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## Molten ionic oxides for CO<sub>2</sub> capture at medium to high temperatures†

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The development of efficient low cost  $CO_2$  capture systems is a critical challenge for mitigating climate change while meeting global energy demand. Herein, we demonstrate the first liquid absorbents for  $CO_2$  capture at medium to high temperatures (500 to 700 °C). Molten ionic oxides based on sodium borate and the mixed alkali-metal borates show remarkably fast kinetics and intrinsic regenerability, with no observable deterioration in performance over multiple absorption—desorption cycles under both temperature—and pressure—swing operations. The behavior of the molten ionic oxides is ascribed to the instantaneous formation of carbonate ions in the molten oxides without the diffusional transport restrictions imposed by solid product layers characteristic of solid adsorbents. The new liquid absorbents will enable continuous processing and thermal integration *via* a simple absorber—desorber arrangement, thereby overcoming the challenges previously restraining high temperature  $CO_2$  capture and opening up new opportunities in clean energy production.

#### Introduction

Mitigation of the substantial risks associated with changes in global climate patterns due to the accumulation of anthropogenic CO<sub>2</sub> in the atmosphere requires the development of energy efficient carbon capture, utilization and storage (CCUS) technologies. Globally deployed CCUS technologies would enable the continued use of existing energy infrastructure in a carbon constrained world, stabilizing the energy landscape for the foreseeable future. The cost and operability of CO<sub>2</sub> capture systems are key to determining the future pathway of global technology pervasion. Innovative breakthroughs in sorbent materials and/or processes are expected to accelerate future developments in, and implementation of, advanced low-cost CO<sub>2</sub> capture methods.

The sorbents required for CO<sub>2</sub> capture can be classified into several groups in accordance with their operating temperature.<sup>5</sup> For low temperature operations, various types of aqueous amines (*e.g.*, MEA, DEA, *etc.*) have been considered as promising absorbents,<sup>6,7</sup> and utilized in most of the current or planned CO<sub>2</sub> capture systems at industrial facilities.<sup>8,9</sup> Amine scrubbing processes, however, face several technological drawbacks due to their high energy demands, degradation of the amines, and equipment corrosion.<sup>3,10,11</sup> As an alternative, various new methods operating at medium to high

temperatures (T > 200 °C) have recently attracted attention for both pre-combustion and post-combustion processes. In precombustion systems, CO2 must be separated from a mixture of H<sub>2</sub> and CO<sub>2</sub> produced by coal gasification or steam methane reforming with water gas shift reactions, and the removal of CO<sub>2</sub> at the temperature of the reforming or shift reactions in the range of 200 to 600 °C leads to lower energy costs for CO2 capture and an enhancement in H2 production yield.12-14 In post-combustion systems, CO2 capture from the exhaust at high temperature, in particular for next generation NGCC power plants, has advantages in both the uptake efficiency and energy economics when compared to the conventional low temperature systems.15,16 Various types of solid adsorbents, such as calcium oxide (CaO),17-19 magnesium oxide (MgO),20-22 lithiumbased mixed oxides (Li<sub>2</sub>ZrO<sub>3</sub>,<sup>23-25</sup> Li<sub>4</sub>SiO<sub>4</sub>,<sup>26-28</sup> LiBO<sub>2</sub>,<sup>29</sup> Li<sub>3</sub>BO<sub>3</sub>,<sup>30</sup> etc.), and layered double hydroxides (LDHs),31,32 have been proposed and studied extensively for CO<sub>2</sub> capture at medium to high temperatures. These solid adsorbents show superior uptake performance with high capacities, thermal stability, and lower environmental impact.5,33 The CO2 capture processes utilizing solid adsorbents, however, face drawbacks with slowuptake kinetics, low cyclic durability and complex system design due to the intrinsic physical behavior of the adsorbents as solid particles.34-36 It is recognized that the development of new advanced sorbents operating via energy efficient processes is necessary for the success of CCUS.

Herein, we identify molten ionic oxides that exist in the liquid phase at temperatures above their melting points as a new class of liquid absorbents operating at medium to high temperatures. Among the many molten ionic oxides available,<sup>37-39</sup> we have focused on sodium and mixed alkali-metal

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borates. These new liquid absorbents show rapid  $\mathrm{CO}_2$  uptake kinetics over the temperature range 500 to 700 °C with excellent cyclic regenerability. The exceptional  $\mathrm{CO}_2$  uptake performance by the molten alkali-metal borates is discussed in detail below, as are the advantages of the potential exploitation of these molten salt absorbents in systems for  $\mathrm{CO}_2$  capture relative to those of applications using aqueous amines and solid oxide adsorbents.

### Experimental

#### 1. Material

Lithium hydroxide (LiOH, 98%), sodium hydroxide (NaOH, 97%), potassium hydroxide (KOH, 99.9%), and boric acid (H<sub>3</sub>BO<sub>3</sub>, 99.5%) were purchased from Sigma-Aldrich. All chemicals were used as received without further purification. All water utilized in the experiments was Milli-Q (Millipore) deionised water.

#### 2. Sample preparation

A series of alkali-metal borate samples  $(A_xB_{1-x}O_y)$ , A: Alkalimetal), were prepared by calcination of mixed precipitants of alkali-metal hydroxide and boric acid prepared in a molar ratio, x, of 0.75. In a typical procedure for the preparation of  $Na_{x^-}B_{1-x}O_y$  (x=0.75, note that electro-neutrality dictates that y=1.5-x), 0.075 mol of sodium hydroxide and 0.025 mol of boric acid were dissolved in 30 ml of distilled water under vigorous stirring. The aqueous solution was dried at 120 °C for several hours to precipitate out the mixed hydroxides. The dried precipitant was ground with an agate mortar and calcined in the oven at 400 °C in air for 4 h. The obtained composite was reground into powders and calcined at 650 °C under a flow of 100%  $N_2$  for 1 h to obtain the final product.

#### 3. Sample characterization and analysis

The CO<sub>2</sub> uptake performance by alkali-metal borates under isothermal conditions was determined by the weight variation of the borate samples under 60 ml min<sup>-1</sup> of 100% dry CO<sub>2</sub> (Airgas) at atmospheric pressure measured by a thermogravimetric analyzer (TGA, TGA Q50; TA Instrument). Here, samples after pre-calcination at 400 °C followed by regrinding were placed on a Pt sample pan in the TGA chamber, and underwent calcination for 1 h at a final temperature of 650 °C under a flow of 100% N2. After calcination at 650 °C, the temperature in the sample chamber was adjusted to the set temperature for the uptake analysis. Measurements of CO2 uptake were initiated by switching the gas introduced to the sample chamber from 100% N<sub>2</sub> to 100% dry CO<sub>2</sub>. The uptake of CO<sub>2</sub> was calculated from the change in sample weight per 1 g of the sample before the introduction of CO<sub>2</sub> gas to the sample chamber by the molecular weight of  $CO_2$  ( $M_{CO_2} = 44.0$ ). The desorption performance at different temperatures was tracked through the decrease in weight under a flow of 100% N2 after the uptake of CO2 at 600 °C under a flow of 100% CO2 for 10 min. CO2 uptake with increasing temperature was examined at a temperature ramp rate of 5 °C min<sup>-1</sup> from 200 °C to 800 °C under the same sample

and gas flow conditions as used in the above isothermal uptake analyses. Cyclic regenerability of CO<sub>2</sub> uptake and desorption by alkali-metal borates in a temperature-swing process was evaluated by repeated switching of the temperature in the TGA sample chamber coupled with changes in the gas flow between 100% CO2 and 100% N2. Pre-treatment of the samples was performed under the same procedure as in the isothermal CO<sub>2</sub> uptake analysis described above. Cyclic regenerability analysis by pressure-swing was performed by switching the gas introduced to the sample chamber between 100% CO2 and 100% N2 at a constant temperature (700 °C). Pre-treatment of the samples was performed as for the isothermal CO2 uptake analysis described above. The melting points of the alkali-metal borates were identified by endothermic peaks in the heat flow with increasing temperature under a flow of 100% N<sub>2</sub> measured by differential scanning calorimetry (DSC, SDT-Q600; TA Instrument). The phase composition and crystallographic features of the samples before and after reaction with CO<sub>2</sub> at the reaction temperature were examined by temperature controlled in situ powder X-ray diffractometry (XRD: PANalytical X'Pert Pro Multipurpose Diffractometer with Cu-K $\alpha$  X-ray ( $\lambda = 1.541$  Å)), in which the samples were placed on a Pt-sheet substrate. The peaks in the XRD spectra were identified by referring to the ICDD PDF-4+2016RDB database. Fluidity of the molten alkalimetal borates at high temperature was evaluated by viscosity measurements on a rheometer equipped with an Environmental Testing Chamber (AR 2000-ETC; TA Instrument). The variation in viscosity as a function of shear rate was measured at a constant temperature (600 °C).

#### Results and discussion

CO<sub>2</sub> uptake by molten alkali-metal borates was examined first with sodium borate,  $Na_xB_{1-x}O_v$  (x = 0.75). Fig. 1(a) shows the uptake of  $CO_2$  (Q [mmol  $g^{-1}$ ]) as the temperature increased at a rate of 5  $^{\circ}$ C min $^{-1}$  under a flow of 100% CO $_2$ . It is clear that the absorption of CO2 by sodium borate at this composition increased dramatically once the temperature was raised to the onset temperature ( $T_{\rm on}$ ) at around 570 °C. After the sharp increase at  $T_{\rm on}$ , the uptake rapidly reached full capacity ( $\sim$ 4.6 mmol g<sup>-1</sup>) in a few minutes and leveled off at higher temperatures. Fig. 1(b) shows the isothermal CO<sub>2</sub> uptake by the same sample at different temperatures, which proceeded at a rapid rate for temperatures greater than 550 °C. The fastest uptake was recorded at 650 °C, where 99% of the full uptake capacity (~5.26 mmol g<sup>-1</sup>) was attained after 1 min reaction with CO<sub>2</sub>. At 500 °C, an initial induction period of a few minutes was evident before initiation of the rapid uptake. At 450 °C, there was no substantial absorption of CO<sub>2</sub>. The sodium borate uptake characteristics at 600 °C are shown in Fig. 1(c) to be significantly different from those of previously-reported solid adsorbents CaO and Li3BO3; the results are normalized by the uptake capacities attained following 1 h reaction with CO<sub>2</sub>. With the solid adsorbents, the uptake rate decreased as the reaction proceeded, resulting in a gradual approach to the full uptake capacity with increasing reaction time. In contrast, the uptake by sodium borate jumped to the full capacity without any

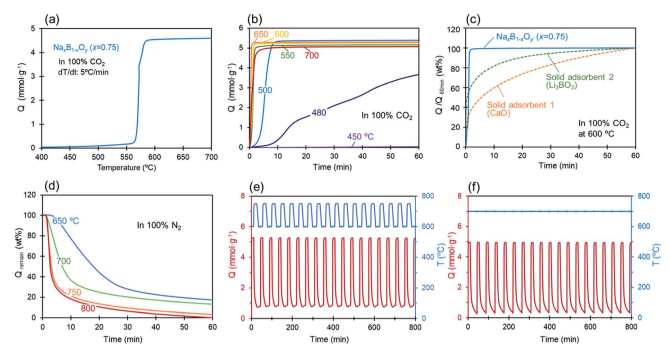


Fig. 1 (a)  $CO_2$  uptake (Q [mmol  $g^{-1}$ ]) by  $Na_xB_{1-x}O_y$  (x=0.75) with temperature (increased at a rate of 5 °C min<sup>-1</sup>) under a flow of 100%  $CO_2$  at atmospheric pressure (1 bar), (b) isothermal  $CO_2$  uptake by  $Na_xB_{1-x}O_y$  (x=0.75) at different temperatures under a flow of 100%  $CO_2$ , (c)  $CO_2$  uptake by  $Na_xB_{1-x}O_y$  (x=0.75), CaO, and  $Li_3BO_3$  at 600 °C, normalized by the uptake capacity after 1 h of reaction, (d)  $CO_2$  desorption curves at different temperatures under a flow of 100%  $N_2$  for  $Na_xB_{1-x}O_y$  (x=0.75) after the uptake of  $CO_2$  under a flow of 100%  $CO_2$  at 600 °C for 10 min, (e) cyclic regenerability of  $CO_2$  uptake and the desorption by  $Na_xB_{1-x}O_y$  (x=0.75); by temperature swing operation with  $\Delta T=150$  °C (at 600 °C under a flow of 100%  $CO_2$ , and at 750 °C under a flow of 100%  $N_2$ ) and, (f) by pressure swing operation on repeated switching of the gas stream between 100%  $CO_2$  ( $pCO_2=1$  bar) and 100%  $N_2$  ( $pCO_2=0$  bar) at 700 °C.

slowing of the uptake rate. Fig. 1(d) shows the CO<sub>2</sub> desorption curves at different temperatures under a flow of 100% N2 for the same sodium borate sample (Na<sub>x</sub>B<sub>1-x</sub>O<sub>y</sub> (x = 0.75)) following CO<sub>2</sub> uptake, from which it is evident that desorption of CO<sub>2</sub> occurs at temperatures higher than 650 °C. Here, CO2 was first released rapidly and then more slowly as desorption proceeded. With higher regeneration temperatures, the desorption proceeded at a faster rate. At 800 °C, desorption of 80% preabsorbed CO2 was completed within 10 min following the introduction of N2 gas to the test chamber. The cyclic regenerability of the sorbents and the extent of desorption are important characteristics of these materials if they are to be used in practical CO<sub>2</sub> capture systems. From Fig. 1(e) and (f) it is clear that  $Na_x B_{1-x} O_v$  (x = 0.75) exhibits excellent cyclic regenerability with no deterioration in uptake capacity nor in reaction rate during cyclic temperature- or pressure-swing operations. Although a residual amount of CO2 was retained by the molten salt at the end of the desorption process, the absorption and desorption performance in later cycles was essentially identical to that of the initial cycle. The excellent cyclic behavior was confirmed up to 100 cycles experimentally, as shown in the ESI (Fig. S1).†

Fig. 2(a) shows the uptake of  $CO_2$  by alkali-metal borates with different alkali-metal species at a temperature increase rate of 5 °C min<sup>-1</sup> under 100%  $CO_2$ . Binary, A = ( $Li_{0.5}Na_{0.5}$ ), and ternary, A = ( $Li_{0.33}Na_{0.33}K_{0.33}$ ), mixtures of the alkali-metals were prepared with equimolar mixtures of these alkali-metal

species. The onset temperature at which the rapid uptake begins shifts to lower temperatures when the sodium is replaced by lithium (i.e.,  $\text{Li}_x B_{1-x} O_y$ ) or by a mixed species that includes lithium ( $(Li_{0.5}Na_{0.5})_xB_{1-x}O_y$ ,  $(Li_{0.33}Na_{0.33}K_{0.33})_xB_{1-x}O_y$ ). With  $\text{Li}_x \text{B}_{1-x} \text{O}_y$ , the uptake accelerated as the temperature increased to a maximum at around 590 °C, followed by a decline at higher temperatures. With the mixed alkali-metal species the uptake rate increased even more sharply, and showed a broad maximum starting at around 570 °C. In the case of potassium borate  $(K_x B_{1-x} O_y)$ , however, the uptake was low over the entire temperature range studied. Cyclic regenerability with no deterioration in capacity upon repeated cycling of CO2 absorption and desorption for the most promising mixed alkali-metal species  $(\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_y$  is evident in Fig. 2(b). Here, the uptake was quite fast, as was the case for sodium borate, and the full uptake capacity of 7.27 mmol  $g^{-1}$  was attained in a few minutes under reaction with CO2.

The results given above demonstrate that, apart from potassium borate, the alkali-metal borates perform well with high  $CO_2$  capacities and uptake rates beyond a well-defined onset temperature ( $T_{\rm on}$ ). A clue as to the temperature dependence of the  $CO_2$  uptake by these alkali-metal borates was provided by DSC results given in Fig. 3(a)–(c) that show changes in the physical state of the different alkali-metal borates with increasing temperature. A clear endothermic peak was observed at  $\sim$ 570 °C with  $Na_xB_{1-x}O_y$  (x=0.75), whereas no peak or only a weak broad peak was apparent in the scans for  $Li_xB_{1-x}O_y$  (x=0.75)

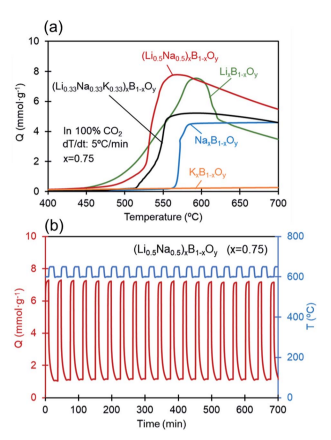


Fig. 2 (a)  $CO_2$  uptake by alkali-metal borates  $(A_xB_{1-x}O_y)$  with different alkali-metal species at x=0.75 with temperature (increased at a rate of 5 °C min<sup>-1</sup>) under a flow of 100%  $CO_2$ , (b) cyclic regenerability under repeated cycles of  $CO_2$  uptake and desorption by  $(Li_{0.5}Na_{0.5})_xB_{1-x}O_y$ .

0.75) and  $K_x B_{1-x} O_y$  (x = 0.75). These results are an indication that the sodium borate melted to form a molten oxide at  $\sim$ 570 °C, whereas the lithium borate and potassium borate did not melt or only melted partially. The fact that the onset temperature at which the steep jump in uptake capacity was observed for the sodium borate coincided with its melting point suggests that the CO2 uptake depends strongly on the physical state of the sorbent. Lithium borate remained in the solid phase up to at least 650 °C. Thus, the moderate rate at which uptake of  $CO_2$  by  $Li_xB_{1-x}O_y$  (x = 0.75) increases with temperature was due to a gas-solid reaction on the surfaces of solid particles, as discussed in the previous report.30 Potassium borate, which was similarly in a solid state, most likely separated into K<sub>2</sub>O and KBO<sub>2</sub>, as inferred from a database in which stable compounds of  $K_x B_{1-x} O_y$  in the range of 0.5 < x < 1.0 appear not to have been confirmed.40 Here, K2O should react instantly with atmospheric CO2 and H2O to form KHCO3 which can be transformed further into K<sub>2</sub>CO<sub>3</sub> during pre-calcination at 650 °C; this carbonate does not decompose back to K2O and CO2. The low CO2 uptake performance by  $K_x B_{1-x} O_y$  (x = 0.75) can be attributed to the low CO<sub>2</sub> uptake capacity of the mixture of KBO<sub>2</sub> and K<sub>2</sub>CO<sub>3</sub>. The physical state and phase composition of two different sodiumcontaining alkali-metal borates,  $Na_xB_{1-x}O_y$  (x = 0.75) and  $(\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_y$  (x = 0.75), at the reaction temperature for

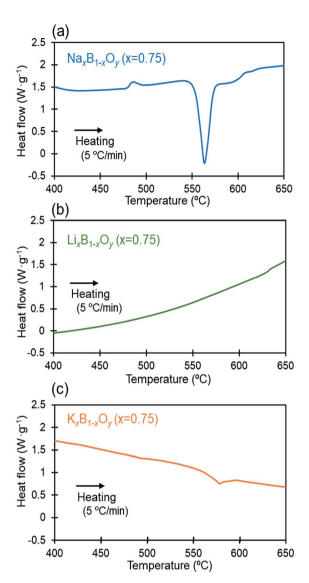


Fig. 3 Heat flow required to increase temperature at a rate of 5 °C min<sup>-1</sup> under 100%  $N_2$  measured by DSC; (a) on  $Na_xB_{1-x}O_y$  (x=0.75), (b) on  $Li_xB_{1-x}O_y$  (x=0.75) and (c) on  $K_xB_{1-x}O_y$  (x=0.75).

CO<sub>2</sub> uptake were examined further by temperature controlled in situ XRD. Fig. 4(a) and (b) show the XRD spectra at 600 °C before reaction with CO2. No distinctive diffraction peaks of solid crystals were observed in the XRD spectra for either compound at 600 °C; only the peaks for the Pt-sheet used as a substrate for XRD analysis registered on the spectra. These results demonstrate that the alkali-metal borates were in a molten (liquid) state at 600 °C before their reaction with CO2. Following reaction with CO2 for 30 min, clearly defined multiple peaks appeared in the XRD spectrum at the same temperature with  $Na_xB_{1-x}O_v$  (x = 0.75), as shown in Fig. 4(c), consistent with solid precipitant generation by the reaction with CO2. Identification of the peaks within the spectrum revealed that solid crystals of α-Na<sub>2</sub>CO<sub>3</sub> and Na<sub>2</sub>B<sub>4</sub>O<sub>7</sub> formed at 600 °C. In contrast, no clear peaks emerged for  $(Li_{0.5}Na_{0.5})_xB_{1-x}O_y$  (x = 0.75) even after reaction with CO<sub>2</sub> for 30 min, as shown in Fig. 4(d), indicating that the uptake of CO<sub>2</sub> by  $(\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_v$  (x = 0.75) proceeds

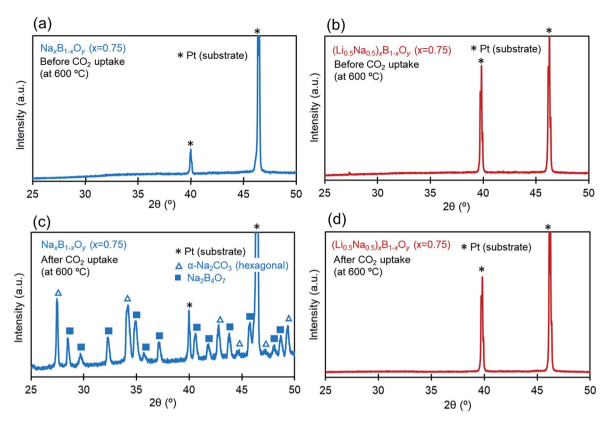


Fig. 4 In situ XRD spectra at  $600 \,^{\circ}$ C for  $Na_xB_{1-x}O_y$  (x = 0.75) and  $(Li_{0.5}Na_{0.5})_xB_{1-x}O_y$  (x = 0.75), before ((a) and (b)) and after ((c) and (d)) the uptake of  $CO_2$  for 30 min.

without the generation of solid precipitants. Since the melting point of the mixed lithium–sodium carbonate is  $\sim 500~^{\circ}\text{C}$  at around the equimolar eutectic composition,<sup>41</sup> it is evident that the mixed carbonates generated by the reaction with  $\text{CO}_2$  existed as ionized molten liquids. Here, the carbonate ions  $(\text{CO}_3^{2^-})$  produced by the reaction of  $\text{CO}_2$  with oxygen anions  $(\text{O}^{2^-})$  in the molten oxides could be stabilized by the coordination of double alkali-metal ions (Li<sup>+</sup> and Na<sup>+</sup>) to the carbonate ions at the reaction temperature (600  $^{\circ}\text{C}$ ).

The dynamic variation in the uptake rate was examined for two of the alkali-metal borates,  $Na_xB_{1-x}O_y$  (x = 0.75) and  $(\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_v$  (x = 0.75), in which solid precipitants were and were not generated by the reaction with CO2, respectively. Fig. 5(a) shows expanded plots of the uptake by these borates at 600 °C, focusing on the initial 2 min of absorption. The uptake rates for the two samples were significantly different. For the case of  $Na_x B_{1-x} O_v$  (x = 0.75), the absorption began at a low rate initially, and then accelerated after a few seconds (t > 0.3 min) contact before leveling off after t > 0.7 min to complete the uptake reaction. In contrast, with  $(\text{Li}_{0.5}\text{Na}_{0.5})_x \text{B}_{1-x} \text{O}_{v}$  (x = 0.75), the uptake increased promptly following a quite short initial induction period, and leveled off on approach to the full uptake capacity, albeit more gradually than in the case of  $Na_xB_{1-x}O_v$  (x = 0.75). The initial induction time with a low uptake rate observed for  $Na_xB_{1-x}O_v$  (x = 0.75) is due to the time required for dissolution of CO<sub>2</sub> to accumulate carbonate ions (CO<sub>3</sub><sup>2-</sup>) in the molten oxides at sufficient levels to exceed the supersaturation

threshold for nucleation and growth of nuclei to form solid sodium carbonate particles (Na<sub>2</sub>CO<sub>3</sub>(s)). The accelerating increase after the initial induction period (t > 0.3 min) was described well by the Avrami-Erofeyev (Erofe'ev or Erofeev) model for nucleation and nucleus-growth, <sup>20,42-44</sup> written as,

$$-\ln(1-\alpha) = kt^n \tag{1}$$

where  $\alpha$  is the conversion ratio ( $Q/Q_{60 \text{ min}}$ ), k is a reaction constant, n is a constant representing the nucleation occurrence and nucleus dimension. Fig. 5(b) shows the left side term of eqn (1) for the uptake data with  $Na_xB_{1-x}O_y$  (x = 0.75) in the accelerating period (0.2 min < t < 0.7 min) as a function of reaction time (t) on a double logarithmic plot. The data in this period are fitted well by the model (shown by the line in Fig. 5(b)), where a power index n of 3.98 was obtained from the linear fitting. This high n value suggests that the growth of the nuclei proceeds three dimensionally with a constant rate consecutive nucleation. For the case of  $(\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_y$  (x=0.75), the uptake increased following a quite short initial induction period, indicating that the accumulation of carbonate ions in the molten oxides proceeded almost immediately as, in contrast to the former case, an induction period for the nucleation of solid precipitants did not occur. The uptake variation with  $(\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_y$  (x=0.75) is consistent with the predictions of the 2D reaction control model, written as43,45

$$[1/(1 - \alpha)] - 1 = kt \tag{2}$$

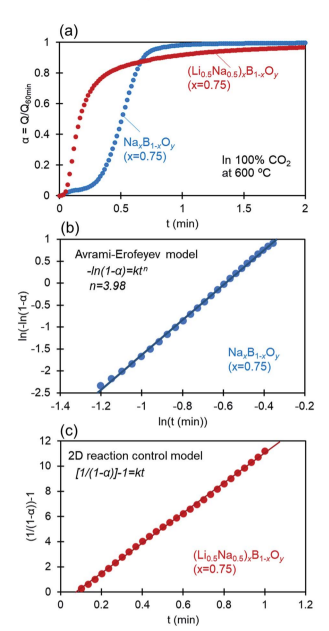


Fig. 5 (a) Variation of CO<sub>2</sub> uptake ratio ( $\alpha=Q/Q_{60~\text{min}}$ ) with Na<sub>x</sub>B<sub>1-x</sub>O<sub>y</sub> (x=0.75) and (Li<sub>0.5</sub>Na<sub>0.5</sub>)<sub>x</sub>B<sub>1-x</sub>O<sub>y</sub> (x=0.75) under a flow of 100% CO<sub>2</sub> at 600 °C over 2 min, (b) double logarithmic plot of the Avrami–Erofeyev equation (eqn (1)) for CO<sub>2</sub> uptake by Na<sub>x</sub>B<sub>1-x</sub>O<sub>y</sub> (x=0.75) over the second period (0.3 min < t<0.7 min). (c) 2D reaction control model (eqn (2)) for CO<sub>2</sub> uptake by (Li<sub>0.5</sub>Na<sub>0.5</sub>)<sub>x</sub>B<sub>1-x</sub>O<sub>y</sub> (x=0.75) as a function of time.

Fig. 5(c) shows the left-hand term of eqn (2) based on the uptake data over the period 0.1 min < t < 1.0 min for  $(\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_y$  (x = 0.75). The linearity of this plot demonstrates that the uptake rate was controlled by a second order reaction, presumably through the coordination of double alkali-metal ions to stabilize the carbonate ions in the molten oxide.

The excellent cyclic regenerability realized by the alkalimetal borates can also be attributed to the liquid behavior of the molten oxides. It is well recognized that deterioration of the uptake performance in repeated uptake and desorption cycles

by solid adsorbents is due to the morphological deformation of solid grains to induce pore-blockage or interparticle sintering. 5,34,46 For the molten oxides, however, deterioration due to morphological deformation does not occur intrinsically due to the fluidic nature of the molten oxide. The rapid mass transfer of ionic species in the molten oxides averages out any defects or inhomogeneities in molecular distribution instantly to maintain the original uptake performance over all operation cycles. Therefore, the cyclic regenerability of the molten oxide is said to be intrinsic to the material.

Viscosity is an important property of any engineering liquid. Fig. 6 shows the viscosity of  $Na_xB_{1-x}O_y$  (x=0.75) and (Li-Na) $_xB_{1-x}O_y$  (x=0.75) in the molten state at 600 °C. The molten ionic oxides exhibit pseudo-plastic shear-thinning behavior; the viscosities for  $Na_xB_{1-x}O_y$  (x=0.75) and (Li-Na) $_xB_{1-x}O_y$  (x=0.75) at a shear rate of 1000 (s<sup>-1</sup>) are 790 cP and 170 cP, respectively. These fluid absorbents can be transferred through transfer lines between the absorber and desorber via simple transfer pumps, and should allow low-cost continuous  $CO_2$  capture systems operating at medium-high temperature to be established.

Table 1 compares the characteristic features of molten ionic oxides for CO2 capture to those of two representative conventional sorbents, aqueous amines and solid oxides. 6,19,25,28,36,47-52 Aqueous amines typically use a gas scrubber operating at nearambient temperatures with simple transfer pumps to deliver the sorbent between the absorber and desorber. Solid oxides operating at medium to high temperatures are far more challenging to transfer between the vessels. Significant effort has been made to address solids handling issues, such as dual fluidized bed reactors with cyclones and conveyer-like moving beds,36,53 but solids handling and system complexity, in addition to uptake deterioration upon cycling, remain key challenges for solid oxides. Molten ionic oxides combine the benefits of the simple liquid transfer system of aqueous amines with the high uptake capacity and high operating temperature of solid sorbents. In contrast to low temperature systems, the streams leaving the process at high temperatures contain valuable heat content which could be recovered in

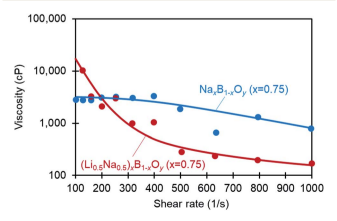


Fig. 6 Viscosity of  $Na_xB_{1-x}O_y$  (x=0.75) and  $(Li_{0.5}Na_{0.5})_xB_{1-x}O_y$  (x=0.75) in the molten state at 600 °C.

Table 1 Comparison of the performances between different sorbents for CO<sub>2</sub> capture

Class of material	Molten ionic oxides	Amines	Solid oxides		
Materials	$Na_xB_{1-x}O_v$ , $(Li-Na)_xB_{1-x}O_v$	MEA, DEA, piperazine, KS-1	CaO, MgO, Li <sub>4</sub> SiO <sub>4</sub> , Li <sub>2</sub> ZrO <sub>3</sub> , Li <sub>3</sub> BO <sub>3</sub>		
Reactor system	Liquid circulation	Liquid circulation	Dual fluidized bed (cyclone, moving bed)		
	(wet scrubbing, bubbling,)	(wet scrubbing, bubbling, EMAR,)			
State of sorbent	Molten	Dissolved in solution	Solid particles		
(active material wt%)	(100 wt%)	(20-30 wt%)	(60–100 wt%)		
Process temperature	500−700 °C	40-60 °C	250-350 °C	500-600 °C	600-700 °C
(sorber)			(MgO)	(Li <sub>4</sub> SiO <sub>4</sub> )	(CaO)
Process temperature	550-800 °C	100–120 °C	350-450 °C	600-700 °C	700–900 °C
(desorber)			(MgO)	(Li <sub>4</sub> SiO <sub>4</sub> )	(CaO)
Enthalpy of reaction	$212 \pm 13 \; { m kJ \; mol^{-1}}$	70–120 kJ mol <sup>–1</sup>	101 kJ⋅mol <sup>-1</sup>	142 kJ⋅mol <sup>-1</sup>	178 kJ⋅mol <sup>-1</sup>
$(\Delta H^0)$	$((\text{Li}_{0.5}\text{Na}_{0.5})_x\text{B}_{1-x}\text{O}_y, (x=0.75))$		(MgO)	(Li <sub>4</sub> SiO <sub>4</sub> )	(CaO)
Material capacity	$\sim$ 5 mmol·g <sup>-1</sup>	$0.5 - 0.75 \text{ mmol} \cdot \text{g}^{-1}$	$\sim$ 15 mmol·g <sup>-1</sup>	$\sim$ 5 mmol·g <sup>-1</sup>	$\sim$ 10 mmol·g <sup>-1</sup>
			(MgO)	(Li <sub>4</sub> SiO <sub>4</sub> )	(CaO)
Working capacity in reactor	$\sim$ 12 mol CO $_2$ per L	$\sim$ 1–2 mol CO $_2$ per L	$\sim$ 1 mol CO $_2$ per L		
	-	-	(with a solids fraction of $\sim$ 2%)		
Rate of uptake	Fast	Moderate	Slow		
	(<1 min to full uptake)	(few minutes to full uptake)	(hours to full uptake)		
Cyclic regenerability	Excellent	Good	Poor		
	(intrinsic to material)	(but material deterioration)	(requires regular supply of fresh material)		
Main Challenges	Gas-liquid contact	uid contact Energy penalty System complexity, grain fragm			nentation,
	-		deterioration of uptake capacity		

a downstream heat recovery steam generator. Therefore, it is expected that molten ionic oxides will have a low energy penalty of capture in an integrated system. Moreover, the volumetric capacity of molten salts under operational conditions (~12 mol  $CO_2$  L<sup>-1</sup>) is significantly greater than that of amine solutions ( $\sim$ 1-2 mol CO<sub>2</sub> L<sup>-1</sup>) and fluidized particles ( $\sim$ 1 mol CO<sub>2</sub> L<sup>-1</sup> with a solids fraction of  $\sim$ 2%) and thus it is anticipated that the equipment size and plant footprint could be reduced significantly. The added benefits of rapid kinetics and intrinsic regenerability are expected to make molten ionic oxides an important class of materials for future CO<sub>2</sub> capture systems.

## Conclusions

Molten alkali-metal borates are proposed as a new class of liquid absorbents that can be exploited at medium to high temperatures, in the range of 500-700 °C, with recovery of high grade heat in secondary steam generators. These sorbents exhibit rapid kinetics for the uptake of CO<sub>2</sub> to attain full uptake capacity of 5.2 to 7.3 mmol g<sup>-1</sup> within 1 min under a flow of CO<sub>2</sub> at atmospheric pressure, with intrinsic cyclic regenerability over repeated cycles of CO<sub>2</sub> uptake and desorption under both temperature- and pressure-swing operations. The rapid CO2 uptake and superior cyclic performance are attributed to the ionic interactions in the melts and the fluidic nature of the molten oxides.

Technologies based on the molten ionic oxides will be applicable to both post- and pre-combustion CO2 capture processes, in particular for next generation high-efficiency power generators, such as integrated combined cycle systems. In post-combustion, CO<sub>2</sub> capture at high temperatures could both minimize the energy penalty by high efficiency heat recovery and reduce the size of the carbon capture system,

resulting in low cost power generation without CO<sub>2</sub> emissions. In pre-combustion, the appropriate installation of capture units for CO2 separation from mixtures of H2 and CO2 could be realized at the temperatures at which coal gasifiers and steam methane reformers with shift reactors operate and thereby improve H<sub>2</sub> production rates through shifts in the reaction equilibria. Thus, molten ionic oxides represent a new paradigm for carbon capture, enabling energy efficient process configurations not previously feasible at high temperatures, and could be well positioned to reshape major segments of the CCUS and energy landscape in years to come.

#### Conflicts of interest

There are no conflicts to declare.

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#### References

- 1 G. C. Institute, The Global Status of CCS: 2017, 2017.
- 2 M. Bui, C. S. Adjiman, A. Bardow, E. J. Anthony, A. Boston, S. Brown, P. S. Fennell, S. Fuss, A. Galindo, L. A. Hackett, J. P. Hallett, H. J. Herzog, G. Jackson, J. Kemper, S. Krevor, G. C. Maitland, M. Matuszewski, I. S. Metcalfe, C. Petit, G. Puxty, J. Reimer, D. M. Reiner, E. S. Rubin, S. A. Scott, N. Shah, B. Smit, J. P. M. Trusler, P. Webley, J. Wilcox and N. Mac Dowell, Energy Environ. Sci., 2018, 11, 1062-1176.

- 3 D. Y. C. Leung, G. Caramanna and M. M. Maroto-Valer, Renewable Sustainable Energy Rev., 2014, 39, 426–443.
- 4 E. Blomen, C. Hendriks and F. Neele, *Energy Procedia*, 2009, 1, 1505–1512.
- 5 S. Choi, J. H. Drese and C. W. Jones, *ChemSusChem*, 2009, 2, 796–854.
- 6 G. T. Rochelle, Science, 2009, 325, 1652-1654.
- 7 B. Dutcher, M. Fan and A. G. Russell, ACS Appl. Mater. Interfaces, 2015, 7, 2137–2148.
- 8 O. Miyamoto, C. Maas, T. Tsujiuchi, M. Inui, T. Hirata, H. Tanaka, T. Yonekawa and T. Kamijo, *Energy Procedia*, 2017, **114**, 5616–5623.
- E. Oko, M. Wang and A. S. Joel, *Int. J. Coal Sci. Technol.*, 2017,
   4, 5–14.
- 10 G. T. Rochelle, Curr. Opin. Chem. Eng., 2012, 1-2, 183-190.
- 11 A. Veawab, P. Tontiwachwuthikul and A. Chakma, *Ind. Eng. Chem. Res.*, 1999, 38, 3917–3924.
- 12 D. Jansen, M. Gazzani, G. Manzolini, E. Van Dijk and M. Carbo, Int. J. Greenhouse Gas Control, 2015, 40, 167–187.
- 13 X. Zhu, Y. Shi, S. Li, N. Cai and E. J. Anthony, in *Pre-combustion Carbon Dioxide Capture Materials*, ed. Q. Wang, Royal Society of Chemistry, 2018, pp. 281–334.
- 14 D. P. Harrison, Ind. Eng. Chem. Res., 2008, 47, 6486-6501.
- 15 D. Berstad, R. Anantharaman and K. Jordal, *Int. J. Greenhouse Gas Control*, 2012, **11**, 25–33.
- 16 J. Strohle, A. Lasheras, A. Galloy and B. Epple, *Chem. Eng. Technol.*, 2009, 32, 435–442.
- 17 A. M. Kierzkowska, R. Pacciani and C. R. Müller, ChemSusChem, 2013, 6, 1130–1148.
- 18 C. Courson and K. Gallucci, in *Pre-combustion Carbon Dioxide Capture Materials*, ed. Q. Wang, The Royal Society of Chemistry, 2018, pp. 144–237.
- 19 V. Manovic and E. J. Anthony, *Environ. Sci. Technol.*, 2009, **43**, 7117–7122.
- 20 T. Harada, F. Simeon, E. Z. Hamad and T. A. Hatton, *Chem. Mater.*, 2015, 27, 1943–1949.
- 21 J. Wang, L. Huang, R. Yang, Z. Zhang, J. Wu, Y. Gao, Q. Wang, D. O'Hare and Z. Zhong, *Energy Environ.Sci.*, 2014, 7, 3478–3518.
- 22 W. Gao, L. Sun and Q. Wang, in *Pre-combustion Carbon Dioxide Capture Materials*, ed. Q. Wang, The Royal Society of Chemistry, 2018, pp. 61–143.
- 23 K. Nakagawa and T. Ohashi, J. Electrochem. Soc., 1998, 145, 1344–1346.
- 24 Q. Xiao, Y. Liu, Y. Zhong and W. Zhu, *J. Mater. Chem.*, 2011, 21, 3838–3842.
- 25 D. J. Fauth, E. A. Frommell, J. S. Hoffman, R. P. Reasbeck and H. W. Pennline, *Fuel Process. Technol.*, 2005, **86**, 1503–1521.
- 26 M. Kato, S. Yoshikawa and K. Nakagawa, *J. Mater. Sci. Lett.*, 2002, 21, 485–487.
- 27 P. V. Subha, B. N. Nair, P. Hareesh, a. P. Mohamed, T. Yamaguchi, K. G. K. Warrier and U. S. Hareesh, *J. Mater. Chem. A*, 2014, 2, 12792–12798.

- 28 M. Seggiani, M. Puccini and S. Vitolo, *Int. J. Greenhouse Gas Control*, 2013, 17, 25–31.
- 29 A. Román-Tejeda and H. Pfeiffer, J. Therm. Anal. Calorim., 2012. 110, 807-811.
- 30 T. Harada and T. A. Hatton, J. Mater. Chem. A, 2017, 5, 22224-22233.
- 31 M. K. Ram Reddy, Z. P. Xu, G. Q. Lu and J. C. D. Da Costa, *Ind. Eng. Chem. Res.*, 2006, **45**, 7504–7509.
- 32 Q. Wang, H. H. Tay, Z. Zhong, J. Luo and A. Borgna, *Energy Environ. Sci.*, 2012, 5, 7526.
- 33 J. Wang, L. Huang, R. Yang, Z. Zhang, J. Wu, Y. Gao, Q. Wang, D. O'Hare and Z. Zhong, *Energy Environ. Sci.*, 2014, 7, 3478–3518.
- 34 R. Barker, J. Appl. Chem. Biotechnol., 1973, 23, 733-742.
- 35 B. Feng, W. Liu, X. Li and H. An, *Energy Fuels*, 2006, **20**, 2417–2420.
- 36 F. Fang, Z. S. Li and N. S. Cai, *Ind. Eng. Chem. Res.*, 2009, **48**, 11140–11147.
- 37 D. Araten, J. Appl. Chem., 1968, 18, 118-121.
- 38 L. Shartsis, W. Capps and S. Spinner, *J. Am. Ceram. Soc.*, 1953, **36**, 319–326.
- 39 O. L. G. Alderman, M. Liška, J. Macháček, C. J. Benmore, A. Lin, A. Tamalonis and J. K. R. Weber, *J. Phys. Chem. C*, 2016, 120, 553–560.
- 40 M. Kowalski, P. J. Spencer and D. Neuschütz, in *Slag Atlas*, ed. Verein Deutscher Eisenhuettenleute (VDEh), Verlag Stahleisen GmbH, 2nd edn, 1996, p. 51.
- 41 G. J. Janz and M. R. Lorenz, *J. Chem. Eng. Data*, 1961, **6**, 321–323.
- 42 T. J. W. De Bruijn, W. A. De Jong and P. J. Van Den Berg, *Thermochim. Acta*, 1981, 45, 315–325.
- 43 A. Khawam and D. R. Flanagan, J. Phys. Chem. B, 2006, 110, 17315–17328.
- 44 M. Avrami, J. Chem. Phys., 1939, 7, 1103.
- 45 Y. S. Ho, J. Hazard. Mater., 2006, B136, 681-689.
- 46 H. Lu, E. P. Reddy and P. G. Smirniotis, *Ind. Eng. Chem. Res.*, 2006, **45**, 3944–3949.
- 47 H. F. Svendsen, E. T. Hessen and T. Mejdell, *Chem. Eng. J.*, 2011, **171**, 718–724.
- 48 S. Ziaii, G. T. Rochelle and T. F. Edgar, *Ind. Eng. Chem. Res.*, 2009, **48**, 6105–6111.
- 49 F. Closmann, T. Nguyen and G. T. Rochelle, *Energy Procedia*, 2009, **1**, 1351–1357.
- 50 I. Kim and H. F. Svendsen, Int. J. Greenhouse Gas Control, 2011, 5, 390-395.
- 51 N. Rodríguez, M. Alonso and J. C. Abanades, *AIChE J.*, 2011, 57, 1356–1366.
- 52 W. Liu, H. An, C. Qin, J. Yin, G. Wang, B. Feng and M. Xu, *Energy Fuels*, 2012, **26**, 2751–2767.
- 53 T. Okumura, K. Yoshizawa, S. Nishibe, H. Iwasaki, M. Kazari and T. Hori, *Energy Procedia*, 2017, 114, 2322–2329.