leaching of the metal ions from the substrate. The peaks at 745, 1046 and 1400 cm$^{-1}$ are known to be nitrate vibrational modes [47].

The EDS study of the biohybrid before sonication and after sonication using 4 M acid for 30 min is shown in Fig. 7 (b). The study demonstrated a reduction of U content in the biohybrid from 21% (w/w) to 3% (w/w) after 30 min of sonication.
In order to study the effect of operating parameters in the intensified dissolution, the effect of acid concentration, temperature, operating frequency and acoustic intensity were subsequently studied and now discussed.

3.3. Effect of acid concentration on intensified dissolution

The plots given in the Fig. 8 (a) explains the dependence of the dissolution ratios with time at different concentrations of HNO$_3$. Nitric acid is a strong oxidizing agent that oxidizes the U(IV) to U(VI) and it is expected that the rate of oxidation will be dependent on the
concentration. With 2 M HNO$_3$, minimal extents of dissolution were seen for both SiO$_2$M and BFHM which increased almost linearly with the loading of HNO$_3$ at the same temperature of 45 °C for a sonication time of 30 min. As the acid concentration increases, metal recovery from SiO$_2$M shows a linear increase due to faster dissolution in acid. Comparatively, BFHM shows a gradual rise till 12 M acid, beyond which the U recovery increases dominantly which may be attributed to breakdown of the cellular composition in high acid concentration and faster release of the adsorbed uranium into the solution. The same was confirmed by the disintegration and collapse of the doughnut shaped structure as demonstrated in Fig. 2 (f) for the case of 16 M acid strength. Even in the case of U recovery from graphite, too high acid strength was observed to lead to pitting corrosion of the substrate [32,39]. Therefore, though the leaching of uranium from SiO$_2$M would be fastest in 16 M acid but due to the damage to the structure of SiO$_2$M, high release of NO$_x$ gases from the dissolution process and stringent requirement on the
material of construction for the ultrasonic bath, very high concentration of acid is considered to be difficult to handle on a large scale. Though a lower acid concentration is always beneficial, it sacrifices the rate and extent of recovery. At 45 °C and 20 kHz acoustic field, the recovery of uranium is 10% at 2 M and 17% at 4 M acid concentration. Therefore, 4 M acid was recommended for uranium recovery for both SiO\textsubscript{2}M and \textsuperscript{3}BHM.

3.4. Effect of temperature on intensified dissolution

The experiments to study the effect of temperature were carried out using different operating temperatures selected over the range of 40 °C to 60 °C using 4 M HNO\textsubscript{3} solution under 30 ± 3 kHz ultrasonic field with an acoustic intensity of 8 W cm\textsuperscript{-2} (80 W power) applied for 30 min. The results for the effect of temperature on the dissolution of uranium are illustrated in Fig. 8 (b). The experimental results showed that while the temperature of the acid leachant had little effect (a change of less than 5% over the experimental range) on the silica nanoparticles, the dissolution from the \textsuperscript{3}BHM was a more kinetically controlled process. Quantitatively, the extent of dissolution from SiO\textsubscript{2}M only increased from 31% to 39% whereas dissolution from \textsuperscript{3}BHM increased from about 40% to more than 60% with an increase in temperature from 40 °C to 60 °C for the case of 8 M HNO\textsubscript{3} (as seen in Fig. 8 (b)). An increase in the temperature usually increases the kinetic rate of dissolution which is evident in the present case. The ease of generation of cavities also increases due to the increase in the vapour pressure of the solution, also leading to faster dissolution at 8 W cm\textsuperscript{-2} [36,37]. The dampening of the overall sonication effect at higher temperatures due to cushioning effect of the bubbles is not observed in the present study giving a continuous increase in the dissolution rate with an increase in the temperature over the investigated range. Further, the leaching of SiO\textsubscript{2}M was less sensitive to the temperature changes compared to \textsuperscript{3}BHM. This may be explained by the fact that though silica remains unaffected in the studied temperature range, the temperature sensitive constituents (especially proteins) of the cells are affected by the temperature of the leachant. Therefore, the cells lose their binding capacity and release the adsorbed metal faster compared to SiO\textsubscript{2}M. Considering the observed results, the recommended temperature for both SiO\textsubscript{2}M and \textsuperscript{3}BHM would be 60 °C in terms of faster dissolution or 45 °C in terms of better recovery of sorbent. In this work, the studies were carried out at the lowest temperature 45 °C to prevent masking of the effects of other parameters by the thermal effects. It is interesting to note that the effect of temperature on the dissolution from microstructures was similar to the trend observed in case of uranium recovery from the pores of graphite [24].

3.5. Effect of operating frequency of ultrasound

The available transducers of frequencies as 20 kHz, 30 kHz and 40 kHz (in the low frequency range) were used to characterize the effect of frequency on uranium recovery. The acoustic intensity maintained for all the operating frequency was fixed at 8 W cm\textsuperscript{-2} (80 W as the actual supplied power). The dissolution tests for different frequencies were
conducted using 8 M acid at 40 °C. It was observed that the recovery of uranium from SiO₂M decreases with an increase in the operating frequency. The recovery was 40% at 20 kHz as the frequency, 30% at 30 kHz as the frequency and 25% at 40 kHz frequency (Fig. 8 (c)). At lower frequencies, the bubbles formed are larger, collapse violently and hence they render higher intensity mechanical effect on the substrate to dislodge the contaminants. The strong cavitation effect assisting the reaction with the acid dislodges the uranium from the surface of SiO₂M at a faster rate. Based on the obtained results, the recommended operating frequency for SiO₂M is 20 kHz. Similar trend was also reported for the case of uranium and yttria recovery from graphite substrate [32,39].

In the case of BHM, the recovery showed hardly any change within the tested frequency range. The recovery of uranium from BHM is more sensitive to chemical effects than mechanical effects of ultrasound. Therefore, no significant change was observed over the narrow range of frequency tested.

3.6. Effect of acoustic intensity

The acoustic intensity must exceed a certain threshold intensity to induce cavitation at desirable intensity and bring out the desired intensification. The obtained data for effect of acoustic intensity on the recovery in the case of SiO₂M and BHM is shown in Fig. 8 (d). It is clearly established that the metal recovery from SiO₂M increases linearly with an increase in the applied acoustic intensity. The intensification in the reaction rate is a strong function of input power as increase in acoustic intensity increases the cavitation bubble density. These bubbles act as microreactors of high temperature and pressure and accelerate the dissolution process. Increase in acoustic intensity also increases the mechanical effect of ultrasound, as bubbles formation and subsequent implosion increases in turn accelerating the dissolution process.

It was also observed that the metal recovery from the BHM is faster giving higher extent of recovery compared to that obtained in the case of SiO₂M. It is also important to note that at much higher vibrational amplitude, a large number of bubbles are generated which build up near the source of ultrasound and act as barrier in the transfer of acoustic energy to the liquid. This phenomenon is called source decoupling leading to a lowering of the effect of cavitation. The trend was clearly seen for the case of BHM where the recovery was seen saturating beyond 5 W cm⁻² for an operating frequency of 20 kHz in the tested intensity range. Beyond the acoustic intensity of 5 W cm⁻² not much difference in the extent of recovery is seen. Interestingly, no such decoupling or cushioning effects were observed for SiO₂M over the range of power intensity investigated in the current work. Lower extents of recovery in the case of SiO₂M can be one of the reasons for the observed continuous increase in the extent of dissolution meaning that there is still some way to go before saturation is seen. In the case of recovery of uranium from graphite pores [31], the extent of dissolution also reached a peak indicating that the trends are dependent on the specific substrate materials making the study presented in the current work important.

3.7. Recyclability of the silica substrates after sonication

As discussed in section 3.2, the SiO₂M retains its original shape and size after sonication and appears to be less affected by the combined action of cavitation and acid compared to its biohybrid. Considering this analysis, SiO₂M was used for adsorption studies after ultrasound assisted leaching just to confirm the reusability. It was found that the SiO₂M adsorbs 69 ± 7% of uranyl nitrate solution after 24 h of agitation. The FTIR spectrum of the SiO₂M after U adsorption confirms the presence of Si-O-Si bond at ~1090 cm⁻¹ and Si-OH bond at 900 cm⁻¹ (Fig. 9 (a)). The broad centered peak at 3200 to 3500 cm⁻¹ was assigned to OH stretch. The peak around 600–800 cm⁻¹ may be attributed to the presence of UO₂. Thus it can be clearly said that SiO₂M is effectively reused for uranium adsorption.

Reusability studies on biohybrid treated with 1 M acid was found to yield removal of 60 ± 3% of the uranyl nitrate mother liquor, while biohybrid sonicated in 4 M acid adsorbs 58 ± 2%. These values are less than 92 ± 2% uranium uptake, earlier reported [29] which may be attributed to the loss of adsorption sites due to the acid treatment.

The reusability studies on biohybrid treated beyond 4 M acid were not repeatable. The FTIR study of the BHM before sonication treatment has been reported by Mishra et al. [29]. Comparing the FTIR spectra of BHM after sonication in 20 kHz ultrasonic field (Fig. 9 (b)) using 16 M acid (without adsorbing U) with that before sonication, we find that the Si-O-Si group (~1090 cm⁻¹) and the C = O group (~1700 cm⁻¹) are missing after treatment. The only detectable silica signature was at 900 cm⁻¹ as the asymmetric vibration of Si-OH. This change may be attributed to the reaction of acid and silanol groups after the breakdown of the biohybrid. The –CH and C = C stretch at ~1400 cm⁻¹, –NH and NO₂ stretch at ~1550 cm⁻¹, =C-H, –CH around 3100 cm⁻¹, and –OH and –NH stretch at ~3200–3500 cm⁻¹ (due to water and trapped acid) are the few other dominant peaks observed. It can be concluded that due
to the structural deformation and disintegration of the \( ^6\)BHM mainly in high acid concentration, the major adsorption sites like Si-O-Si, C=O, were missing. Therefore, the reusability studies on the recycled \( ^6\)BHM beyond 4 M acid concentration were not repeatable and yielded quite low adsorptive removals of uranium.

4. Modelling of the dissolution kinetics

A kinetic study of the dissolution process was carried out based on the well-known shrinking core model and all three possible mechanisms as film diffusion control, surface reaction control and product layer diffusion control were studied [31] for possible fitting in the current work. The shrinking core model assumes that the reaction proceeds at

Fig. 10. Comparison of the control mechanisms governing the uranium recovery from (a) SiO\(_2\)M (b) \(^6\)BHM.
the solid–liquid interface which moves into the solid core, that remains unreacted. Each of the controlling mechanism based model equation can be expressed as shown in Eqs (1–3):

\[ x = k_d t \] for film diffusion control, 

\[ 1 - (1 - x)^{1/2} = k_d t \] for surface reaction control

\[ 1 + 2(1-x)-3(1-x)^{2/3} = k_d t \] for product layer diffusion control

Where \( x \) is the reacted fraction at time \( t \) and \( k_d, k_s, k_0 \) are apparent rate constants expressed as in Eqs (4)–(6):

\[ K_F = \frac{3bC_{\text{HNO}_3}}{\rho} \]  \hspace{1cm} (4) 

\[ k_s = \frac{bC_{\text{HNO}_3}}{\rho \rho_0} \]  \hspace{1cm} (5) 

\[ k_0 = \frac{2bMD_{\text{HNO}_3}}{\rho \rho_0^2} \]  \hspace{1cm} (6)

In Eqs (4)–(6), \( b \) is the stoichiometric coefficient; \( M \) is the molecular weight of the reacted substance; \( \rho \) is the density of the reacted particle; \( \rho_0 \) is the initial particle radius; \( k \) is the intrinsic rate constant; \( D_0 \) is the effective diffusivity and \( C_{\text{HNO}_3} \) is the bulk concentration of \( \text{HNO}_3 \). The fitting results for the shrinking core model for \( \text{SiO}_2\text{M} \) and \( \text{SiO}_2\text{M} \), respectively. Comparing the figures, it can be seen that the product layer diffusion control model is the best fit with correlation coefficient of 0.973 and standard deviation 0.0084 rather than surface reaction control model (correlation coefficient 0.9693 and standard deviation 0.0119) or film diffusion model (correlation coefficient 0.9679 and standard deviation 0.023) for both adsorbents as \( \text{SiO}_2\text{M} \) and \( \text{SiO}_2\text{M} \). The conclusion of best fitting model is based on the relative values of the obtained correlation coefficients (highest value) and standard deviation (lowest value) for the product layer diffusion control model.

The data was also used to determine the value of \( k_0 \) according to Eq. (7) at different values of temperature, similar to our previous work [31]. The rate constant, \( k_0 \) can be expressed as follows:

\[ k_0 = k_0 e^{\frac{E_a}{RT}} \]  \hspace{1cm} (7)

The linear form of Eq. (7) can be expressed as shown in Eq. (8):

\[ \ln(k_0) = \ln(k_0) - \frac{E_a}{RT} \]  \hspace{1cm} (8)

In Eq. (8), \( k_0 \) is the Arrhenius constant, \( E_a \) is the activation energy, \( T \) is the temperature and \( R \) is the universal gas constant [31]. The activation energy was calculated using the plots of rate constants at different temperatures using the linear form of Arrhenius equation. The summary of the results are given in Table 1. The activation energy in both cases is less than 20 kJ mol\(^{-1}\) (confirming the diffusion-controlled mechanism). Interestingly, the activation energy required to recover uranium from \( \text{SiO}_2\text{M} \) are almost comparable to \( \text{SiO}_2\text{M} \).

| S no. | Substrate | Activation Energy (kJ/mol) | Control mechanism |
|-------|-----------|---------------------------|-------------------|
| 1     | \( \text{SiO}_2\text{M} \) | 17.8                      | Product layer diffusion control: \( 1 + 2(1-x) - 3(1-x)^{2/3} = -17842 e^{-\frac{k}{RT}} \) |
| 2     | \( \text{SiO}_2\text{M} \) | 19.6                      | Product layer diffusion control: \( 1 + 2(1-x) - 3(1-x)^{2/3} = -10956 e^{-\frac{k}{RT}} \) |

5. Conclusions

The present work established the intensified recovery of uranium from the substrates. Based on the demonstrated results, it was observed that the process could be utilized widely in nuclear industries to recover the radioactive material uranium by intensified dissolution using ultrasound from functionalized and non-functionalized microparticles. It was observed that in 45 min, the uranium recovery was 34% for the \( \text{SiO}_2\text{M} \) and 39% for the \( \text{SiO}_2\text{M} \). The intensification due to ultrasound was more pronounced (five times) in case of \( \text{SiO}_2\text{M} \), than that (two times) in case of \( \text{SiO}_2\text{M} \) compared to the silent approach. The uranium metal recovery from the substrates was determined and the dissolution of uranium from biohybrid showed slightly higher activation energy than the silica microparticles.

CRediT authorship contribution statement

S. Lahiri: Conceptualization, Methodology, Investigation, Writing - original draft. A. Mishra: Methodology, Resources, Investigation, Writing - original draft. D. Mandal: Supervision. R.L. Bhardwaj: Supervision, Writing - review & editing, Project administration. P.R. Gogate: Supervision, Writing - review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Data availability

The raw/processed data required to reproduce these findings cannot be shared at this time as the data also forms part of an ongoing study.

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