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Cationic Oligomer as an Organic Template for Direct Synthesis of Aluminosilicate ITH Zeolite

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Abstract: Despite great progress in the synthesis of aluminosilicate zeolites, there are still a large number of zeolites that could not be prepared in the aluminosilicate form. One of typical examples is ITH zeolite, where direct synthesis of aluminosilicate ITH is currently still challenging. Herein, for the first time we demonstrate the successful synthesis of aluminosilicate ITH zeolite using a simple cationic oligomer as an organic template. The key to this success is that the cationic oligomer has a strong complexation ability with aluminum species combined with a structural directing ability for the ITH structure similar to that of the conventional organic template. Physicochemical analysis shows that the aluminosilicate ITH zeolite has very high crystallinity, nanosheet-like crystal morphology, large surface area, fully four-coordinated AI species, and abundant acidic

sites. More importantly, methanol-to-propylene (MTP) tests reveal that the AI-ITH zeolite shows much higher selectivity for propylene and longer lifetime than commercial ZSM-5. FCC tests show that Al-ITH zeolite is a good candidate as a shape selective FCC additive for enhancing propylene and butylene selectivity.

Introduction

Zeolites, as a typical family of microporous materials, have shown important roles in the process of oil refining and fine chemicals production because of their uniform channel

distribution, high surface area, and large micropore volume.^[1-22] As an important type of zeolites, germanosilicate-based zeolites provide many new structures, exhibiting excellent performance in various catalytic reactions.^[23-34] For example, BEC zeolite displays good catalytic performance in the isomerization of glucose to fructose;^[24] ECNU-21 zeolite is an efficient shape selective catalyst for ethylene oxide hydration;^[25] ITH zeolite shows excellent performance in the catalytic cracking and methanol-to-propylene (MTP) reaction.^[26-29]

ITH zeolite, firstly discovered by Corma's group,^[30] has a special framework structure (three dimensional 9×10×10membered ring channels; aperture size of 4.0×4.8, 4.8×5.1, and 4.8×5.3 Å). It could be prepared in the form of silicate and borosilicate but it was tough to synthesize the aluminosilicate form due to competitive growth of EUO when aluminum is present in the synthesis gel.^[27,31-33] In order to incorporate aluminum in the structure of ITH zeolite to extend the catalytic applications, the addition of germanium species in the ITH zeolite synthesis was always required.^[26-29] This addition not only greatly increases the synthetic cost, but also significantly reduces the hydrothermal and thermal stabilities of the ITH zeolite.^[34] The drawbacks mentioned above strongly limit its catalytic application. To avoid the use of germanium, a postsynthesis route was developed through alumination of borosilicate ITH zeolite into aluminosilicate ITH zeolite.^[30] Up to now, it was still a challenge to directly synthesize aluminosilicate ITH zeolite.

Recently, it has been reported that many zeolites such as RHO, Beta, and GME could be successfully synthesized using cationic polymers/oligomers as novel organic templates.^[35-37] In these cases, it is worth noting that the zeolite products have more Al species in the framework than those synthesized from conventional organic templates. One possibility is that the cationic polymers/oligomers have a stronger complexation ability with aluminum species than conventional templates. Therefore,

it is suggested that the use of novel cationic oligomer as an efficient organic template might be favourable for direct synthesis of the aluminosilicate ITH zeolite.

In this work, using a simple cationic oligomer as an organic template, the direct synthesis of an aluminosilicate ITH zeolite (AI-ITH) was shown to be successful. The obtained AI-ITH zeolite has a Si/AI ratio as low as 57. More interestingly, the AI-ITH shows excellent hydrothermal stability and good performance in both methanol-to-propylene (MTP) and FCC reactions.

Results and Discussion

In order to search for a suitable cationic oligomer for directing the ITH zeolite structure, theoretical simulations were performed, as presented in Figure 1 and Table S1. Notably, the interaction stability energy between the pure silica ITH zeolite framework and the structure of the cationic oligomer (Figure 1, OSDA1) is very similar to that of the conventional organic template (Figure 1, OSDA2), suggesting the possibility of the cationic oligomer for directing the ITH zeolite. In order to compare the complexation ability of the organic templates with aluminum species, both OSDA1 and OSDA2 solutions were mixed with a certain amount of AIF₃. ICP analysis shows that the concentration of aluminum species in the solution of OSDA1 is much higher than that of the OSDA2, as presented in Table S2. These results indicate that the OSDA1 has indeed stronger complexation ability with aluminum species than the OSDA2. To measure the importance of Al complexation ability for OSDA1, we performed additional experiments for addition of aluminum source (AIF₃) in the final gel (H₂O/SiO₂ at 3) by partial removal of water from the starting gel ($H_2O/SiO_2 > 30$). In the final gel, the



Figure 1 Structures of the cationic oligomer (OSDA1, a) and traditional template (OSDA2, b), and their positions within the all silica ITH zeolite framework.

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complexation ability between OSDA1 with aluminum species should be weak due to shortage of water solution, compared with the starting gel. As a result, the addition of aluminum species in the final gel leads to the formation of amorphous product rather than the AI-ITH zeolite (Figure S1), confirming the importance of the complexation of AI with OSDA1. Therefore, both structural directing ability and strong complexation ability with aluminum species of the OSDA are very important for the successful synthesis of AI-ITH zeolite.

Figure 2A shows X-ray diffraction (XRD) pattern of as-made AI-ITH zeolite from the initial gel with Si/AI ratio at 100, exhibiting a series of characteristic peaks associated with ITH structure (Si/AI ratio of 70), in good agreement with the simulated XRD pattern (Figure S2). The scanning electron microscopy (SEM) images in Figure 2B and Figure S3 show that the as-made AI-ITH zeolite has a nanosheet-like morphology with a thickness of about 30 nm. Figure 2C displays nitrogen sorption isotherm of calcined AI-ITH zeolite, giving BET surface area and micropore volume of about 400 m²/g and 0.15 cm³/g, respectively. In addition, the measured external surface area of 64 m²/g and mesopore volume of 0.22 cm³/g, might be related to the nanosheet-like morphology of AI-ITH zeolite. Furthermore, Si/AI ratio of AI-ITH zeolite is determined to be 70 by inductively coupled plasma (ICP) technique. Figure 2D shows ²⁹Si MAS

NMR spectrum of as-made AI-ITH zeolite, giving four peaks with the chemical shift centered at -116.8, -114.2, -111.3, and -105.4 ppm. The first three peaks are assigned to Si(4Si) species, and the fourth one is attributed to the Si(3Si, OH) and/or Si(3Si, 1Al). Figure 2E exhibits ¹⁹F MAS NMR spectrum of as-made AI-ITH zeolite, displaying two peaks at -35.5 and -63.8 ppm, which are assigned to fluoride species in the double four member rings and the [415262] cage, respectively.[33] Figure 2F gives 27AI MAS NMR spectrum of as-made AI-ITH zeolite, showing a peak at 53.3 ppm associated with four-coordinated aluminum species in the ITH zeolite framework. We cannot observe the peak at round 0 ppm, implying the absence of six-coordinated aluminum species in the AI-ITH zeolite product. Figure 2G shows temperature programmed desorption of ammonia (NH₃-TPD) curve of the H-AI-ITH zeolite, showing two desorption peaks centered at about 185 °C and 390 °C, which are very similar to those of AI-Ge-ITH zeolite synthesized by direct method reported in the literature.^[28] Thermal analysis (TG-DTA, Figure 2H) of as-made AI-ITH zeolite shows a weight loss of 15.69 % in the region of 200-600 °C, which is attributed to the decomposition of OSDA1. ¹³C nuclear magnetic resonance (NMR) spectrum of the liquid phase obtained by dissolving asmade AI-ITH zeolite with a dilute HF solution is close to that of the OSDA1 solution (Figure 2I). In addition, many weak peaks



Figure 2 (A) XRD pattern (wavelength: 1.5406 Å), (B) SEM image, (C) N₂ sorption isotherm, (D) ²⁹Si MAS NMR, (E) ¹⁹F MAS NMR, (F) ²⁷Al MAS NMR, (G) NH₃-TPD curve, (H) TG-DTA curve of Al-ITH zeolite product; and (I) ¹³C NMR of a) template solution and b) template dissolved from Al-ITH sample.

have been observed in the dissolved solution treated from Al-ITH zeolite, which could be related to the Hoffmann elimination of a small amount of OSDA1 at relatively high temperature (Figure S4). Combining with the results of NMR and TG-DTA characterizations, it is concluded that the OSDA1 mainly exists in the micropores of the Al-ITH zeolite.

Figure 3 shows selected-area electron diffraction (SAED) patterns and transmission electron microscopy (TEM) images of as-synthesized AI-ITH zeolite. Figure 3a-3c are collected from a nanosheet-like crystal lying flat on the grid. The indexing of SAED pattern with sharp diffraction spots and comparison of experimental and simulated high-resolution TEM images indicate that the crystal has a preferred orientation along the c axis. Another crystal standing vertically on the grid was aligned to the [100] zone axis with assistance of Zones program (Figure 3d-f).^[38] The TEM image viewed along the *a* axis shows that the thickness of the crystal is around 30 nm, in good agreement with that of SEM images. Both projections along these two zone axes match well with the simulated images using the ITH zeolite framework (Figures S5-S7). These results confirm the successful synthesis of AI-ITH zeolite with ITH structure in the presence of the novel cationic oligomer.

Quantitative differential phase contrast (qDPC) has emerged as an effective approach to study e-beam sensitive porous materials using scanning transmission electron microscopy (STEM) mode.^[39,40] Compared with high-angle annular dark field (HAADF) image, qDPC-STEM shows advantages on visualizing both heavy and light elements at the same time.^[41-46] As a result, it is possible to obtain information on organic templates in the Al-ITH zeolite micropores using qDPC-STEM images. From the structure model of ITH zeolite (Figure S5), straight channels of 10 MR and 9 MR ring would be clearly seen along [010] and [100] zone axes, respectively. Therefore, qDPC-STEM images should be taken along these two directions to observe organic templates. However, Al-ITH zeolite is a nanosheet-like material and [001] zone axis is vertical to the plate, which means that the Al-ITH crystals along [010] or [100] zone axis would be difficult to find otherwise it is too thick to take qDPC-STEM images clearly. Fortunately, we were able to find a crystal with the thin section along [010] zone axis and thus obtain the HAADF-STEM and qDPC-STEM images of AI-ITH zeolite, as shown in Figure 4. Compared with HAADF image (Figure 4a) without the observation of any organic template due to the limitation of HAADF technique, it would be clearly seen that there are some white contrasts in aDPC-STEM image (Figure 4b), which might be related to organic template molecules occluded in the zeolite micropore channels. Combining with the result of theoretical simulation, NMR and TG-DTA characterizations, and gDPC-STEM images, it would further be concluded that the OSDA1 molecule, which is fully positioned in the micropore of aluminosilicate AI-ITH zeolite, is indeed an organic template for directing the formation of aluminosilicate AI-ITH zeolite structure.



Figure 3 SAED patterns and TEM images of AI-ITH zeolite viewed along the [001] (a, b and c) and [100] (d, e and f) zone axes. The insets in (c) and (f) are symmetry-averaged (left) and simulated images (right), respectively. The settings of defocus values and thickness for (c) and (f) are 500 nm & 5.2 nm and -140 nm & 13 nm, respectively.



Figure 4 HAADF-STEM (a) and qDPC-STEM (b) images of as-synthesized AI-ITH zeolite viewed along [010] zone axis. Circles on Figure 4b with the white contrast spots might represent organic templates molecules inside the zeolite channel.

To understand the AI-ITH crystallization, XRD, SEM, and MAS NMR characterization techniques were used. Before the crystallization, only an amorphous phase is observed (Figure S8a and Figure S9a). After 4 days of crystallization, the XRD pattern of the sample shows very weak peaks associated with the AI-ITH zeolite (Figure S8b). At the same time, the AI-ITH zeolite crystals with nanosheet-like morphology can clearly be seen from the SEM image (Figure S9b). Increasing the crystallization time from 5 to 8 days, the XRD peak intensity of the samples gradually increases (Figures S8c-8f). Correspondingly, more AI-ITH zeolite crystals are observed in the SEM images (Figures S9c-9f). After 10 days, Al-ITH zeolite is obtained with high crystallinity (Figure S8g and Figure S9g). The dependence of the AI-ITH zeolite crystallinity on the crystallization time is shown in Figure S10. Figures S11 and S12 show ²⁷Al and ²⁹Si MAS NMR spectra of Al-ITH zeolite products obtained after the different crystallization times. As shown in Figure S11, the products after 4 to 10 days of crystallization have a peak near 60 ppm associated with 4-coordinated aluminum species, while the aluminum species are 6coordinated at the beginning in the crystallization. These results show that 6-coordinated aluminum species have been completely transferred into 4-coordinated species in the initial stage of the crystallization. As shown in Figure S12 and Tables S3-S4, the maximum percentage of Si(3Si) species (-101.9 ppm) appears at 5 days. Correspondingly, the XRD data show the crystallization happening fast at this stage. Possibly, the rearrangement of silicon species with the intermediate of Si(3Si) under assistance of organic templates plays a critical role in the crystallization.

It is well known that high hydrothermal stability is a key factor for many industrial applications of zeolites.^[47,48] Therefore, it is very crucial to investigate the hydrothermal stability of Al-ITH zeolite in this work. After steaming treatment at 800 °C for 5 h with 10% water, the crystallinity, surface area, and pore volume of aged H-Al-ITH zeolite are very close to those of the fresh one (Figure S13, Tables S5-S6). These results demonstrate that the

Al-ITH zeolite has extraordinarily hydrothermal stability, which would be favorable for its industrial application.

Figure 5 and Figure S14 show the catalytic performances in the MTP reaction over AI-ITH-70, AI-ITH-94, SiGeAI-ITH, SiAI(B)-ITH, and ZSM-5 zeolites. Table S7 presents the catalytic data after 2 h time on stream. Obviously, the AI-ITH zeolites combine the advantages of high selectivity for propylene and long catalytic lifetime among the tested zeolite samples. In addition, it is worth noting that the absence of Ge species would stabilize the structure of AI-ITH zeolite against the irreversible hydrothermal effect of the water produced in the MTO reaction. Therefore, the above results are potentially important for selective production of propylene in the industrial applications.



Figure 5 Dependences of methanol conversion and product selectivity on reaction time in MTP reaction over the Al-ITH-70 zeolite at 480 °C. \oplus : $C_2^{=}$ selectivity, \blacktriangle : $C_3^{=}$ selectivity, \bigstar : $C_4^{=}$ selectivity, \bigstar : Conversion of methanol.

In addition, it has been reported that the structural and acidic properties of AI-ITH make it a good alternative to ZSM-5 as shape selective FCC additive for producing propylene.^[27,49] Thus,

the catalytic behaviour of the AI-ITH sample and two ZSM-5 samples have been compared for catalytic cracking of a vacuum gasoil (VGO, see Table S8 for VGO properties) in a microactivity test unit.^[28] The results are shown in Figures S15-S17. The presence of the additives results in all cases in a decrease of the gasoline fraction and an increase of the gases produced due to cracking of the olefins and paraffins present in the gasoline. The drop in gasoline yield is higher for the ZSM-5 additive with the highest Al content and lowest for the H-Al-ITH. Diesel and coke selectivity are not affected. The highlight is the AI-ITH additive increases propylene yield to levels close to those obtained with ZSM-5 (113) but without increasing the yield of propane. Moreover, it also increases the yield to the linear butenes to values comparable to those given by the ZSM-5 zeolites with no increase of the saturated butanes. This results in higher olefin to paraffin ratios in the C₃ and C₄ fractions. When comparing the catalytic behaviour of AI-ITH with that of a AIGe-ITH (Si/Ge=10, (Si+Ge)/Al=50) described previously,[27] it can be seen that the latter, with a higher Al content, produces more gases by gasoline and even diesel cracking (see Figure S18a). However, Al-ITH, although yielding less propene, presents comparable propene/propane ratio (see Figure S18b) and the additional advantage of its higher thermal and hydrothermal stability due to the absence of Ge. Therefore, the above results show that Al-ITH zeolite is indeed a good candidate as an FCC additive.

After the successful synthesis of AI-ITH zeolite with a Si/AI ratio of 70 in the final product (Table S9, Run 1), AI-ITH zeolites with various Si/AI ratios (57, 94, and 120) are also synthesized (Table S9, Run 2-4). All products have a good crystallinity, four-coordinated aluminum species, and crystals with nanosheet-like morphology (Figures S19-S21). The details for all the synthesis process are presented in Table S9.

Conclusion

In summary, the direct synthesis of AI-ITH zeolite with different Si/AI ratios is successful in the presence of a cationic oligomer as an organic template. The key to the successful synthesis is the synergistic feature of the cationic oligomer with both the structural directing ability for crystallization of ITH zeolite and the strong complexation ability of aluminum species. The obtained AI-ITH zeolite has superior hydrothermal stability. Catalytic tests in MTP reaction show that the AI-ITH zeolite is active and selective for the production of propylene together with a long catalytic lifetime. In addition, AI-ITH zeolite is a good candidate as a shape selective FCC additive. The use of the cationic oligomers might open an alternative way in the future to synthesize other zeolites directly that up to now could not yet be crystallized as aluminosilicates.

Experimental Section

As a typical example for the organic template synthesis, 17.231 g of N,N,N',N'-tetramethylhexane-1,6-diamine (0.1 mol) was mixed with 21.591 g of 1,4-dibromobutane (0.1 mol) and 50 mL of ethanol. The mixture was put under reflux for 12 h. Then, the solvent was evaporated, and the white sediment was washed with ether and dried under vacuum. Gel permeation chromatography (GPC) analysis showed that the bromide salt form of the template had a molecular weight between 4291 and 8669 (n=11-22). The bromide form was converted to the hydroxide

form using an anion exchange resin (IRN-78) in water, and the obtained solution was titrated using 0.1 M HCl.

As a typical run for synthesis of Al-ITH zeolite, boehmite (0-24 mg, 0-0.16 mmol Al₂O₃) and 10.67 g of organic template solution (OSDA1, 0.36 mol/L OH) were mixed. Then, 0.935 g of fumed silica (15.58 mmol) was added under stirring. After stirring the mixture for another 6 h, 0.05 g of siliceous ITH zeolite seeds and 0.190 mL of HF (40%, 3.3 mmol) were added. After partial water evaporation from starting gel (H₂O/SiO₂ > 30), the final molar composition of the mixture was 1.0 SiO₂ : 0-0.01 Al₂O₃ : 0.005 OSDA1 : 0.2 HF : 1-3 H₂O. Finally, the mixture was transferred into a Teflon-lined autoclave and heated at 160 °C for 10 days under rotation condition (50 rpm). The solid zeolite product, which was denoted as Al-ITH-x, where x was the Si/Al ratios in the starting gel, was then filtered, washed with deionized water, dried at 100 °C and calcined at 550 °C for 5 h and named as H-Al-ITH-x. After hydrothermal treatment of H-Al-ITH zeolite samples at 800 °C with 10% H₂O for 5 h, the aged zeolite product was obtained.

To evaluate the complexation ability of organic templates with aluminum species, 0.05 g AlF₃ and 10.67 g of template solution (0.36 mol/L OH⁻ in OSDA1 and OSDA2 aqueous solution) were mixed at room temperature. After stirring for 30 min, the Al concentration of the supernatant was analysed using inductively coupled plasma (ICP). Considering the AlF₃ is not soluble in water and does not react with base, if the supernatant has higher Al species concentration, it means the better complexation with Al species.

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Keywords: zeolite • aluminosilicate ITH • cationic oligomer • direct synthesis • MTP reaction

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This work shows a way to directly synthesize aluminosilicate ITH (AI-ITH) zeolite using a cationic oligomer as an organic template, where the strong complexation ability with aluminum species is a key factor for the successful preparation.