



Cite this: *RSC Adv.*, 2017, 7, 18883

Lithium adsorption performance of a three-dimensional porous H_2TiO_3 -type lithium ion-sieve in strong alkaline Bayer liquor†

Xin Xu,^{abc} You Zhou,^{ab} Maohong Fan,^c Zijian Lv,^a Yang Tang,^{ab} Yanzhi Sun,^{ab} Yongmei Chen^{*ab} and Pingyu Wan^{ab}

Removing lithium from the Bayer liquor for ensuring good alumina product quality demands a special lithium ion-sieve (LIS) with good stability in a strong alkaline medium. In this study, a three-dimensional porous H_2TiO_3 -type LIS (porous-HTO) prepared by a polystyrene (PS) colloidal microspheres template was applied to adsorb Li^+ from the strong alkaline Bayer liquor. XRD and SEM results confirm the fine stability of porous-HTO in strong alkaline medium, and the regeneration tests show that more than 64 mg g^{-1} lithium adsorption capacity still remains even after 5 cycles of lithiation–delithiation in the simulation Bayer liquor. The lithium adsorption processes of porous-HTO and bare H_2TiO_3 -type LIS (bare-HTO) both fit the pseudo-second-order model, but the adsorption capacity and the adsorption rate of the porous-HTO are much better than those of the bare-HTO. For porous-HTO, the adsorption rate constant is $0.02357 \text{ g mg}^{-1} \text{ h}^{-1}$ and the equilibrium adsorption capacity is 76.3 mg g^{-1} , while for bare-HTO, the adsorption rate constant is $0.009682 \text{ g mg}^{-1} \text{ h}^{-1}$ and the equilibrium adsorption capacity is only 44.8 mg g^{-1} . The lithium selectivity tests demonstrate that the coexisting ions including Na^+ , K^+ , AlO_2^- , SiO_3^{2-} in the simulation Bayer liquor have low influence on lithium adsorption. The simulation Bayer liquors with Li^+ ions of 56.00, 30.00 and 5.00 mg L^{-1} are all reduced to below 1.00 mg L^{-1} by virtue of one-time-adsorption of porous-HTO at various solid to liquid ratios of 1.0, 0.5 and 0.1 g L^{-1} , respectively.

Received 24th January 2017

Accepted 20th March 2017

DOI: 10.1039/c7ra01056g

rsc.li/rsc-advances

Introduction

The Bayer process for producing refined smelter grade alumina has been applied for more than 100 years in industry.^{1–6} In the Bayer leaching process, the Bayer liquor containing a high concentration caustic sodium solution (*ca.* 200 g L^{-1}) is circularly used for alumina leaching. With cycle number increases, some impurities (mainly oxides and hydroxides of Si, Ga, and V, *etc.*) brought from bauxite would accumulate in the Bayer liquor, which might reduce the purity of the alumina product in

industry. In the past decades, researchers have used several methods such as fractional precipitation method, electrochemical method, solvent extraction method, and ion exchange method, *etc.* to remove Si, Ga, and V from the Bayer liquor.^{7–13} Recently, it has been reported that a kind of lithium-containing bauxite used for alumina production suffers from lithium enrichment ($50\text{--}100 \text{ mg L}^{-1}$) in the Bayer liquor, which has seldom been encountered in previous studies.^{14,15} Due to high dissolubility of most lithium salts, it is hard to remove lithium impurities by precipitation/co-precipitation methods from the Bayer liquor. In addition, considering the economic applicability in alumina industry, some reported conventional methods for lithium recovery from salt-lake brines or seawater such as solar evaporation method, solvent extraction method and hydrometallurgical method are all unsuitable.^{16–19}

In the past few years, some materials were found to possess selective lithium adsorption performance by virtue of special Li^+ screen crystal structures (either spinel structure or layered structure); correspondingly, these materials were called lithium ions sieves (LISs). The LISs have been proven to be efficient for recovering lithium ions from aqueous solution.^{20–25} However, most reported studies about LISs focused on neutral and weak alkaline medium although some of them involved the effect of alkalinity on lithium adsorption. Wang *et al.*²⁶ confirmed that

^aNational Fundamental Research Laboratory of New Hazardous Chemicals Assessment & Accident Analysis, Beijing University of Chemical Technology, Chaoyang District, No. 15, North 3rd Ring Road East, 100029 Beijing, P. R. China. E-mail: chenym@mail.buct.edu.cn; Tel: +86-13683294397

^bInstitute of Applied Electrochemistry, Beijing University of Chemical Technology, 100029 Beijing, P. R. China

^cDepartment of Chemical and Petroleum Engineering, University of Wyoming, Laramie, WY 82071, USA

† Electronic supplementary information (ESI) available: Details regarding XPS image of full spectra for porous-LTO, porous-HTO, and Li^+ -loaded porous-HTO, SEM images of 3D order self-assembly colloidal PS microspheres, kinetic fitting plots, lithium adsorption performance of Mn-LIS in strong alkaline solution, and lithium and sodium adsorption comparison by both porous-HTO and bare-HTO in the simulation Bayer liquor. See DOI: 10.1039/c7ra01056g



manganese-type LISs (Mn-LISs) were unstable in alkaline solution: the lithium adsorption capacity increased as alkalinity grew in the range of pH 7–12 but more than 1% of Mn dissolved when pH reached 13. The instability of Mn-LISs in strong alkaline medium might be caused by the serious Jahn–Teller distortion of Mn^{3+} during the Li^+ insertion process in spinels.^{27,28} In this regard, titanium-type LISs (Ti-LISs) display better stability in strong alkaline medium because no Jahn–Teller distortion occurs and the Ti–O bond energy is higher than the Mn–O bond energy.²⁵ Till now, although a few of studies have reported the lithium adsorption activities of Ti-LISs in weak alkaline solution (e.g. 0.1 mol L⁻¹ LiOH, alkali-added seawater, etc.),^{29,30} the lithium adsorption performance in strong alkaline solution has not been explored.

The objective of the present study is trying to remove lithium ions from the Bayer liquor by the layered H_2TiO_3 -type LIS (HTO), which is a typical Ti-LIS. In order to improve the lithium recovery performance in strong alkaline Bayer liquor, a three-dimensional (3D) porous H_2TiO_3 -type LIS (porous-HTO) was prepared by a polystyrene (PS) colloidal microspheres template. The lithium adsorption kinetics, regeneration performance and lithium selectivity of the porous-HTO were investigated by comparing those of the bare H_2TiO_3 -type LIS (bare-HTO). In addition, some suitable solid to liquid ratios of both porous-HTO and bare-HTO were suggested for lithium removal from the simulation Bayer liquors with different lithium concentrations in industry. This study offers a feasible way for lithium removal from the Bayer liquor and widens the practical application area of Ti-LIS.

Experimental section

LISs preparation

Bare-HTO. The precursor of the bare-HTO was obtained by a sol–gel method. The synthesis process started from dissolving $\text{Ti}[\text{O}(\text{CH}_2)_3\text{CH}_3]_4$ and LiNO_3 (mole ratio 1 : 2) in 150 mL ethyl alcohol, followed by adding 25 mL CH_3COOH to the above solution dropwise under constant stirring. After stirring for 30 min, a mixed solution containing 10 mL deionized water and 30 mL ethyl alcohol was dropped into this blend to turn the transparent solution into white sol. Then the sol was heated at 60 °C for 24 h in water bath environment to form light yellow gel. The light yellow dry gel powder was obtained by heating the gel at 120 °C for 12 h. In the crystallization process, the dry gel powder was heated to a final temperature of 800 °C at a rate of 2 °C min⁻¹ and kept at 800 °C for 4 h in the air to form the bare precursor (bare-LTO). After cooling to the ambient temperature, the bare-LTO was treated by 0.2 mol L⁻¹ HCl solution with occasional shaking for 4 h at 60 °C (1 g of solid in 1 L solution). Finally, the white bare-HTO powder was obtained by filtrating and drying.

Porous-HTO. The PS microspheres were synthesized by an emulsion polymerization method.^{31,32} First, 0.5 g cetane trimethyl ammonium bromide (CTAB) was dissolved in 150 mL deionized water. Second, 0.1 g ammonium persulfate (APS) as the initiator and 20 g styrene as the monomer were added to the above solution. Third, N_2 was piped into the reactor for 0.5 h to

remove the residual O_2 . After stirring for 24 h at 65 °C, a white stable emulsion was formed. Finally, the emulsion was placed in a vacuum freezing drying oven and kept for 120 h, thus obtaining white PS microspheres powders.

The porous-HTO was prepared by using PS microspheres as template. The precursor solution was obtained by adding solution A (6.81 g LiNO_3 dissolved in 100 mL ethanol and 0.6 mL H_2O) to solution B (3.40 g $\text{Ti}[\text{O}(\text{CH}_2)_3\text{CH}_3]_4$, 50 mL ethanol and 1.80 g acetic acid) dropwise under vigorous stirring for 1 h. Then the PS microspheres powders were soaked in the above solution for 0.5 h, followed by centrifugation and drying under vacuum at 40 °C for 2 h. The above operations were repeated for 3 times until the precursor solution penetrated the voids of PS templates completely. The obtained powder was calcined in a muffle to remove the PS template with a heating rate of 2 °C min⁻¹ and kept at 300 °C for 2 h, then heated up to 800 °C at the same rate and kept for 4 h. After that the precursor of porous-HTO (porous-LTO) was obtained. Finally, the porous-LTO was immersed in 0.2 mol L⁻¹ HCl for 4 h at 60 °C, then the porous-HTO was obtained after being filtered, washed and dried.

Characterizations

Structures and morphologies. The X-ray diffract (XRD) pattern of the samples was performed using Rigaku Smartlab X-ray diffractometer system equipped with graphite-monochromatized $\text{Cu}/\text{K}\alpha$ radiation ($\lambda = 1.5406 \text{ \AA}$). The scanning rate was 5° min⁻¹ in the 2θ range of 10–90°, operating at 40 kV and 40 mA. The X-ray photoelectron spectroscopy (XPS) spectra was measured with a Physical Electronics ESCA5800 spectrometer equipped with a monochromatic Al $\text{K}\alpha$ X-ray source ($E = 1486.6 \text{ eV}$). The scanning step and working pressure were 0.1 eV and $2 \times 10^{-7} \text{ Pa}$, respectively. The morphologies of the materials were studied by scanning electron microscopy (SEM, HITACHI, TM3000) using an accelerating voltage of 5 kV with a vacuum of $1 \times 10^{-6} \text{ Pa}$ and a resolution of 30 nm.

Lithium recovery process. The regeneration performance of bare-HTO and porous-HTO was tested in the procedure as follows: 0.1 g bare-HTO or porous-HTO was immersed in 1.0 L Li^+ -containing strong alkaline solution (6.0 mol L⁻¹ NaOH containing 56.00 mg L⁻¹ Li^+) and kept under magnetic stirring for 24 h at room temperature. The suspensions were filtered and the Li^+ concentration in the filter liquor was determined by an Inductively Coupled Plasma-Atomic Emission Spectrometry (ICP-AES, Shimadzu ICPS-75000). The residual solids were washed with deionized water for further delithiation. In delithiation process, the residues were treated with 0.2 mol L⁻¹ HCl solution at 60 °C and kept under magnetic stirring for 4 h. The lithiation–delithiation operations were repeated for 5 cycles. The titanium contents in bare-HTO and porous-HTO were determined by the methylene blue–ascorbic acid colorimetric method after it was dissolved in 40 mL nitric acid and 20 mL hydrochloric acid in a heating condition.³³

The adsorption kinetics and the adsorption selectivity were tested in the simulation Bayer liquor, which is a 6.0 mol L⁻¹ NaOH solution containing 66.81 mg L⁻¹ Li^+ , 56.79 mg L⁻¹ K^+ ,



70.46 mg L⁻¹ AlO₂⁻, and 56.79 mg L⁻¹ SiO₃²⁻. For the adsorption kinetics, Li⁺ ions concentration was determined at a certain time ($t = 0, 1, 2, 4, 6, 8, 12, 18, 24$ h). The adsorption selectivity behavior of LISs was determined by analyzing the concentration change of all ions in the simulation Bayer liquor. The distribution coefficient (K_d), separation factor (α_{Me}^{Li}) and concentration factor (CF) were calculated by the eqn (1)–(3):

$$K_d = \frac{C_0(\text{Me}) - C_e(\text{Me})}{C_e(\text{Me})} \times \frac{V}{W} \quad (1)$$

$$\alpha_{Me}^{Li} = \frac{K_d(\text{Li})}{K_d(\text{Me})} \quad (2)$$

$$\text{CF} = \frac{Q_e(\text{Me})}{C_0(\text{Me})} \quad (3)$$

where $C_0(\text{Me})$ is the initial concentration of each metal ion, $C_e(\text{Me})$ is the equilibrium concentration of each metal ion, V is the volume of the simulation solution, W is the mass of LIS and $Q_e(\text{Me})$ is the equilibrium adsorption capability of each metal ions by LISs.

Results and discussion

Lithium extraction from porous-LTO

The lithium extraction from porous-LTO experiments were designed for choosing the optimal acid pickling condition for preparing the porous lithium ion-sieve. The results are shown in Fig. 1. Fig. 1a presents the lithium extraction and titanium dissolution upon treatment of porous-LTO with different HCl solution. When the hydrochloric acid concentrations are lower than 0.2 mol L⁻¹, the extracted lithium amounts increase with the hydrochloric acid concentration and the titanium dissolution amounts keep at a low level (less than 0.1 mg g⁻¹). In this concentration range, the activity of H⁺ in solution plays a vital role for the chemical potentials equilibrium of lithium in both solid and solution, more Li⁺ can be replaced by the H⁺ in high H⁺ activity solution upon Li⁺-H⁺ ion exchange reaction. Meanwhile, low titanium dissolution indicates that the material is stable enough in this concentration range. When hydrochloric acid concentrations are not lower than 0.2 mol L⁻¹, the extracted lithium amounts keep at maximum values of approximately 127.3 mg g⁻¹ (the theoretical value of Li⁺ content in the Li₂TiO₃) and the titanium dissolution amounts increase significantly. Nearly 100% extracted Li⁺ amounts demonstrate that the Li⁺-H⁺ ion exchange reaction is completed in this range, and the increasing dissolved titanium dissolution amounts imply that the layered structure is not stable in high concentration hydrochloric acid solution. Therefore, the optimal concentration of hydrochloric acid solution for treating porous-LTO should be 0.2 mol L⁻¹. On the other hand, the bare Li₂TiO₃ (bare-LTO) treated in 0.2 mol L⁻¹ hydrochloric acid solution at 60 °C for 24 h also exhibits 99.9% extracted Li⁺ amount and only 0.82% dissolved Ti⁴⁺ amount, indicating 0.2 mol L⁻¹ hydrochloric acid solution is also fitted for the pickling process of bare-LTO. Fig. 1b shows the lithium extraction rates from porous-LTO in 0.2 mol L⁻¹ hydrochloric

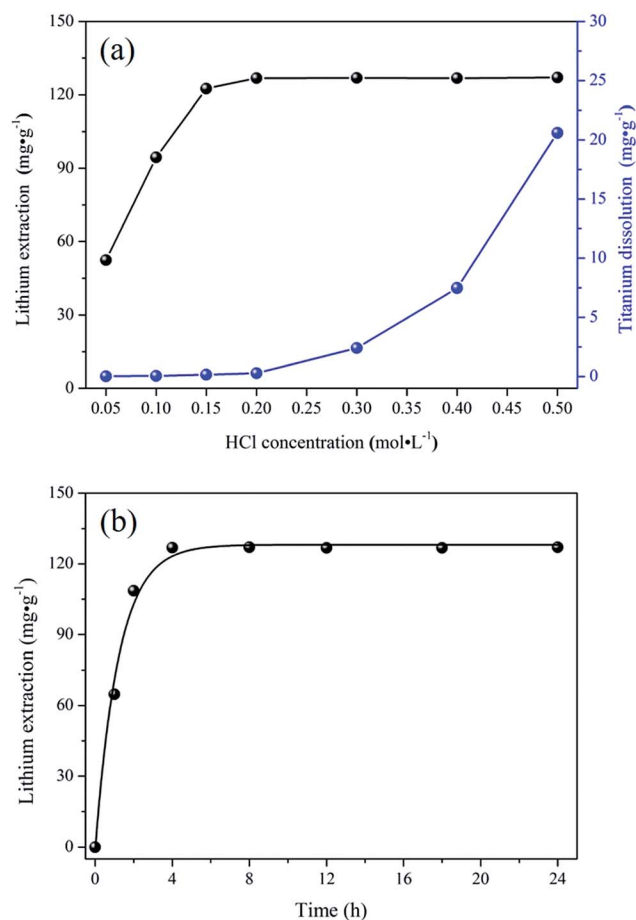


Fig. 1 (a) Lithium extraction and dissolution of titanium upon treatment of porous-LTO with different HCl solutions; (b) the lithium extraction rate from porous-LTO with 0.2 mol L⁻¹ HCl solution.

acid concentration. Obviously, the lithium extraction amount increases with time in the initial 4 h and keeps in maximum value (ca. 127.3 mg g⁻¹) in the following time, indicating pickling for 4 h in 0.2 mol L⁻¹ hydrochloric acid solution at 60 °C is enough to ensure a complete Li⁺ extraction from the porous-LTO. Also, this condition is fitted for completely converting the bare-LTO to the bare-HTO. In summary, the optimal acid pickling condition should be treating the porous-LTO in 0.2 mol L⁻¹ of hydrochloric acid solution for 4 h at 60 °C.

Structure characterization

Crystal structures. The XRD patterns of both bare and porous materials are shown in Fig. 2. For the bare-LTO (curve a), the diffraction peaks at $2\theta = 18.469^\circ, 20.165^\circ, 35.861^\circ, 43.582^\circ, 47.754^\circ, 57.565^\circ, 63.486^\circ, 66.735^\circ$ are assigned to the monoclinic Li₂TiO₃ (*C2/c* space group, JCPDS card 33-0831).³⁴ Similarly, the XRD pattern of porous-LTO (curve b) could also be indexed as the monoclinic Li₂TiO₃, indicating the existence of PS templates during the preparation does not affect the crystallization of Li₂TiO₃. After acid treatment, the XRD patterns of both bare-HTO (curve c) and porous-HTO (curve d) change dramatically in comparison to their precursors: the peak of



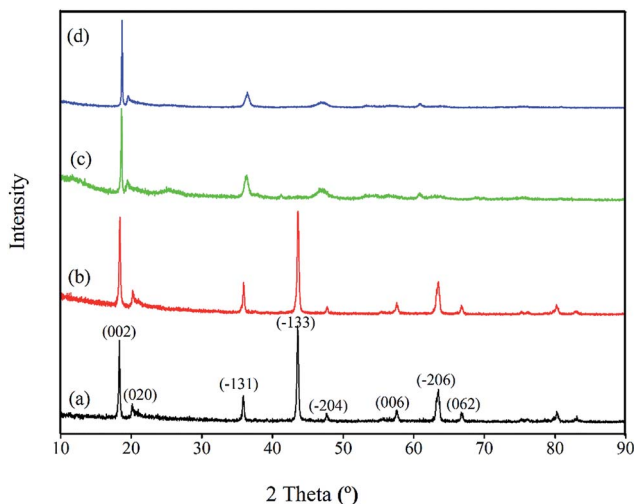


Fig. 2 XRD patterns of (a) the bare-LTO, (b) the porous-LTO, (c) the bare-HTO, and (d) the porous-HTO.

(020) shifts to left but the peaks of (002) and (-131) shift to right slightly, the peaks of (-131) and (-133) turn wide, and the peaks of (-133), (-206) and (062) all disappear. This change is well consistent with previous work,^{20,29,30} which is possibly ascribed to different bonding ways between Li and H in Ti-O layers. Meanwhile, curve c and d are quite similar, implying the first pickling process on porous-LTO is similar to that on bare-LTO, the PS-modified precursor also shows the same Li-H exchange process in Ti-O layered.

The further XPS examination (shown in Fig. 3) verifies that the patterns of HTO before and after loading lithium ions from strong alkaline solution ($6.0 \text{ mol L}^{-1} \text{ NaOH}$ containing $56.00 \text{ mg L}^{-1} \text{ Li}^+$) are different (the full spectra see Fig. S1 in ESI[†]). Through precision survey, the Ti_{2p} , O_{1s} , and Li_{1s} peaks are shown in Fig. 3a, b and c, respectively. In Fig. 3a, the binding energy values of $\text{Ti } 2p_{1/2}$ and $\text{Ti } 2p_{3/2}$ peaks for porous-LTO are 463.79 eV and 458.27 eV, respectively, increasing slightly for both porous-HTO (464.66 and 458.91 eV) and Li^+ -loaded porous-HTO (464.74 and 458.86 eV). The binding energy values of $\text{Ti } 2p_{1/2}$ and $\text{Ti } 2p_{3/2}$ peaks for porous-HTO and Li^+ -loaded porous-HTO are very close.^{35–37} In Fig. 3b, the binding energy value of O_{1s} peak for porous-LTO is 529.53 eV, increasing slightly for both porous-HTO (529.78 eV) and Li^+ -loaded porous-HTO (529.68 eV) likewise. The binding energy values of O_{1s} peak value for porous-HTO and Li^+ -loaded porous-HTO are also quite alike. The similar binding energy variation trend of Ti_{2p} and O_{1s} demonstrates the chemical structure of porous-HTO are different from that of porous-LTO, but similar with that of Li^+ -loaded porous-LTO; in other words, the chemical environments of Ti and O have irreversibly changed after first acid treatment. The reason could be attributed to different impacts of Li and H on Ti-O layers: owing to different radius of Li^+ (*ca.* 0.76 Å) and H^+ (*ca.* 0.012 Å), the space occupancy of H^+ is much smaller than that of Li^+ , and the formed HTO layers is more flexible than the LTO. Thus, the interaction between Ti and O weakens and the Ti-O bond lengthens after first acid treatment, resulting further structural transformation. In the later Li adsorption

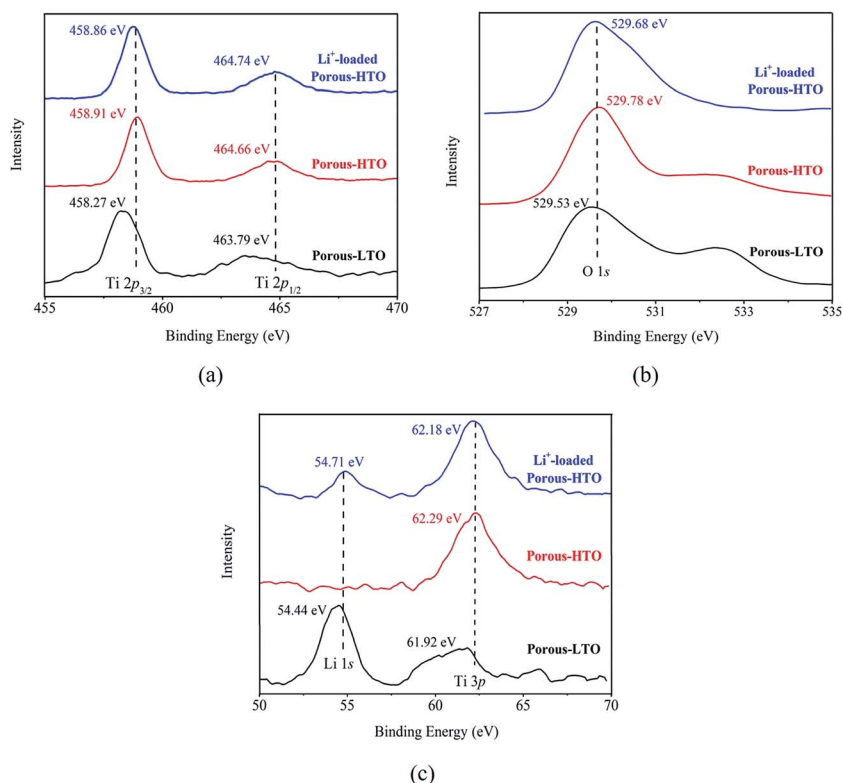


Fig. 3 XPS images of (a) Ti_{2p} spectra, (b) O_{1s} spectra and (c) Li_{1s} spectra.



process, Li^+ ions are probably accommodated into the HTO with the electrostatic force.²⁹ In Fig. 3c, the Li_{1s} exhibits the highest relative intensity (*vs.* Ti_{3p}) on porous-LTO at binding energy of 54.44 eV, while it almost disappears on porous-HTO and reappears on Li^+ -loaded porous-HTO at binding energy of 54.71 eV. However, the relative intensity on Li^+ -loaded porous-HTO is lower than that on porous-LTO, suggesting the lithium content in Li^+ -loaded sample is lower than that in initial precursor sample. The structural change of porous materials is consistent with previous work for bare materials, indicating the PS modification has no impact on Li adsorption mechanism.³⁸

Surface morphology. The SEM images of the materials are shown in Fig. 4. The bare-LTO displays a smooth surface (Fig. 4a), while the porous-LTO exhibits regular honeycomb porous structure (Fig. 4b), which is attributed to the 3D ordered self-assembly colloidal PS microspheres (for details see Fig. S2 in ESI†). Fig. 4c presents the surface morphology of bare-HTO. The results suggest the alterations of the solid particle surface after delithiation are not remarkably visible. In Fig. 4d, the honeycomb-like structure of the porous-HTO is also retained after Li^+ ions are exchanged with protons, suggesting the impact of both water molecules and Li-H ions exchange reaction on the change in porous structure is limited.

Specific surface areas. BET results in Table 1 demonstrate the specific surface area change of bare and porous samples. It is obvious that the specific surface area of porous samples is 8–10 times as large as bare samples. The results turn out that PS modification shows remarkable improvement in surface area of the materials. Besides, the surface areas of both bare-HTO and porous-HTO are larger than that of the corresponding LTO, which may be attributed to the preferential adsorption on acid adsorption sites.²⁹

Table 1 BET surface areas of LTO and HTO

BET surface areas ($\text{m}^2 \text{g}^{-1}$)		
Bare samples	Bare-LTO	5.92
	Bare-HTO	12.87
Porous samples	Porous-LTO	67.61
	Porous-HTO	94.14

Adsorption kinetics

The adsorption kinetics of lithium ions on both bare-HTO and porous-HTO in the simulation Bayer liquor ($6.0 \text{ mol L}^{-1} \text{ NaOH}$, $56.00 \text{ mg L}^{-1} \text{ Li}^+$) were investigated to determine the lithium adsorption rate and establish the equilibrium lithium adsorption capacity. As shown in Fig. 5, the curves of Li^+ uptake by the two adsorbents rise fast in the initial period followed by becoming flat afterwards. However, the Li^+ adsorption equilibrium time and the capacity of the two adsorbents are different: about 4 h are needed for the porous-HTO to reach equilibrium, which is much shorter than that of bare-HTO (*ca.* 8 h). The practical equilibrium lithium capacity on porous-HTO is about 70 mg g^{-1} , which is much higher than that of bare-HTO (*ca.* 40 mg g^{-1}).

Two kinds of kinetics models of pseudo-first-order kinetic model (eqn (4)) and pseudo-second-order kinetic model (eqn (5))^{39,40} were applied to investigate the lithium adsorption kinetic behaviors:

$$\lg(q_e - q_t) = -\frac{k_1 t}{2.303} + \lg q_e \quad (4)$$

$$\frac{t}{q_t} = \frac{t}{q_e} + \frac{1}{k_2 q_e^2} \quad (5)$$

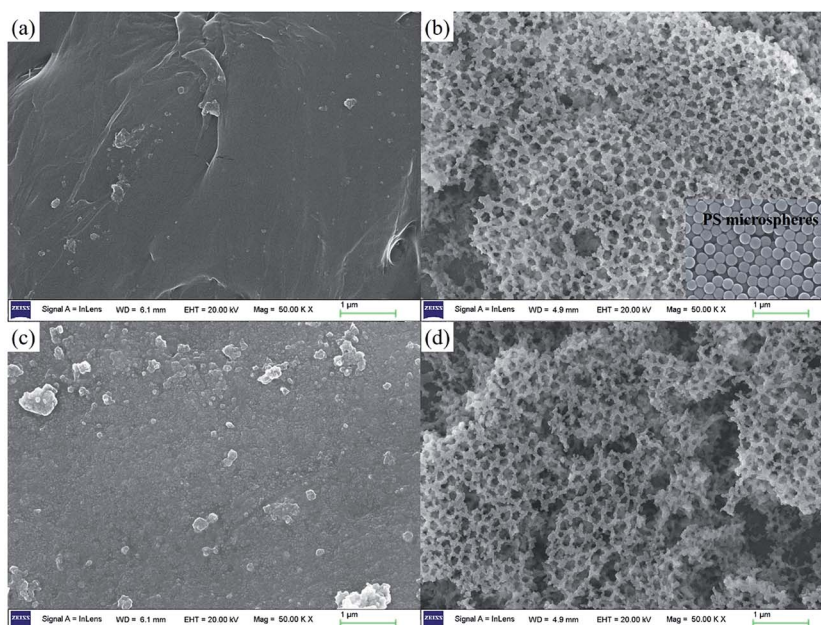


Fig. 4 SEM images of (a) bare-LTO, (b) porous-LTO, (c) bare-HTO, and (d) porous-HTO.



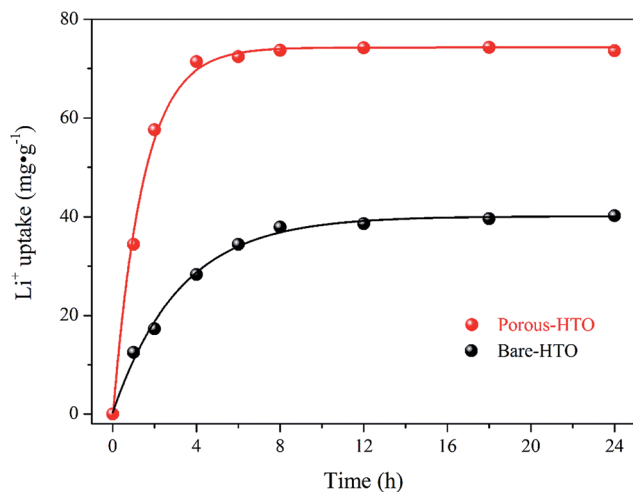


Fig. 5 Li^+ uptake kinetics of both bare-HTO and porous-HTO in the simulation Bayer liquor.

In above equations, q_e and q_t are the equilibrium lithium capacity and the lithium adsorption capacity in the time of t ; k_1 and k_2 represent the adsorption rate constants of pseudo-first-order kinetic model and pseudo-second-order kinetic model respectively. All of the kinetic parameters were calculated according to the slope and the intercept of the fitted-lines. The fitting results are shown in Table 2 (the fitting curves are illustrated in Fig. S3 in ESI†).

As shown in Table 2, it is more likely to describe Li^+ adsorption on porous-HTO in the simulation Bayer liquor as a pseudo-second-order kinetic process because the linear correlation coefficient of the pseudo-second-order dynamic equation (0.9980) is better than that of pseudo-first-order dynamic equation (0.8846). Similarly, the process of Li^+ adsorption on bare-HTO could also be described as a pseudo-second-order dynamic process. The results suggest the rate-determining factors in lithium adsorption on HTO are related with not only the lithium concentration in liquid phase but also the quantity of active lithium adsorption sites in the solid phase. In other words, under the premise of constant lithium concentration in solution, large surface area of the adsorbent is beneficial for improving the lithium adsorption rate in the simulation Bayer liquor because larger surface area has more active Li^+ adsorption sites. Just as the results indicate, the rate constants for the bare-HTO and porous-HTO are 0.009682 and 0.02357 $\text{g mg}^{-1} \text{h}^{-1}$, respectively.

Table 2 Kinetics parameters for Li^+ adsorption on both bare-HTO and porous-HTO in the simulation Bayer liquor at room temperature

Fitting models	Parameters	Bare-HTO	Porous-HTO
Pseudo-first-order	k_1 (h^{-1})	0.2411	0.3749
	q_e (mg g^{-1})	40.2	74.4
	R^2	0.9434	0.8846
Pseudo-second-order	k_2 ($\text{g mg}^{-1} \text{h}^{-1}$)	0.009682	0.02357
	q_e (mg g^{-1})	44.8	76.3
	R^2	0.9946	0.9980

Moreover, the porous-HTO also presents higher practical equilibrium lithium adsorption capacity than the bare one (76.3 mg g^{-1} vs. 44.8 mg g^{-1}). This may be attributed to the superior permeability of porous-HTO because porous-HTO provides a short diffusion pathway for lithium ions so that the $\text{Li}^+/\text{H}_2\text{TiO}_3$ contact is improved effectively. Consequently, the inner solid particles of the porous-HTO play an effective role in lithium ions adsorption, and the apparent lithium adsorption amount could be enhanced as compared with bare-HTO. In summary, by virtue of the PS modification on adsorbent, the mass transfer efficiency between solid adsorbent and lithium ions in simulation Bayer liquor is improved effectively, and the corresponding lithium recovery performance including both the lithium adsorption rate and the practical lithium adsorption capacity is enhanced remarkably.

Regeneration performance

Fig. 6 shows lithium uptake change by both bare-HTO and porous-HTO during 5 cycles of lithiation–delithiation in the simulation Bayer liquor ($6.0 \text{ mol L}^{-1} \text{ NaOH}$, $56.00 \text{ mg L}^{-1} \text{ Li}^+$). It is observed that the lithium adsorption capacities of bare-HTO and porous-HTO both remains at a relatively stable level in 5 cycles' operations, implying the H_2TiO_3 -type lithium ion-sieves have good stability in strong alkaline medium. Notably, the lithium adsorption capacity fading of porous-HTO (about 10.81%) is slight higher than that of bare-HTO (about 4.65%), which is possibly caused by the destruction of pore structures in cyclic operations (Fig. 8a). On the other hand, the dissolution rate of Ti for both bare-HTO and porous-HTO in each cycle are less than 0.5 wt%, indicating the dissolution of both adsorbents are quite limited. However, as the comparison shows, the Li^+ uptake amount of $\lambda\text{-MnO}_2$ in $6.0 \text{ mol L}^{-1} \text{ NaOH}$ solution decreases sharply at the second cycle due to the dissolution of Mn (the detailed lithium recovery performance of $\lambda\text{-MnO}_2$ in strong alkaline medium are illustrated in Fig. S4 in ESI†). The results strongly

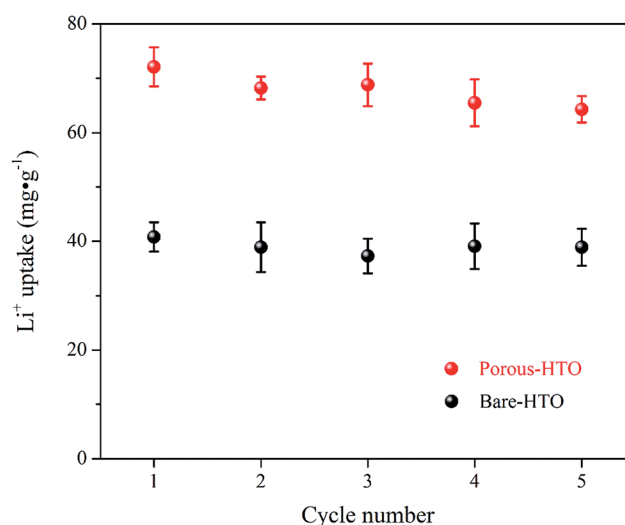


Fig. 6 Regeneration performances of both bare-HTO and porous-HTO in the simulation Bayer liquor.



demonstrate that Ti-LISs are more stable than Mn-LISs in strong alkaline medium, so it is appropriate to choose the Ti-LISs as the lithium adsorbent for recovering Li^+ from strong alkaline medium.

The XRD patterns of Li^+ -loaded bare-HTO and Li^+ -loaded porous-HTO after 5 cycle operation are shown in Fig. 7. All of the diffraction peaks derived from (002), (020), (-131), and (-204) crystal faces are the same as those of porous-HTO (shown in Fig. 2d), and no extra peaks appear. The results manifest both Li^+ -loaded bare-HTO and Li^+ -loaded porous-HTO keep fine crystallinity even if the operation is at the 5th cycle. On the other hand, in Fig. 8, the SEM images of porous-HTO after 5 times of Li^+ -loading still possess the porous structure which is almost similar to the as-prepared one shown in Fig. 4 except for occurrence of a slight collapse of the porous structure. Based on the above results, it is concluded that both two adsorbents present fine regeneration performance in the simulation Bayer liquor, although the porous-HTO shows slight higher Li^+ adsorption capacity fading degree than the bare-HTO, the practical Li^+ adsorption capacity of porous-HTO is higher than another one.

Adsorption selectivity

The results of adsorption selectivity of lithium ions to other coexisted ions in the simulation Bayer liquor on both porous-HTO and bare-HTO are listed in Table 3. For both adsorbents, the equilibrium distribution coefficient (K_d) for Li^+ and other ions are evaluated as $\text{Li}^+ \gg \text{K}^+ > \text{SiO}_3^{2-} \approx \text{AlO}_2^-$. Apparently, the Li^+ ions are preferentially adsorbed by both adsorbents on the whole. In addition, the separation factors ($\alpha_{\text{Me}}^{\text{Li}}$) of Li^+ to other coexisting ions in the simulation Bayer liquor are much higher than 1.00, and the Li^+ concentration factor (CF) value of both adsorbents are far above those of other ions (577.49 of bare-HTO and 908.40 of porous-HTO), indicating the existence of K^+ , AlO_2^- , and SiO_3^{2-} does not interfere with Li^+ during the adsorption process. The above results demonstrate that (1) the

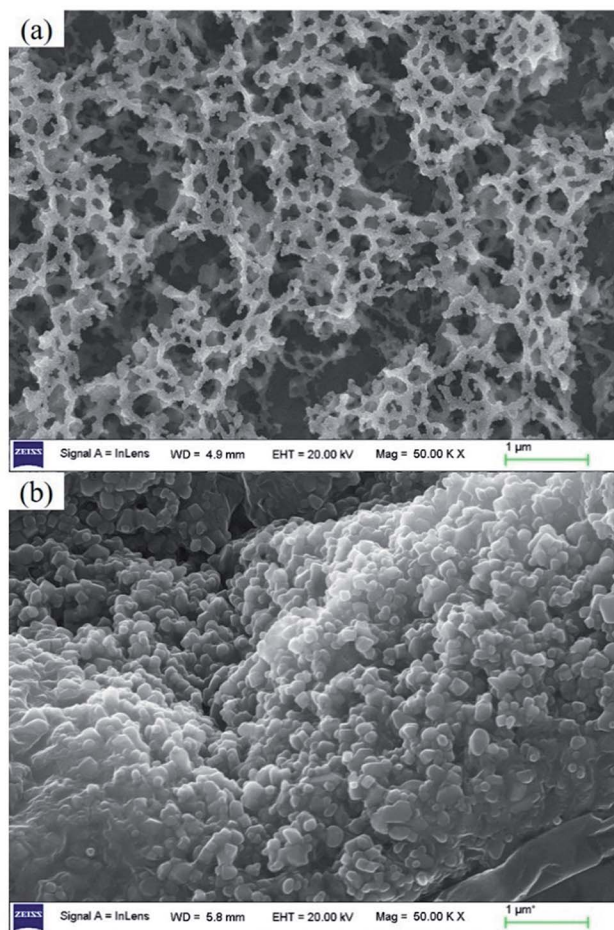


Fig. 8 SEM images of (a) Li^+ -loaded porous-HTO and (b) Li^+ -loaded bare-HTO after 5 cycle operation.

layered H_2TiO_3 -type ion-sieve has high selectivity to Li^+ but much low to all other ions, (2) the modification of PS microspheres almost has no effect on lithium selectivity of the adsorbent.

On the other hand, it is hard to determine the change in concentration of Na^+ ions in the simulation Bayer liquor because the concentration of Na^+ is far higher than that of Li^+ (mass ratio $\text{Na} : \text{Li} = 2066 : 1$). In order to investigate the influence of Na^+ ions on lithium adsorption, an additional experiment was performed: same quantity of sodium ions and lithium ions (*ca.* 60.0 mg L^{-1}) was dissolved in 1 L 6.0 mol L^{-1} KOH aqueous solution, then 1.00 g of adsorbent was added and kept for 24 h under stirring. The results show that the equilibrium adsorption capacity of Li^+ on the porous-HTO is 53.67 mg L^{-1} and that of Na^+ is only 0.65 mg L^{-1} . The $\alpha_{\text{Na}}^{\text{Li}}$ value is 770.99, and the CF value of Li and Na is 893.90 and 10.81 respectively, revealing that Na^+ does not interfere with Li^+ during the adsorption process. That is to say, the huge quantity of sodium ions existing in the simulation Bayer liquor presents low influence on lithium adsorption, and the modification of PS microspheres has no adverse impact on the lithium selectivity from the simulation Bayer liquor (the results were listed in Table S1 in ESI†).

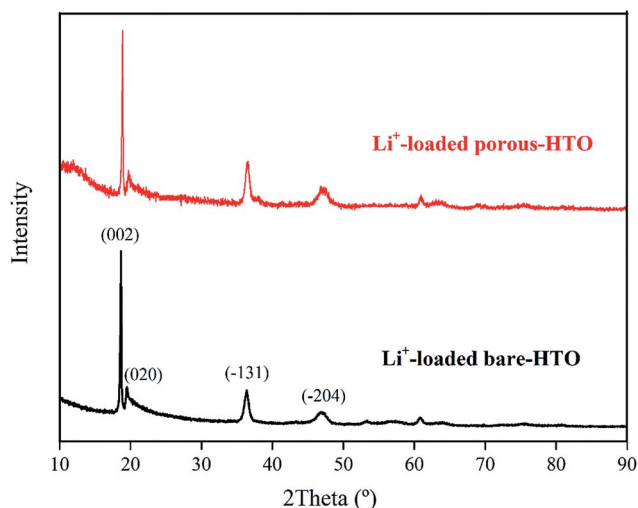


Fig. 7 XRD patterns of both Li^+ -loaded porous-HTO and Li^+ -loaded bare-HTO after 5 cycle operation.



Table 3 Lithium selectivity on bare-HTO and porous-HTO in the simulation Bayer liquor^a

Ions	Bare-HTO						Porous-HTO					
	C_0 (mg L ⁻¹)	C_e (mg L ⁻¹)	Q_e (mg g ⁻¹)	K_d (mL g ⁻¹)	α_{Me}^{Li}	CF (L mg ⁻¹)	C_0 (mg L ⁻¹)	C_e (mg L ⁻¹)	Q_e (mg g ⁻¹)	K_d (mL g ⁻¹)	α_{Me}^{Li}	CF (L mg ⁻¹)
Li ⁺	66.81	28.23	38.58	1366.63	1.00	577.49	66.81	6.12	60.69	9916.67	1.0	908.40
K ⁺	56.79	54.12	2.67	49.33	27.70	47.02	56.79	54.28	2.51	46.24	214.45	44.20
AlO ₂ ⁻	70.46	69.83	0.63	9.02	151.48	8.94	70.46	69.11	1.35	19.53	507.66	19.16
SiO ₃ ²⁻	84.57	82.25	2.32	28.21	48.45	27.43	84.57	84.01	0.56	6.67	1487.68	6.62

^a Notes: C_0 is the initial concentration of each metal ion; C_e is the equilibrium concentration of each metal ion; Q_e is the calculated equilibrium adsorption capability of each metal ions by LISs; K_d is the calculated equilibrium distribution coefficient and α_{Me}^{Li} is the calculated separation factors.

Dosage

Different amount of the bare-HTO and porous-HTO was added to the 1.0 L simulation Bayer liquors with different Li⁺ concentrations. In dosage tests, the criterion of residual Li⁺ concentration in simulation Bayer liquor is set as below 1.0 mg L⁻¹, which is acceptable in alumina industrial production. The results of lithium removal after one-time adsorption are listed in Table 4. For the initial Li⁺ concentration of 56.00 mg L⁻¹ (no. 1, 3 and 6), the remaining Li⁺ is decreased to below 1.0 mg L⁻¹ when using 1.0 g L⁻¹ porous-HTO of at least 1.0 g L⁻¹. In comparison, for achieving the same Li removal effect, at least 2.0 g L⁻¹ of bare-HTO are required. If the initial concentration of Li⁺ ions is lower, the required dosage of the LIS would be smaller. For instance, for the simulation Bayer liquors with Li⁺ ions of 40.00 mg L⁻¹ (no. 4 and 7) and 30.00 mg L⁻¹ (no. 5, 8 and 11), in order to reduce Li⁺ ions to below 1.00 mg L⁻¹, 1.0 and 0.5 g L⁻¹ porous-HTO are needed respectively, while 1.0 g L⁻¹ bare-HTO is needed for both liquors. Notably, for the simulation Bayer liquor with 5.00 mg L⁻¹ of initial Li⁺ ions, although the residual Li⁺ concentration by using larger doses (no. 10 and no. 14) is slight higher than that by using 0.1 g porous-HTO (no. 15), which is

caused by small measuring errors in practical operations, only 0.1 g L⁻¹ porous-HTO is enough to ensure the residual Li⁺ concentration decreases to below 1.0 mg L⁻¹ (no. 15).

Conclusions

The H₂TiO₃-type lithium ion-sieve was proven to have fine structural stability in strong alkaline medium, and the adsorption kinetic studies show Li⁺ ions adsorbed by HTO follows the pseudo-second-order kinetic model, suggesting that the adsorption kinetic is correlated with the quantity of active sites on LIS. As one of promising methods to remove lithium ions from the strong alkaline Bayer liquor, a 3D porous-HTO prepared by PS template method was prepared. Comparing with the bare-HTO, the equilibrium lithium adsorption capacity and the rate constant of the porous-HTO have been improved significantly. In addition, the adsorption selectivity, regeneration performance of both porous-HTO and bare-HTO were proven to be fine in simulation Bayer liquor. Under optimum conditions, 56.00 mg L⁻¹ of Li⁺ ions in Bayer liquor can be decreased to less than 1.0 mg L⁻¹ by adding 1.0 g L⁻¹ porous-HTO after just one time adsorption. In our future work, technology improvement of the recovery utilization rate of the powder materials and reduction of energy consumption will be studied.

Table 4 Lithium removal effects from simulation Bayer liquor by the porous-HTO and bare-HTO with different dosage

No.	Initial Li ⁺ concentration (mg L ⁻¹)	Dosage of LIS (g L ⁻¹)	Remaining Li ⁺ concentration (mg L ⁻¹)	
			Porous-HTO	Bare-HTO
1	56.00	2.0	0.04	0.06
2	60.00	1.0	1.17	30.04
3	56.00	1.0	0.98	26.45
4	40.00	1.0	0.02	0.57
5	30.00	1.0	0.04	0.09
6	56.00	0.5	32.58	34.88
7	40.00	0.5	12.43	23.63
8	30.00	0.5	0.82	11.47
9	20.00	0.5	0.01	0.62
10	5.00	0.5	0.05	0.02
11	30.00	0.2	19.90	22.61
12	20.00	0.2	7.80	11.06
13	10.00	0.2	0.04	1.95
14	5.00	0.2	0.01	0.01
15	5.00	0.1	0.04	1.28

Conflict of interest

The authors declare no competing financial interest.

Acknowledgements

The authors greatly appreciate the support from the National Natural Science Foundation of China (No. 51374016, 21506010), the Fundamental Research Funds for the Central Universities of China (No. YS1406, JD1515) and BUCT Fund for Disciplines Construction and Development (No. XK1525, XK1531).

References

- 1 G. Power, J. S. C. Loh, J. E. Wajon, F. Busetti and C. Joll, *Anal. Chim. Acta*, 2011, **689**, 8–21.



- 2 S. Ruan, L. Shi, J. Li and A. R. Gerson, *Hydrometallurgy*, 2016, **163**, 1–8.
- 3 Y. L. Sidrak, *Ind. Eng. Chem. Res.*, 2001, **40**, 1146–1156.
- 4 E. VanDalen and L. G. Ward, *Anal. Chem.*, 1974, **46**, 1826.
- 5 A. Buvári-barcza, M. Rozsahegyi and L. Barcza, *J. Mater. Chem.*, 1998, **8**, 451–455.
- 6 E. VanDalen and L. G. Ward, *Anal. Chem.*, 1973, **45**, 2248–2251.
- 7 X. Pan, H. Yu, G. Tu and S. Bi, *Hydrometallurgy*, 2016, **165**(2), 261–269.
- 8 L. Zeng and Z. Li, *Ind. Eng. Chem. Res.*, 2012, **51**, 15193–15206.
- 9 Z. Zhao, Y. Yang, Y. Xiao and Y. Fan, *Hydrometallurgy*, 2012, **125–126**, 115–124.
- 10 P. Selvi, M. Ramasami, M. H. P. Samuel, P. Adaikkalam and G. N. Srinivasan, *Ind. Eng. Chem. Res.*, 2004, **43**, 2216–2221.
- 11 B. Gupta, N. Mudhar and S. N. Tandon, *Ind. Eng. Chem. Res.*, 2005, **44**, 1922–1927.
- 12 Z. Zhao, H. Long, X. Li, Y. Fan and Z. Han, *Hydrometallurgy*, 2012, **115–116**, 52–56.
- 13 M. D. Okudan, A. Akcil, A. Tuncuk and H. Deveci, *Hydrometallurgy*, 2015, **152**, 76–83.
- 14 G. Kuang, H. Li, S. Hu, R. Jin, S. Liu and H. Guo, *Hydrometallurgy*, 2015, **157**, 214–218.
- 15 P. Meshram, B. D. Pandey and T. R. Mankhand, *Hydrometallurgy*, 2014, **150**, 192–208.
- 16 Z. Zhou, W. Qin, S. Liang, Y. Tan and W. Fei, *Ind. Eng. Chem. Res.*, 2012, **51**, 12926–12932.
- 17 Z. Xu, H. Zhang, R. Wang, W. Gui, G. Liu and Y. Yang, *Ind. Eng. Chem. Res.*, 2014, **53**, 16502–16507.
- 18 T. Tran and V. T. Luong, in *Lithium Process Chemistry*, ed. J. Świątowska, Elsevier, Amsterdam, 2015, pp. 81–124, DOI: 10.1016/B978-0-12-801417-2.00003-7.
- 19 J. W. An, D. J. Kang, K. T. Tran, M. J. Kim, T. Lim and T. Tran, *Hydrometallurgy*, 2012, **117–118**, 64–70.
- 20 R. Chitrakar, Y. Makita, K. Ooi and A. Sonoda, *Dalton Trans.*, 2014, **43**, 8933–8939.
- 21 R. Chitrakar, H. Kanoh, Y. Miyai and K. Ooi, *Chem. Mater.*, 2000, **12**, 3151–3157.
- 22 S.-Y. Sun, X. Song, Q.-H. Zhang, J. Wang and J.-G. Yu, *Adsorption*, 2011, **17**, 881–887.
- 23 Q. Feng, Y. Miyai, H. Kanoh and K. Ooi, *Chem. Mater.*, 1993, **5**, 311–316.
- 24 R. Chitrakar, Y. Makita, K. Ooi and A. Sonoda, *Ind. Eng. Chem. Res.*, 2014, **53**, 3682–3688.
- 25 X. Xu, Y. Chen, P. Wan, K. Gasem, K. Wang, T. He, H. Adidharma and M. Fan, *Prog. Mater. Sci.*, 2016, **84**, 276–313.
- 26 L. Wang, C. G. Meng and W. Ma, *Colloids Surf., A*, 2009, **334**, 34–39.
- 27 S. Ghose, M. Kersten, K. Langer, G. Rossi and L. Ungaretti, *Phys. Chem. Miner.*, 1986, **13**, 291–305.
- 28 A. Yamada and M. Tanaka, *Mater. Res. Bull.*, 1995, **30**, 715–721.
- 29 C. P. Lawagon, G. M. Nisola, J. Mun, A. Tron, R. E. C. Torrejos, J. G. Seo, H. Kim and W.-J. Chung, *J. Ind. Eng. Chem.*, 2016, **35**, 347–356.
- 30 X.-c. Shi, Z.-b. Zhang, D.-f. Zhou, L.-f. Zhang, B.-z. Chen and L.-l. Yu, *Trans. Nonferrous Met. Soc. China*, 2013, **23**, 253–259.
- 31 L. J. Fu, T. Zhang, Q. Cao, H. P. Zhang and Y. P. Wu, *Electrochem. Commun.*, 2007, **9**, 2140–2144.
- 32 Y. Cao, Y. Wang, Y. Zhu, H. Chen, Z. Li, J. Ding and Y. Chi, *Superlattices Microstruct.*, 2006, **40**, 155–160.
- 33 M. H. Zavvar and N. Pourreza, *J. Chin. Chem. Soc.*, 2008, **55**, 750–754.
- 34 Y.-z. Hao, Q.-l. Zhang, J. Zhang, C.-r. Xin and H. Yang, *J. Mater. Chem.*, 2012, **22**, 23885–23892.
- 35 G. He, J. Gao, H. Chen, J. Cui, Z. Sun and X. Chen, *ACS Appl. Mater. Interfaces*, 2014, **6**, 22013–22025.
- 36 G. He, J. Liu, H. Chen, Y. Liu, Z. Sun, X. Chen, M. Liu and L. Zhang, *J. Mater. Chem. C*, 2014, **2**, 5299–5308.
- 37 J. W. Zhang, G. He, L. Zhou, H. S. Chen, X. S. Chen, X. F. Chen, B. Deng, J. G. Lv and Z. Q. Sun, *J. Alloys Compd.*, 2014, **611**, 253–259.
- 38 G. He, L. Zhang, D. Zhou, Y. Zou and F. Wang, *Ionics*, 2015, **21**, 2219–2226.
- 39 T. Ryu, J. Shin, D.-H. Lee, J. Ryu, I. Park, H. Hong, Y. S. Huh, B.-G. Kim and K.-S. Chung, *Hydrometallurgy*, 2015, **157**, 39–43.
- 40 Y. S. Ho and G. McKay, *Water Res.*, 2000, **34**, 735–742.

