RSC Advances

PAPER

Cite this: RSC Adv., 2022, 12, 35115

Received 11th May 2022 Accepted 19th November 2022 DOI: 10.1039/d2ra02981b

rsc.li/rsc-advances

Introduction

Air pollution has been a chronic problem and raised increasing concerns since the coronavirus disease 2019 (COVID-19) pandemic.¹⁻³ Nitrogen dioxide $(NO₂)$, as one of the six critical air pollutants according to the United States Environmental Protection Agency (EPA), can result in severe respiratory diseases and devastating environmental issues, such as acid rain, smog, fine particulate matter ($PM_{2.5}$), etc.^{4,5} Therefore, nitrogen oxide reduction $(deNO_x)$ has been a long-term goal with increasingly strict emission standards worldwide. Selective catalytic reduction (SCR) is one of the most common approaches to eliminate NO_x . In SCR systems, NO_x is reduced by ammonia (NH₃) to environment-friendly water (H_2O) and nitrogen (N_2) .⁶ However, NH₃ as a hazardous gas has faced several challenges in its storage and release in automotive SCR systems.^{7,8}

Calcium/strontium chloride impregnated zeolite A and X granules as optimized ammonia sorbents†

Zhejian Cao, \mathbf{D} \ddagger *^a Xiaoping Cai,^b A[na](http://orcid.org/0000-0003-4888-6237) Carolina Feltrin,^a Peizhong Feng,^b Andreas Kaiser \mathbf{P}^c and Farid Akhtar \mathbf{P}^{*a}

Calcium chloride (CaCl₂) impregnated zeolite A and strontium chloride (SrCl₂) impregnated zeolite A and X composite granules were evaluated as ammonia sorbents for automotive selective catalytic reduction systems. The SrCl₂-impregnated zeolite A granules showed a 14% increase in ammonia uptake capacity (8.39 mmol g⁻¹) compared to zeolite A granules (7.38 mmol g⁻¹). Furthermore, composite granules showed 243% faster kinetics of ammonia sorption (0.24 mmol g^{-1} min⁻¹) compared to SrCl₂ (0.07 mmol g^{-1} min⁻¹) in the first 20 min. The composite CaCl₂/SrCl₂ impregnated zeolite A granules combined the advantages of the zeolites and CaCl₂/SrCl₂, where the rapid physisorption from zeolites can reduce the ammonia loading and release time, and chemisorption from the CaCl₂/SrCl₂ offers abundant ammonia capacity. Moreover, by optimizing the content of SrCl₂ loading, the composite granules maintained the granular form with a crushing load of 17 N per granule after ammonia sorption–desorption cycles. Such structurally stable composite sorbents offer an opportunity for fast ammonia loading/release in automotive selective catalytic reduction systems. **PAPER**
 CALCOMP/Strontium chloride impregnated zeolite

CRICIAL CONDINEARTY CONDINEART

Conventional urea-based SCR systems utilize urea $((NH₂)₂CO)$ as an indirect ammonia source, requiring high exhaust temperature and producing carbon dioxide $(CO₂)$ as a byproduct during the hydrolysis reaction.⁹ Furthermore, with a series of problems, such as freezing at low temperatures and catalyst poisoning by the urea residuals, urea-based SCR systems have been replaced by solid SCR systems in several countries.^{10–12} In solid SCR systems, $NH₃$ is stored in solid form, typically in the form of metal ammine complexes by alkaline earth metal halides (AEMHs).^{13,14} The AEMHs demonstrate excellent ammonia storage capacity.¹⁵ With direct ammonia dosing, the deNO_x efficiency of the solid SCR system has been enhanced at low exhaust temperatures.¹⁴ Therefore, various research and applications on AEMHs as ammonia carriers have been studied, including hydrogen storage, heat pumps, etc.¹⁶⁻¹⁹ Nevertheless, AEMHs as ammonia carriers emerge several shortcomings impeding the applicable scope. For instance, $CaCl₂$ and $SrCl₂$ expand up to 4 times by volume after complete ammonia absorption and generate 70% porosity.^{20,21} These dramatic volume changes can result in the disintegration of the structured AEMHs into powder. In automotive SCR systems, this poor structural stability of the ammonia carriers can lead to safety risks in the vehicles and uncontrollable ammonia dosing performance, such as inefficient use of space and pressure drop.22,23 Moreover, it has been reported that in these applications AEMHs with slow kinetics require considerable time (up to 36 hours for $CaCl₂$) to achieve a complete ammonia sorption cycle.²⁴ Slow kinetics of ammonia absorption and desorption in AEMHs can result in long ammonia loading and release time;

[&]quot;Division of Materials Science, Luleå University of Technology, 971 87, Luleå, Sweden. E-mail: zhejian@chalmers.se; farid.akhtar@ltu.se

b School of Materials Science and Physics, China University of Mining and Technology, 221116, Xuzhou, People's Republic of China

Department of Energy Conversion, Technical University of Denmark, 2800 Kgs. Lyngby, Denmark

[†] Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d2ra02981b>

[‡] Present Address: Division of Systems and Synthetic Biology, Department of Biology and Biological Engineering, Chalmers University of Technology, 41296 Gothenburg, Sweden.

especially in the first 5–10 min when the vehicles have a cold start where the tank of ammonia sorbents is far below the threshold temperature, the long ammonia release time can lead to unexpected NO_x escape, which can hinder the development of the NO_x emission standard.^{23,25,26} Materials providing rapid ammonia sorption and releasing kinetics, therefore, is of desire to elevate the performance of SCR systems.

Microporous materials, including zeolites, metal–organic frameworks (MOFs), activated carbon, etc., have been widely studied for gas adsorption applications owing to their high specific surface area.²⁷⁻³⁰ Among them, zeolites A (Linde type A) and X (faujasite type X) have been intensively investigated due to their excellent chemical/thermal stability, and industrial maturity. By tailoring the zeolites pore opening with the ionexchange method, the gas uptake capacity and separation can be modified for specific applications, such as methane CH_4) $CO₂$ separation in biogas.^{31,32} However, the study of zeolites as ammonia carriers have been rarely discussed due to their relatively low ammonia uptake capacities (9.3 mmol g^{-1} in NaX, 7.8 mmol g^{-1} in CaA) compared to AEMHs (63.0 mmol g^{-1} in CaCl₂, 50.5 mmol g^{-1} in SrCl₂).^{15,33–35} Ammonia sorption mechanisms are reported to be different in zeolites compared to AEMHs. In AEMHs, ammonia molecules are strongly absorbed via the formation of coordination complexes in a chemisorption process. For example, $CaCl₂$ absorbs 8 ammonia molecules according to eqn (1)–(3), where the ammonia desorption energy (69 kJ mol⁻¹) is higher than 40 kJ mol⁻¹.^{15,36-38} Zeolites, as physisorbents, absorb ammonia with weak interaction, where the major part of the ammonia molecules are released with desorption energies below less than 40 kJ mol−¹ (for example 20 kJ mol⁻¹ for LTA).^{39,40} The physisorption of zeolites allows a lower energy penalty of ammonia release and rapid ammonia sorption kinetics.⁴¹–⁴³ Therefore, the fast kinetics of gas sorption in zeolites and their excellent structural stability present RSC Advances Wexture of the different interest are ended a position to overcome some limitations of the use of the state of animalism and th

a potential solution to overcome some limitations of the use of pure AEMH structures under fast $NH₃$ sorption/desorption cycles.⁴⁴

$$
CaCl2 + 2NH3 \rightleftharpoons Ca(NH3)2Cl2
$$
 (1)

$$
Ca(NH_3)_2Cl_2 + 2NH_3 \rightleftharpoons Ca(NH_3)_4Cl_2 \tag{2}
$$

$$
Ca(NH3)4Cl2 + 4NH3 \rightleftharpoons Ca(NH3)8Cl2
$$
 (3)

In this study, we designed zeolite–AEMH composites by impregnating $CaCl₂/SrCl₂$ into zeolite A and X granules. The zeolite retained the crystal structure after the ion-exchange and chloride-impregnation process. The resulting zeolite–AEMH composite granules were characterized by various methods to evaluate the structural stability after ammonia sorptiondesorption cycles. Furthermore, the changes in the ammonia uptake capacity and ammonia sorption kinetics of the composite granules were compared, analyzed, and discussed regarding the pristine zeolites and AEMH materials.

Experimental section

Materials and methods

Zeolite granules CaA and NaX (granule size 1.6–2.5 mm, Luoyang Jalong Micro-nano New Materials Co., Ltd., Henan, China), calcium chloride (anhydrous, 93% purity, Alfa Aesar) and strontium chloride (anhydrous, 99% purity, Alfa Aesar) were purchased and used as pristine materials. To reduce the formation of unexpected salts, e.g., NaCl, during the impregnation process, CaA and NaX granules were first treated by ion exchange to replace the Ca^{2+} and Na⁺ cations with Sr^{2+} . The impact of time on the ion-exchange process and the concentration of the $SrCl₂$ solution during the ion-exchange process were investigated (Section S1, ESI†). All the ion-exchanged

Fig. 1 Fabrication process of zeolite–AEMH composites. (a)–(c) The ion-exchange process of CaA and NaX granules, including stirring, rinsing, and drying. (d)–(f) The AEMHs impregnation process of the selected zeolite granules, including granule selection, AEMHs impregnation, and drying.

Table 1 The code for different impregnated granules and their processing parameters

Granule code	Composition	Ion-exchange conditions		Impregnation AEMHs loading $(g^{-1}$ granule)		
Sr_X Sr_A Ca_A_L Ca_A_M Ca_A_H	$SrCl2$ impregnated X SrCl ₂ impregnated A $CaCl2$ (low loading) impregnated A CaCl ₂ (medium loading) impregnated A $CaCl2$ (high loading) impregnated A		3 times with 0.40 g mL^{-1} SrCl ₂ solution 3 times with 0.27 $g \text{ mL}^{-1}$ SrCl ₂ solution	0.81 g $SrCl2$ (45 wt%) 0.27 g SrCl ₂ (21 wt%) 0.14 g CaCl ₂ (12 wt%) 0.27 g CaCl ₂ (21 wt%) 0.54 g CaCl ₂ (35 wt%)		
	granules were dried at $150 \degree C$ for 1 h. As-received CaA granules were directly impregnated with CaCl ₂ solution since the formation of byproduct salts was not expected. The impregna- tion process was done by dripping the AEMHs solution to the granules as shown in Fig. 1(f). The loading of the AEMHs solution was from 12 wt% to 45 wt% (Section S2, ESI [†]). After the impregnation, the granules were dried at 150 \degree C for 1 h. The processing conditions of the obtained impregnated granules			sorption, and then back to high vacuum at the fast speed of the machine for the desorption to characterize the kinetics perfor- mance, especially in the first 10 min regime. The equilibrium of the ammonia sorption-desorption was set until the standard deviation of mass was less than 0.1 mg per 10 min. The falling AEMHs and the loose AEMHs due to volume changes on the		
are listed in Table 1.			(a)			
	Structure characterization			A Zeolite X		
	The microstructure and elemental composition of the pristine materials and the impregnated zeolite granules were charac- terized by scanning electron microscopy-energy dispersive X-ray		Intensity (a.u.)	SrCl, SriX		
	spectroscopy (SEM-EDS, JSM-IT300LV, JEOL GmbH, Germany), with a 15 nm platinum coating (Leica EM ACE 200, Germany) on the tested granules to avoid charging up from incident elec-					

Structure characterization

The microstructure and elemental composition of the pristine materials and the impregnated zeolite granules were characterized by scanning electron microscopy-energy dispersive X-ray spectroscopy (SEM-EDS, JSM-IT300LV, JEOL GmbH, Germany), with a 15 nm platinum coating (Leica EM ACE 200, Germany) on the tested granules to avoid charging up from incident electrons. The crystal structure of the pristine materials and the impregnated zeolite granules was characterized by a Cu Ka radiation X-ray diffractometer (XRD, Empyrean, PANalytical, United Kingdom). All the granules were crushed and ground to a fine powder for XRD measurements. The specific surface area of the granules was obtained by N₂ adsorption at −196 °C with the Brunauer–Emmett–Teller (BET) model using a surface area analyzer (Gemini VII 2390, Micromeritics, Norcross, USA). All measured materials were degassed at 300 °C under a dynamic vacuum overnight before the BET surface area measurements. The crushing load of the granules was measured by loading one granule (diameter 2.2 ± 0.1 mm) for a compression test using a universal machine (WDW-100, Jinan Hensgrand Instrument Co., Ltd., China), with the loading of strain rate at 1.5% s^{-1} . 5 granules of each composition were measured for an average crushing load to obtain statistical reliability.

Ammonia sorption and desorption measurement

The ammonia sorption and desorption performance were characterized by an IsoSORP® sorption analyzer (TA Instruments, United States), which consists of a magnetic suspension balance inside a chamber, an electrical heater for degassing, and a chemistry diaphragm vacuum pump to reach high vacuum. All granules were degassed at 300 °C under a high vacuum for 3 hours, following a buoyancy test with helium at 22 °C to determine the mass and volume of the tested material. Then, the ammonia sorption–desorption performance was measured with ammonia dosing from high vacuum to 1 bar (above the $Sr(NH₃)₈Cl₂$ equilibrium pressure of 0.4 bar) for the

Fig. 2 The XRD pattern of the zeolites. (a) Pristine NaX and Sr_X before ammonia test; (b) pristine NaX, Sr^{2+} ion-exchanged zeolite X, and Sr_X after 2-cycle and 10-cycle ammonia test; (c) pristine CaA, Sr^{2+} ionexchanged zeolite A and Sr_A after ammonia test.

RSC Advances **Paper** Paper **Paper Paper Paper**

surface of granules after the ammonia cycles were removed by a 12-mesh sieve for the second cycle of ammonia measurement. For the cyclic stability characterization, 10 cycles (counting after removing the detached AEMHs) of the ammonia test were performed after removing the falling AEMHs, with ammonia dosing from high vacuum to 3 bar for the sorption, and then back to high vacuum for the desorption.

Results and discussion

The crystal structure of the zeolites was characterized by XRD. Sr_X contains a high $SrCl_2$ loading of 45 wt%, resulting in one main peak of $SrCl₂$ and weak characteristic peaks from zeolite X marked in red triangles, as shown in the X-ray diffractograms in Fig. 2(a). To verify the stability of the crystal structure of zeolites, the XRD of the impregnated granules was measured after ammonia sorption, where most of the $SrCl₂$ on the surface of granules disintegrated due to volume changes and was removed by sieving. Characteristic peaks of $SrCl₂$ were observed with low intensity resulting from small amounts of $SrCl₂$ inside the granules and attached to the surface of the granules. After the Sr^{2+} ion exchange and $SrCl₂$ impregnation process, the crystal structure of the zeolite X and A maintained the crystallinity. This is attributed to the crystal stability of the zeolite frameworks and the robust ion-exchange method.45,46 However, the macro-structure of the obtained granules was different before and after the ammonia sorption–desorption measurement, as shown in Fig. 3. Pristine NaX and CaA granules showed a smooth surface and a spherical morphology. Due to the high loading (45 wt%) of the $SrCl₂$ in Sr_X granules, the zeolite X granules were covered by the $SrCl₂$ shell (Fig. 3(c)). The amount of the SrCl₂ shell could be controlled by the SrCl₂ loading during the impregnation process (Section S2, ESI†). A high loading was chosen in Sr_X to achieve high ammonia uptake capacity from the AEMHs. However, due to the dramatic volume expansion

(400%) of the SrCl₂ during ammonia absorption, the thick $SrCl₂$ shell fell from the zeolite X granules.⁴⁷ Furthermore, when the $SrCl₂$ inside the granules expanded, it resulted in the formation of cracks in the granule, as shown in Fig. 3(d). When the ionexchange process was performed in high concentrated $SrCl₂$ solution, cracks were formed in the CaA granules (Section S1, ESI†). Therefore, impregnation was performed at lower concentration (SrCl₂ solution with 0.27 g mL⁻¹) for the preparation of the Sr_A sample. After lowering the concentration of the impregnation solution, the $SrCl₂$ loading was reduced to 21 wt% and few $SrCl₂$ agglomerations were observed on the zeolite A granules as shown in Fig. $3(e)$. After the ammonia sorption–desorption test (Fig. $3(f)$), SrCl₂ agglomerates seem to partially detach from the surface of zeolite A granules, but crack formation in the granules was not observed, which suggests that a reduced $SrCl₂$ loading in the zeolite A granules can provide better structural stability. However, in the Ca_A_L granules, with low impregnation loading (12 wt%), granule crack formation was observed, as shown in Fig. 3(h). When the $CaCl₂$ loading increased as in Ca_A_M (21 wt%) and Ca_A_H $(35 wt%)$, the degree of destruction of the granules after the ammonia test was exacerbated (Section S3, ESI†). This could be explained by the lower density of CaCl₂ (2.15 $\rm g$ cm^{−3}) compared with SrCl₂ (3.05 g cm⁻³). For the same mass loading, CaCl₂ requires more space to expand, which would result in higher stress in the granules.^{15,47} RSC Advances

Super Access Article and Super Access Article 2022. The stationary of the Stationary and the Stationary and the Common Access Article is licensed to the article is are also that is a stationary and the commo

The influence of the crack formation induced by the expansion on the mechanical performance was investigated by compression test for the granules after ammonia tests. As listed in Table 2, the crushing load reduced in the impregnated granules. Particularly in Sr_X and Ca_A_L, only 43% and 17% crushing load remained after the ammonia sorption-desorption test, respectively. Sr_A retained 74% crushing load (17 N per granule) compared to the pristine CaA granules (23 N per granule), suggesting good structural stability after ammonia

Fig. 3 SEM images of the granules. (a) Pristine NaX; (b) pristine CaA; (c) pristine Sr_X with SrCl₂ shell covering the zeolite X granule; (d) Sr_X after ammonia test with the SrCl₂ shell detaching and crack formation in the granule; (e) pristine Sr_A with SrCl₂ on the granule surface; (f) Sr_A after ammonia test with partial detachment of SrCl₂ shell; (g) pristine Ca_A_L with CaCl₂ on the granule surface; (h) Ca_A_L after ammonia test with partial CaCl₂ shell detachment and crack formation in the granule.

Table 2 The crushing load of the granules after the ammonia sorption–desorption measurement

Granules	NaX	CaA	Sr X Sr A	Ca A L
Crushing load (N/granule) 21 ± 4 23 ± 4 9 ± 3 17 ± 5 4 ± 1				

sorption–desorption cycles. The diametral compressive strength of the Sr_A was 4.5 MPa, which is higher than the mechanical strength reported for zeolite monoliths and granules after other treatments.^{31,44} The mechanical performance loss compared to the pristine granules could be attributed to 2 processes. First, as also reported in other studies, the repeated immersion process of the granules in the solution during the ion-exchange process might lead to a dissolution of binding points of the zeolite crystals in the granules, resulting in a drop in the compressive strength.³¹ Secondly, as shown in the SEM images in Fig. 3, the Sr_X and Ca_A_L granules showed crack formation in the granules after the ammonia sorption measurements, resulting in a lower compressive strength. The Sr A material did not reveal such obvious crack formation and retained most of the crushing load. The Ca_A_M and Ca_A_H granules with the high $CaCl₂$ impregnation loading were broken into pieces after the ammonia test, which was attributed to the large expansion during ammonia sorption–desorption cycles.

The ammonia sorption and desorption performance of the granules were evaluated by the ammonia uptake capacity and the kinetics as shown in Fig. 4. The experimental ammonia uptake capacity of the NaX and CaA was 10.36 mmol g^{-1} and 7.38 mmol g^{-1} , respectively, which was much lower than the measured uptake of SrCl₂ with 46.97 mmol g^{-1} . After the ionexchange process, the partially Sr^{2+} ion-exchanged zeolite X

(IE X) and A (IE A) granules showed a small drop of 8% in IE X and 3% in IE A in the ammonia uptake capacity, which could be attributed to the surface area decrease after the ion-exchange process, as indicated in Fig. 4(d). On the contrary, the ammonia uptake capacity of the impregnated granules Sr_X and Sr_A increased in the first cycle despite the decrease in the surface area by 74% (18.01 mmol $\rm{g}^{-1})$ and 19% (8.80 mmol $\rm{g}^{-1})$, respectively. This can be attributed to the high ammonia capacity for samples highly impregnated with SrCl₂.⁴⁸ After the first ammonia sorption-desorption cycle, it was observed that the $SrCl₂$ particles on the surface detached from the zeolite granules. These loose $SrCl₂$ particles were removed by a 12mesh sieve and separated from the granules. As a result of the loss of $SrCl₂$ from the surface of the $SrCl₂$ –zeolite composite granules, the ammonia uptake capacity decreased signicantly to 10.98 mmol g^{-1} and 8.39 mmol g^{-1} in Sr_X and Sr_A (green bar in Fig. 4(a)), resulting in a moderate 6% and 14% increase of the ammonia capacity compared to the pristine zeolite granules, respectively. AEMHs and zeolites were reported with an excellent cyclic performance of ammonia sorption and desorption.33,47,49,50 The zeolite crystal structure maintains identical after 10 cycles in Sr_X according to the XRD patterns as shown in Fig. 2, and additional detachment of AEMH material from the composite granules was not observed after the second ammonia sorption–desorption cycle, suggesting Sr_X and Sr_A possess stable structure and cyclic performance of ammonia sorption and desorption after removing the falling AEMHs. After the removal of the loose, detached S_{IC} particles from the composite surface by sieving, the amount of $SrCl₂$ that was well attached to the Sr_X and Sr_A composite granules could be estimated from the increase in ammonia uptake compared to the ion-exchanged zeolites (without $SrCl₂$ impregnation). The **Paper**
 Totics 2 The critical control and the particle article on 02 C-11/2 and A(E article is limited and dop of 9% in Hz interactions

to 2022. Note that the common particle is limited and the common particle is limi

Fig. 4 (a) The ammonia uptake capacity of the granules; (b) the ammonia sorption curves in zeolite X granule series; (c) the ammonia desorption curves in zeolite X granule series; (d) the BET surface area of the granules; (e) the ammonia sorption curves in zeolite A granule series; (f) the ammonia desorption curves in zeolite A granule series.

Table 3 Ammonia desorption capacity and the corresponding desorption percentage of the granules and SrCl₂

 $SrCl₂$ loading in Sr X and Sr A after ammonia sorption were 4 wt% and 3 wt%, respectively (for details on the calculations, see Section S4, ESI†). Fig. 4(b) and (e) reveal the kinetics of the granules in the ammonia sorption process. After the first 20 min, the pristine zeolite and both ion-exchanged zeolites, the X and the A zeolite, reached about 80% of their final, saturated ammonia uptake capacity, showing an excellent ammonia sorption kinetics. In contrast, the pure $SrCl₂$ did not start to absorb ammonia before the ammonia pressure reached the equilibrium ammonia vapor pressure of 0.4 bar after 23 min.¹⁵ For the Sr_X and Sr_A composite granules, we observed a two-stage process of ammonia sorption. In the first 20 min (stage one), the second cycle Sr_X and Sr_A granules showed ammonia uptake capacities of 3.6 mmol g^{-1} and 4.8 mmol g^{-1} , resulting in a rate of ammonia sorption at 0.18 mmol g^{-1} min $^{-1}$ and 0.24 mmol g^{-1} min $^{-1}$, respectively. In stage two (after 50 min), we observed ammonia uptake with a reduced sorption rate, which reflects the slower chemisorption process of ammonia sorption in $SrCl₂$ compared to the physisorption process of the zeolite material in the composite. Notably, the rate of ammonia sorption in Sr_A composite granules with 0.24 mmol g^{-1} min⁻¹ was 243% faster compared to SrCl₂ with a rate of 0.07 mmol g^{-1} min⁻¹ in the first 3 hours at 1 bar ammonia pressure. Such rapid kinetics in ammonia sorption in Sr_X and Sr_A composite granules can offer a quick loading of the ammonia cartridges and increase the cycle efficiency.²² The ammonia desorption curves of the granules were plotted in Fig. $4(c)$ and (g) . With the pressure swing adsorption (PSA) method, no prominent difference in the desorption rate was found between granules and $SrCl₂$, due to the instant high vacuum condition. However, the rapid kinetics of the physisorption of ammonia has been reported before with a temperature swing adsorption (TSA) method, RSC Advances www.water.components are the correspondent detection percentage of the granules are $\frac{3\pi\lambda}{24}$ and $\frac{3\pi\lambda}{24}$ and $\frac{3\pi\lambda}{24}$ are the components are the components are $\frac{3\pi\lambda}{24}$ and $\frac{3\pi\lambda}{$

where it took a relatively long time to reach the decomposition temperature for the AEMHs ammines.^{34,51,52} In our previous study, the zeolite X demonstrated 50% higher ammonia release in the first 10 min at low temperature (35 °C), suggesting the physisorbents can offer a rapid ammonia dosing in SCR at a lower temperature compared to the chemisorbents of AEMHs.⁵³ The TSA measurement results show that zeolite X releases 4 times higher ammonia (0.69 mmol $\rm g^{-1})$ than \rm{SrCl}_2 (0.14 mmol g^{-1}) in the first 10 min before reaching 60 °C (Section S5, ESI†). Therefore, combining physisorption and chemisorption in the zeolite–AEMH composites can expand the working temperature window of SCR systems. We observed that with the PSA method, the ammonia in the pristine zeolite and ion-exchanged zeolite granules demonstrated a relatively low desorption efficiency, as shown in Table 3. Less than 50% of the absorbed ammonia was desorbed in pristine NaX and CaA granules, while $SrCl₂$ possessed tremendous desorption efficiency at 89%. By combining the 2 parts from zeolite and SrCl₂, Sr_X and Sr_A granules yielded 64% and 48% desorption efficiency, respectively.

To simulate the practical conditions for a further cyclic stability characterization, 10 cycles of ammonia sorption tests were performed at ammonia pressure at 3 bar for Sr_X. As shown in Fig. 5(a), the ammonia uptake capacity was maintained at over 92% after 10 cycles of ammonia sorption and desorption. The ammonia sorption percentage curves are identical in ammonia sorption and desorption for 10 cycles according to Fig. 5(b) and (c). The XRD patterns (Fig. 2(b)) and SEM images (Section S6, ESI†) of the Sr_X maintained stable after 10 cycles after removing the detaching $SrCl₂$. All these results indicate good cyclic stability of the zeolite–AEMH composites as a long-term practical ammonia sorbent.

Fig. 5 The cyclic performance of ammonia sorption and desorption in Sr_X. (a) The ammonia uptake capacity of Sr_X at 22 °C, 3 bar in 10 cycles. (b) The ammonia sorption percentage of Sr_X in the ammonia absorption for 10 cycles from high vacuum to 3 bar. (c) The ammonia sorption percentage of Sr_X in the ammonia desorption for 10 cycles from 3 bar to high vacuum.

Based on the micro- and macrostructure, mechanical test, and the ammonia adsorption–desorption measurement, the Sr_A granule demonstrated an excellent combination between the zeolite and AEMHs, offering a solution of structural stable ammonia sorbents with rapid kinetics and elevated ammonia capacity.

Conclusions

A simple impregnation method to combine calcium/strontium chloride and commercial zeolite A and X granules was designed. The optimized Sr_A granule demonstrated the best structural stability in terms of crystal structure and mechanical strength after ammonia sorption-desorption cycles. Moreover, Sr_A granules possessed 14% higher ammonia uptake capacity compared to the pristine CaA zeolites, and rapid ammonia kinetics with a 243% faster sorption rate than the $SrCl₂$ in the first 20 min. The results confirmed two stages for the ammonia sorption in Sr_A, first physisorption followed by chemisorption. This might open a new view to solve the current challenges of the AEMHs as ammonia sorbents in SCR systems. Moreover, by adjusting the ratio of the AEMHs loading, the performance of composite granule ammonia sorbents can be tailored for various other potential applications, such as clean fuels, and hydrogen storage. Paper

Based on the microsoft on distribution interactions, the based on 12 December 2022. At iterations

and the microsoft of the continue o

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

This work was supported by grants from Formas (No. 2016- 01099). The authors acknowledge the lab assistance from Kyllian Demenus, Pauline Léonard, and Yang Yu. The authors acknowledge the Wallenberg Foundation "Jubileumsanslaget" travel grants for supporting Z. Cao's visiting trip to the China University of Mining and Technology (CUMT). F. A. acknowledges the support from Swedish Research Council, grant no. 2018-04407.

References

- 1 M. Kampa and E. Castanas, Environ. Pollut., 2008, 151, 362– 367.
- 2 C. Copat, A. Cristaldi, M. Fiore, A. Grasso, P. Zuccarello, S. S. Signorelli, G. O. Conti and M. Ferrante, Environ. Res., 2020, 191, 110129.
- 3 Y. Ogen, Sci. Total Environ., 2020, 726, 138605.
- 4 T. W. Hesterberg, W. B. Bunn, R. O. McClellan, A. K. Hamade, C. M. Long and P. A. Valberg, Crit. Rev. Toxicol., 2009, 39, 743–781.
- 5 T. Boningari and P. G. Smirniotis, Curr. Opin. Chem. Eng., 2016, 13, 133–141.
- 6 G. Ramis, L. Yi, G. Busca, M. Turco, E. Kotur and R. J. Willey, J. Catal., 1995, 157, 523–535.
- 7 D. Damma, P. R. Ettireddy, B. M. Reddy and P. G. Smirniotis, Catalysts, 2019, 9, 349.
- 8 J. J. Renard, S. E. Calidonna and M. V. Henley, J. Hazard. Mater., 2004, 108, 29–60.
- 9 T. Feng and L. Liu, J. Ind. Eng. Chem., 2015, 28, 97–109.
- 10 M. Koebel, M. Elsener and M. Kleemann, Catal. Today, 2000, 59, 335–345.
- 11 K. Shimizu and A. Satsuma, Appl. Catal., B, 2007, 77, 202– 205.
- 12 J. Han, T. Kim, H. Jung, S. Pyo, G. Cho, Y. Oh and H. Kim, Int. J. Automot. Technol., 2019, 20, 1153–1160.
- 13 F. Lacin, A. Kotrba, G. Hayworth, H. Sullivan, M. Tatur, J. Jacques, D. Tomazic and H. Cho, SOLID, SCR®: Demonstrating an Improved Approach to NOx Reduction via a Solid Reductant, SAE Technical Paper 2011-01-2207, 2011, DOI: [10.4271/2011-01-2207](https://doi.org/10.4271/2011-01-2207).
- 14 T. D. Elmøe, R. Z. Sørensen, U. Quaade, C. H. Christensen, J. K. Nørskov and T. Johannessen, Chem. Eng. Sci., 2006, 61, 2618–2625.
- 15 G. Fulks, G. B. Fisher, K. Rahmoeller, M.-C. Wu, E. D'Herde and J. Tan, SAE Tech. Pap., 2009, 01, 907–908.
- 16 C. Hviid Christensen, R. Zink Sørensen, T. Johannessen, U. J. Quaade, K. Honkala, T. D. Elmøe, R. Køhler and J. K. Nørskov, J. Mater. Chem., 2005, 15, 4106–4108.
- 17 A. Klerke, C. H. Christensen, J. K. Nørskov and T. Vegge, J. Mater. Chem., 2008, 18, 2304.
- 18 Z. Yang, M. Qu and K. R. Gluesenkamp, Appl. Therm. Eng., 2020, 179, 115674.
- 19 S. A. H. Mofidi and K. S. Udell, J. Energy Resour. Technol., 2017, 139, 032005.
- 20 H. S. Jacobsen, H. A. Hansen, J. W. Andreasen, Q. Shi, A. Andreasen, R. Feidenhans'l, M. M. Nielsen, K. Ståhl and T. Vegge, Chem. Phys. Lett., 2007, 441, 255–260.
- 21 J. S. Hummelshøj, R. Z. Sørensen, M. Yu. Kustova, T. Johannessen, J. K. Nørskov and C. H. Christensen, J. Am. Chem. Soc., 2006, 128, 16–17.
- 22 T. Johannessen, H. Schmidt, A. M. Frey and C. H. Christensen, Catal. Lett., 2009, 128, 94–100.
- 23 D. Chakraborty, H. N. Petersen, C. Elkjær, A. Cagulada and T. Johannessen, Fuel Cells Bull., 2009, 2009, 12–15.
- 24 A. B. Hart and J. R. Partington, J. Chem. Soc., 1943, 104–118.
- 25 J. Chang, G. Kalghatgi, A. Amer, P. Adomeit, H. Rohs and B. Heuser, SAE Int. J. Engines, 2013, 6, 101–119.
- 26 J. Johansen, J. Oechsle, H. Schmidt, T. Johannessen and J. Svagin, Release of stored ammonia at start-up. Classifications: C01C1/006 Storage or handling of ammonia making use of solid ammonia storage materials, e.g. complex ammine salts, U.S. Pat., US8491842B2, 2013, <https://patents.google.com/patent/US8491842>.
- 27 O. M. Yaghi, ACS Cent. Sci., 2019, 5, 1295–1300.
- 28 S. Stankovich, D. A. Dikin, G. H. B. Dommett, K. M. Kohlhaas, E. J. Zimney, E. A. Stach, R. D. Piner, S. T. Nguyen and R. S. Ruoff, Nature, 2006, 442, 282.
- 29 O. Cheung, Z. Bacsik, Q. Liu, A. Mace and N. Hedin, Appl. Energy, 2013, 112, 1326–1336.
- 30 X. Han, W. Lu, Y. Chen, I. da Silva, J. Li, L. Lin, W. Li, A. M. Sheveleva, H. G. W. Godfrey, Z. Lu, F. Tuna,

E. J. L. McInnes, Y. Cheng, L. L. Daemen, L. J. M. McPherson, S. J. Teat, M. D. Frogley, S. Rudić, P. Manuel, A. J. Ramirez-Cuesta, S. Yang and M. Schröder, J. Am. Chem. Soc., 2021, 143, 3153–3161.

- 31 K. Narang, K. Fodor, A. Kaiser and F. Akhtar, RSC Adv., 2018, 8, 37277–37285.
- 32 O. Cheung and N. Hedin, RSC Adv., 2014, 4, 14480–14494.
- 33 J. Helminen, J. Helenius, E. Paatero and I. Turunen, J. Chem. Eng. Data, 2001, 46, 391–399.
- 34 H. G. W. Godfrey, I. da Silva, L. Briggs, J. H. Carter, C. G. Morris, M. Savage, T. L. Easun, P. Manuel, C. A. Murray, C. C. Tang, M. D. Frogley, G. Cinque, S. Yang and M. Schröder, Angew. Chem., Int. Ed., 2018, 57, 14778– 14781. PSC Advances Weekendon Commons, L. B. McTures Articles. Published on 07 December 2022. Downloaded on 2022. Downloaded on 2022. Downloaded on 2022. The State Att December 2022. Downloaded on 11/2000 New Article is licensed
	- 35 S. Wu, T. X. Li and R. Z. Wang, Energy, 2018, 161, 955–962. 36 J. J. Pitts, J. Cell. Plast., 1971, 7, 202–207.
	- 37 C. Y. Liu and K. Aika, Bull. Chem. Soc. Jpn., 2004, 77, 123–131.
	- 38 G. An, L. Wang, J. Gao and R. Wang, Sci. Rep., 2019, 9, 1563.
	- 39 N. E. Gordina, T. N. Borisova, K. S. Klyagina, I. A. Astrakhantseva, A. A. Ilyin and R. N. Rumyantsev, Membranes, 2022, 12, 147.
	- 40 G. G. Martens, J. W. Thybaut and G. B. Marin, Ind. Eng. Chem. Res., 2001, 40, 1832–1844.
	- 41 J.-B. Lin, T. T. T. Nguyen, R. Vaidhyanathan, J. Burner, J. M. Taylor, H. Durekova, F. Akhtar, R. K. Mah, O. Ghaffari-Nik, S. Marx, N. Fylstra, S. S. Iremonger,
		- K. W. Dawson, P. Sarkar, P. Hovington, A. Rajendran,

T. K. Woo and G. K. H. Shimizu, Science, 2021, 374, 1464– 1469.

- 42 T. Zeng, H. Huang, N. Kobayashi and J. Li, Nat. Resour., 2017, 8, 611–631.
- 43 M. Thommes, K. Kaneko, A. V. Neimark, J. P. Olivier, F. Rodriguez-Reinoso, J. Rouquerol and K. S. W. Sing, Pure Appl. Chem., 2015, 87, 1051–1069.
- 44 F. Akhtar, Q. Liu, N. Hedin and L. Bergström, Energy Environ. Sci., 2012, 5, 7664–7673.
- 45 B. Coughlan and R. G. Shaw, Proc. R. Ir. Acad., Sect. B, 1976, 76, 191–210.
- 46 B. Ilić and S. G. Wettstein, Microporous Mesoporous Mater., 2017, 239, 221–234.
- 47 Z. Cao and F. Akhtar, Adv. Funct. Mater., 2021, 31, 2008505.
- 48 Z. Wang, L. Wang, P. Gao, Y. Yu and R. Wang, Appl. Therm. Eng., 2018, 128, 1382–1390.
- 49 Z. Cao, N. Grimaldos Osorio, X. Cai, P. Feng and F. Akhtar, J. Environ. Chem. Eng., 2020, 8, 103584.
- 50 C. Y. Liu and K.-I. Aika, Res. Chem. Intermed., 2002, 28, 409– 417.
- 51 Z. Cao, K. N. Landström and F. Akhtar, Catalysts, 2020, 10, 1444.
- 52 V. E. Sharonov and Y. I. Aristov, React. Kinet. Catal. Lett., 2005, 85, 183–188.
- 53 Z. Cao, R. A. Laisequilla and F. Akhtar, Energy Proceedings, 2021, 18, 014.