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Title: Improvement of low-temperature catalytic activity over hierarchical Fe-Beta catalysts for selective catalytic reduction of  $NO_x$  with  $NH_3$ 

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To appear in:

Chinese Chemical Letters



Please cite this article as: Zhu N, Lian Z, Zhang Y, Shan W, He H, Improvement of low-temperature catalytic activity over hierarchical Fe-Beta catalysts for selective catalytic reduction of  $NO_x$  with  $NH_3$ , *Chinese Chemical Letters* (2019), https://doi.org/10.1016/j.cclet.2019.03.011

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#### Communication

### Improvement of low-temperature catalytic activity over hierarchical Fe-Beta catalysts for selective catalytic reduction of NO<sub>x</sub> with NH<sub>3</sub>

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### **Graphical Abstract**



Hierarchical Fe-Beta exhibited higher low-temperature NH3-SCR activity than conventional Fe-Beta, due to more active sites and better dispersion of Fe species.

#### ARTICLE INFO

Article history: Received 27 January 2019 Received in revised form 1 March 2019 Accepted 5 March 2019 Available online

*Keywords:* Hierarchical structure Fe-Beta NH<sub>3</sub>-SCR Low-temperature activity Kinetics

#### ABSTRACT

Hierarchical Fe-Beta obtained by hydrothermal synthesis exhibited higher low-temperature NH<sub>3</sub>-SCR activity than conventional Fe-Beta. In order to identify the main factors leading to the difference in catalytic activity, we investigated the pore structure, acidity and Fe sites of the hierarchical Fe-Beta and conventional Fe-Beta. The enhanced activity of hierarchical Fe-Beta was mainly due to the increase of the quantity of active Fe species. NH<sub>3</sub>-TPD and DRIFTS results of NH<sub>3</sub> adsorption clearly verified that hierarchical Fe-Beta had more Lewis acid sites, which is beneficial to the adsorption and activation of NH<sub>3</sub>. The H<sub>2</sub>-TPR, UV-vis DRS, and EPR results confirmed that the hierarchical structure introduced more structural defects as ion-exchange sites. Based on the analysis of kinetics experiments and the above-mentioned characterizations, it can be concluded that the improvement of NH<sub>3</sub>-SCR activity was not due to an intrinsic effect of the specific structural characteristics, but was related to more Fe active sites and better dispersion of Fe species in the hierarchical Fe-Beta.

Iron-based zeolites (mostly Fe-ZSM-5 and Fe-Beta) have been extensively investigated because of their high activity and thermal stability for the selective catalytic reduction of  $NO_x$  with NH<sub>3</sub> (NH<sub>3</sub>-SCR) [1-6]. Compared with ZSM-5, Beta zeolite is an attractive host for iron and Fe-Beta zeolite has higher activity than Fe-ZSM-5 in some catalytic reactions [7, 8]. Many studies focused on Fe-Beta catalysts for N<sub>2</sub>O decomposition [8-11] and NH<sub>3</sub>-SCR, especially on the active sites and hydrothermal stability [1, 12, 13]. Previous studies suggested that the lowtemperature NH<sub>3</sub>-SCR activity of Fe-Beta could be enhanced by bimetallic doping (such as Cu [14], Mn [15], Ce [16, 17]) and high-temperature H<sub>2</sub>-pretreatment [2, 6].

Hierarchical zeolites contain both micro and meso-/macrospores, which can eliminate diffusion limitations and promote the accessibility of reactants to the active sites [18-21]. The predominant approach for the synthesis of hierarchical structure zeolites is demetallation (desiliconization and dealumination) and recrystallization with the surfactants [19, 22]. Previous studies suggested that NH<sub>3</sub>-SCR activity was enhanced by introducing hierarchical structure to Fe-ZSM-5 [23, 24]. However, there are few open literatures about hierarchical Fe-Beta for NH<sub>3</sub>-SCR.

In this work, hierarchical Fe-Beta zeolites catalysts were prepared by hydrothermal synthesis and evaluated for  $NH_3$ -SCR. The preparation procedure and activity tests are shown in the Support information. It was found that the low-temperature activity of hierarchical Fe-Beta was higher than that of Fe-Beta. The structure and activity relationship was discussed in detail.

As shown in Fig. 1, it was clear that the Beta was active for the SCR reaction in a narrow temperature range of 375-500 °C. Acidity may account for its catalytic performance [25, 26]. Hierarchical Beta zeolite showed obviously improved NO conversions in the whole temperature range. This result indicated that hierarchical Beta zeolite may have more acid sites than Beta, which will be verified in the subsequent NH<sub>3</sub>-TPD and NH<sub>3</sub> adsorption sections. After the doping of Fe, the Fe-Beta and hierarchical Fe-Beta indicated highly improved SCR activity as compared with Beta and hierarchical Beta in the whole temperature of 150-500 °C. Hierarchical Fe-Beta showed enhanced low-temperature SCR activity compared with Fe-Beta obtained using the same ion-exchanged procedure. All of these

catalysts presented excellent  $N_2$  selectivity. The relationship of structure and activity was discussed in detail as follows.



**Fig. 1.** NO<sub>x</sub> conversion and N<sub>2</sub> selectivity (inserted) results of Fe-Beta and Beta catalysts. Reaction conditions:  $[NO] = [NH_3] = 500$ ppm,  $[O_2] = 5$  vol%, N<sub>2</sub> balance and GHSV = 200,000 h<sup>-1</sup>.

The XRD patterns (Fig. S1 in Supporting information) show that all samples are phase-pure zeolites with the BEA framework structure. The peak intensity of Fe-Beta and hierarchical Fe-Beta declined, which may be due to the collapse or blocking of the partial microspore of zeolite. FeO<sub>x</sub> species were not detected, which may be due to the Fe species existing as amorphous or high dispersion on the zeolite support [27, 28].

The textural properties in Table S1 and Fig. S2 (Supporting information) show that the total volume, external surface area and BET surface area of hierarchical Beta are larger than those of Beta, indicating that mesopores and/or macropores are introduced into Beta to form hierarchical structure [19, 20]. After doping with Fe, the BET surface area of both Fe-Beta and hierarchical Fe-Beta decreased, indicating that the partial microspore may be blocked, in agreement with the XRD results. ICP analysis revealed that the Fe content of Fe-Beta and hierarchical Fe-Beta was 2.82% and 3.71%, respectively, suggesting that hierarchical Fe-Beta could achieve a large amount of iron at the exchange sites under the identical ion-exchange procedure.

Comparing TEM images of hierarchical Fe-Beta (Fig. S3, c and d) with those of Fe-Beta (Fig. S3, a and b), the smaller crystal size and some hollow Beta structure were obtained during the hydrothermal synthesis, indicating the formation of micromacroporous structure. This result is in accordance with the BET

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results. In Fig. S3c, the hollow zeolite was made of a polycrystalline micro-mesoporous shell and macropore core. Previous studies suggested that hollow Beta single crystals were much more difficult to prepare [18]. The absence of Al zoning prohibited the use of the dissolution-recrystallization strategy employing in the synthesis of hollow Beta zeolite single crystals [29].

UV-vis DR spectroscopy is an effective and common method for determining the nature and distribution of Fe species in Fezeolites. Fig. S4a (Supporting information) shows the UV-vis DR spectra of the Fe-Beta samples. The spectra were deconvoluted into sub-sands, which can be attributed to specific Fe species in the Fe-zeolites. The bands at 211 and 277 nm were respectively assigned to isolated Fe3+ ions in tetrahedral and octahedral coordination environments [30]. The bands at 300-400 nm indicated the presence of small oligomers  $Fe_x^{3+}O_y$  in the zeolite channels [8, 28], and the adsorption bands above 400 nm were attributed to the presence of bulky Fe<sub>2</sub>O<sub>3</sub> nanoparticles on the external surface of the zeolite [31, 32]. Fig. S4b (Supporting information) shows the UV-vis DR spectra of hierarchical Fe-Beta samples, the peak attributions are similar to Fe-Beta samples. The amount of Fe species was estimated semiquantitatively based on the relative intensities of these sub-bands derived from UV-vis DR spectra and the total Fe loading in the zeolite. The data in Table S2 in Supporting information shows that isolated Fe<sup>3+</sup> ions are the predominant species in Fe-Beta and hierarchical Fe-Beta catalysts. Previous studies suggested isolated  $Fe^{3\scriptscriptstyle +}$  species are active sites for  $NH_3\mbox{-}SCR$  on Fe-Beta catalysts [4,5,25,33,34]. When the Fe loadings were considered, the hierarchical Fe-Beta had the larger content of isolated Fe<sup>3+</sup> species than Fe-Beta, resulted in the enhanced NH<sub>3</sub>-SCR performance.

EPR spectroscopy was further employed to identify the nature of different Fe species, the corresponding results are illustrated in Fig. S5 (Supporting information). Signals at g=2.0, g=4.3 and g=8.8 were observed for Fe-Beta and hierarchical Fe-Beta. The signals at g=4.3, g=8.8 and g=2 were generally attributed to isolated Fe species in tetrahedral and distorted tetrahedral coordination [30, 35], isolated Fe species with a distorted octahedral environment [8, 13] and isolated Fe species in a high symmetry octahedral coordination or oligonuclear Fe<sub>x</sub>O<sub>y</sub> clusters [8, 28], respectively. Overall, the EPR signal intensity of hierarchical Fe-Beta was higher than Fe-Beta, illustrating the amount of isolated Fe species on hierarchical Fe-Beta was higher than that of Fe-Beta, in agreement with UV-vis DRS results.



Fig. 2. H<sub>2</sub>-TPR results of the different Beta samples.

In order to investigate the redox performance of the samples, the  $H_2$ -TPR experiments were conducted and the results are

shown in Fig. 2. There is no clear  $H_2$  consumption signal for the Beta and hierarchical Beta. Two reduction peaks appeared on the Fe-Beta and hierarchical Fe-Beta. The overlapped reduction peaks centered at 354 °C (369 °C) and 441 °C (499 °C) were attributed to the reduction of isolated Fe<sup>3+</sup> to Fe<sup>2+</sup> [36] and oligomeric iron oxo species or tiny Fe<sub>2</sub>O<sub>3</sub> into Fe<sub>3</sub>O<sub>4</sub>, respectively [37-39]. The reduction temperature of Fe species over hierarchical Fe-Beta was lower than Fe-Beta, which may be due to the better dispersion of Fe species. The amount of H<sub>2</sub> consumption of hierarchical Fe-Beta catalyst had more Fe active sites. The better redox performance resulted in the enhanced NH<sub>3</sub>-SCR activity on the hierarchical Fe-Beta.

In view of the analysis of UV-vis DRS, EPR and  $H_2$ -TPR results above, we have confirmed that more Fe active species existed on the hierarchical Fe-Beta and dispersed better than those on Fe-Beta, which may be due to the specific hierarchical structure.

To identify the strength of the acid sites of hierarchical Fe-Beta, the NH<sub>3</sub>-TPD experiments were carried out and the results are shown in Fig. 3. Beta zeolite showed NH3 desorption peaks at the range of 100-500 °C, assigning to the desorption of ammonia species from weak Lewis acid sites (167 °C), strong Lewis acid sites (262 °C and 330 °C) and strong Brønsted acid sites (373 °C) [25, 37, 40]. The intensity of the desorption peak at 262 °C was much stronger than that of peak at 373 °C, suggesting that Beta zeolite had more Lewis acid sites because of its structural defects caused by the unsaturated bonding of silicon [41, 42]. When forming the hierarchical structure, the peaks shifted to the hightemperature, demonstrating the formation of the stronger acid sites. The peak at 89 °C on hierarchical Beta may be attributed to physisorbed NH<sub>3</sub> [39, 43]. The number of acid sites of hierarchical Beta also increased, which resulted in the enhanced SCR activity, compared with Beta zeolite (Fig. 1). When Fe was introduced into Beta, the NH3 storage capacity decreased, and the desorption peak shifted to high-temperature. However, when Fe was exchanged to hierarchical Beta, the NH<sub>3</sub> desorption peak shifted to the low-temperature, to form more weak Lewis acid sites (167 °C), which is beneficial to the adsorption and activation of NH<sub>3</sub>.



Fig. 3. NH<sub>3</sub>-TPD profiles of different Beta zeolites.

In order to understand the adsorption performance and reaction mechanism, in situ DRIFTS experiments were carried out for Fe-Beta and hierarchical Fe-Beta, as shown in Fig. S6 in Supporting information. After the adsorption of  $NH_3$  and  $N_2$  purge, the  $NH_3$  adsorbed peaks appeared on both of the catalysts. The negative bands at 3773, 3721, 3604 cm<sup>-1</sup> was assigned to Al-OH groups, terminal Si-OH, isolated bridging acidic hydroxyl stretching

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vibrations (Si(OH)Al), respectively [44-46]. The negative band at 3675 cm<sup>-1</sup> only appeared on hierarchical Fe-Beta, which was attributed to hydroxyls bonded on extra-framework Al species [47], due to hierarchical structure. This result indicated that hierarchical structure introduced more exchanged sites and acid sites, which led to the better SCR activity. In the N-H stretching regions (3100-3500 cm<sup>-1</sup>), the bands corresponding to the asymmetric stretching (3343cm<sup>-1</sup>) and symmetric stretching (3273 and 3192 cm<sup>-1</sup>) of coordinated NH<sub>3</sub> on Lewis acid sites were observed, respectively [48]. The bands at 1478 cm<sup>-1</sup> and 1163 cm<sup>-1</sup> were attributed to the NH4<sup>+</sup> ions adsorbed on the Brønsted acid sites and coordinated NH<sub>3</sub> adsorbed on the Lewis acid sites, respectively [48-50]. The band at 896 cm<sup>-1</sup> could be assigned to T-O-T framework vibrations by Fe species [7]. The peak intensity at 896 cm<sup>-1</sup> of hierarchical Fe-Beta was higher than that of Fe-Beta, illustrating more Fe was incorporated into hierarchical Fe-Beta, in agreement with the ICP, EPR and H<sub>2</sub>-TPR results. Overall, the intensity of NH<sub>3</sub> adsorption peaks for hierarchical Fe-Beta was higher than those for Fe-Beta, suggesting more acid sites existing on hierarchical Fe-Beta than those on Fe-Beta, in accordance with NH<sub>3</sub>-TPD results.

The NO adsorbed spectra were shown in Fig. S6b. No peak for NO<sub>x</sub> adsorbed species was clearly observed, suggesting that Fe-Beta and hierarchical Fe-Beta did not adsorb NO<sub>x</sub> species. The NO<sub>x</sub> adsorbed species were not detected on the Fe-Beta and hierarchical Fe-Beta, but the excellent NO<sub>x</sub> conversion could be obtained, illustrating that the NH<sub>3</sub>-SCR reaction may mainly carried out through Eley-Rideal (E-R) pathway with gaseous NO reacting with adsorbed NH<sub>3</sub> species on both of the catalysts.



Fig. 4. Arrhenius plots of the SCR reaction rates over Fe-Beta and hierarchical Fe-Beta in the feed gas of 500 ppm NO/ 500 ppm NH<sub>3</sub>/ 5.0% vol. % O<sub>2</sub>/ N<sub>2</sub> (500 mL/min, 40 mg catalysts, GHSV: 400,000 h<sup>-1</sup>).

In order to obtain further understanding of the intrinsic effect of hierarchical structure on the NH<sub>3</sub>-SCR reaction, the kinetic experiments were performed on the Fe-Beta and hierarchical Fe-Beta, as shown in Fig. 4. The apparent activation energy obtained from the Arrhenius plots for Fe-Beta and hierarchical Fe-Beta was 56.9 and 56.2 kJ/mol, respectively, in agreement with those for Fe-zeolite reported in the previous literature (Ea = 40~54 kJ/mol) [31, 51, 52]. The two catalysts had similar activation energy, indicating an identical rate controlling mechanism [13]. Since there were no mass transfer related limitations, hierarchical Fe-Beta catalyst had better SCR activity, demonstrating the higher activity was mainly caused by higher amount of Fe active sites [53], in accordance with the H<sub>2</sub>-TPR and UV-vis DRS results.

In summary, hierarchical Beta was prepared by hydrothermal synthesis. The BET and TEM results confirmed that the

meso/macrospores were introduced to Beta zeolite. Hierarchical Fe-Beta exhibited better low-temperature activity than Fe-Beta. The results of UV-vis DRS, EPR and H<sub>2</sub>-TPR indicated that isolated Fe<sup>3+</sup>, oligomers and Fe<sub>2</sub>O<sub>3</sub> coexisted on Fe-Beta and hierarchical Fe-Beta. The enhanced activity of the hierarchical Fe-Beta was related to the prominent presence of highly dispersed Fe active species, which may be because that the hierarchical structure introduced more defects to provide more ion-exchange sites for Fe species. The kinetic experiments demonstrated that the activate energy of Fe-Beta and hierarchical Fe-Beta was 56.2 and 56.9 kJ/mol, respectively, suggesting an identical rate controlling mechanism. The NH<sub>3</sub>-TPD and DRIFTS results of NH<sub>3</sub> adsorption indicated that the hierarchical Fe-Beta had more acid sites for the adsorption and activation of NH<sub>3</sub>. The adsorption of NOx was not clearly observed on Fe-Beta or hierarchical Fe-Beta, indicating that NH3-SCR process of both of the two catalysts mainly followed the Eley-Rideal (E-R) reaction mechanism.

#### Acknowledgments

We gratefully acknowledge the financial supports from the National Natural Science Foundation of China (Nos. 51822811, 21637005), the National Key R&D Program of China (Nos. 2017YFC0212502, 2017YFC0211101), and the Young Talent Project of the Center for Excellence in Regional Atmospheric Environment, CAS (No. CERAE201806).

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