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The promotion effect of manganese on Cu/SAPO for selective catalytic reduction of NO_x with NH₃

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The activity and hydrothermal stability of Cu/SAPO and xMn–2Cu/SAPO for low-temperature selective catalytic reduction of NO_x with ammonia were investigated. An ion-exchanged method was employed to synthesize xMn–2Cu/SAPO, which was characterized by N₂ adsorption, ICP-AES, X-ray diffraction (XRD), NH₃-temperature programmed desorption (NH₃-TPD), NO oxidation, X-ray photoelectron spectrum (XPS), UV-vis, H₂-temperature programmed reduction (H₂-TPR) and diffuse reflectance infrared Fourier transform spectra (DRIFTS). 2Mn–2Cu/SAPO and 4Mn–2Cu/SAPO showed the best SCR activity, in that at 150 °C NO conversion reached 76% and N₂ selectivity was above 95% for the samples. NO oxidation results showed that the 2Mn–2Cu/SAPO had the best NO oxidation activity and the BET surface area decreased as manganese loading increased. XRD results showed that the metal species was well dispersed. NH₃-TPD showed that the acid sites have no significant influence on the SCR activity of xMn–2Cu/SAPO. H₂-TPR patterns showed good redox capacity for xMn–2Cu/SAPO. UV-vis and H₂-TPR showed that the ratio of Mn⁴⁺ to Mn³⁺ increased as manganese loading increased. XPS spectra showed a significant amount of Mn³⁺ and Mn⁴⁺ species on the surface and addition of manganese increased the ratio of Cu²⁺. The promotion effect of manganese to 2Cu/SAPO comes from the generation of Mn³⁺ and Mn⁴⁺ species. Deduced from the DRIFTS spectra, the Elay–Rideal mechanism was effective on 4Mn–2Cu/SAPO.

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1 Introduction

NO_x consists of NO, NO₂, N₂O and their related derivatives, which can lead to numerous environmental and health hazards.¹ By forming nitric acid, NO_x can lead to the generation of acid rain; by forming ammonium nitrate, it can accelerate the generation of PM2.5; by reacting with hydrocarbons under sunlight, it can cause the generation of photo chemical smog and ozone; by infiltrating into lungs, it can result in respiratory morbidity, like impaired host defense and lung inflammation. Thus, legislations on NO_x emission are now becoming more and more stringent. In China, according to the emission standard of air pollutants for thermal power plants (GB 13223-2011), the NO_x emission limit is 100 mg m⁻³ for new natural gas-fired industrial boilers and all oil and coal-fired industrial boilers. For compression ignition and gas fueled positive ignition engines of vehicles, the NO_x emission limit in Chinese National 5 standards is 57% of that in Chinese National 4 standards and 40% of that in Chinese National 3 standards.

The treatment of environmentally harmful NO_x compounds emitted from mobile or stationary sources remains

a challenging task, especially for industrial applications such as cement plants or iron and steel plants, because the suitable process temperature for NO_x abatement is usually around 150 °C, under which commercial SCR catalysts V₂O₅–MoO₃ (WO₃)/TiO₂ cannot work well.

Manganese oxides were reported by Smirniotis *et al.* as potential catalysts for low-temperature NH₃-SCR reactions.^{2–5} They studied the promoted manganese oxides supported on TiO₂ systematically^{3–5} and found that the surface Mn⁴⁺ species was reported to be highly active for the SCR of NO reaction with ammonia at low temperatures.

Recently, copper modified zeolites with a CHA structure attracted much attention owing to their excellent activity, N₂ selectivity and hydrothermal stability for low temperature NH₃-SCR reaction.^{6–8}

However for the Cu/SAPO-34 catalyst, the oxygen activation of transiently formed Cu pairs [Cu^I(NH₃)₂] to [(NH₃)₂Cu^{II}–O₂–Cu^{II}(NH₃)₂] was rate-limiting in the catalytic cycle.⁸ Feng Gao⁹ also thought that the oxidation reaction of Cu⁺ to Cu²⁺ was the rate-determine step, so tuning the redox properties of the active site by introducing a second cation might be a promising approach. Li *et al.*¹⁰ and Chen *et al.*¹¹ modified the Cu/SAPO catalyst with CeO_x for SCR reaction; they found that the introduction of CeO_x increased the SCR activity at high temperatures and cerium helped increase the amount of isolated Cu²⁺ ions.

Leistner *et al.*¹² found that Cu in Cu/SAPO-34 was more easily reduced compared to Cu/SSZ-13, which could facilitate the

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redox processes and increase the SCR activity. Ma *et al.*¹³ also found that Cu-SAPO-34 showed higher DeNO_x catalytic activity than Cu-SSZ-13. The usage of SAPO-34 in low-temperature SCR might be favored in low temperature SCR reactions. When manganese is introduced into the Cu/SAPO system, it not only promotes the oxidation reaction of Cu⁺, but also acts as an active site for low temperature SCR reaction. Thus the manganese modified Cu/SAPO-34 catalyst would probably show better SCR activity.

In this work, the treatment of exhaust from a gas-fired boiler which contains a high concentration of H₂O and a very low concentration of SO₂ was focused upon. *x*Mn-Cu/SAPO catalysts with different manganese loadings were prepared and their performance in SCR reaction was discussed. Various characterization methods (XRD, XPS, H₂-TPR, UV-vis) were used to probe the promotion effect of manganese on Cu/SAPO for low temperature SCR reaction.

2 Experimental

2.1 Catalyst preparation

*x*Mn-2Cu/SAPO catalysts were prepared by an ion-exchange method. The alkali metals in commercial Na/SAPO-34 powder (Jiangsu XFNANO) would decrease SCR activities, so it was transformed to H⁺/SAPO as follows. Na/SAPO-34 powder was ion exchanged using 11% wt NH₄NO₃ (Aladdin, >98.5%) solution whose pH value was adjusted to 3.0–4.0 by 2 M ammonium hydroxide (Aladdin, 25–28%) solution at 80 °C for 4 h. It was then filtered and washed with distilled water three times. Finally, it was dried at 110 °C for 16 h and calcined at 550 °C for 3 h.

*x*Mn-2Cu/SAPO (*x* = 0, 1, 2, 4, 8) was prepared in two steps. Firstly, H⁺/SAPO was mixed with Cu(NO₃)₂ (across, >95%) solution at 80 °C for 6 h under vigorous stirring. Then it was dried and calcined at 550 °C for 3 h. In the second step, it was mixed with Mn(NO₃)₂ (across, >95%) solution at 80 °C for 6 h under vigorous stirring, and at last it was dried and calcined at 550 °C for 3 h.

2.2 SCR activity measurements

Catalytic activity evaluation was carried out using a flow-through powder reactor system equipped with a Fourier transform infrared (FT-IR) spectrometer (THERMO SCIENTIFIC IGS). In order to prevent condensation along upstream tubing, all the gas lines were heated and maintained at 120 °C. The gas mixture was 500 ppm NO + 500 ppm NH₃ + 3% vol O₂ balanced with N₂. The gas hourly space velocity (GHSV) was 65 000 h⁻¹ for the standard SCR reaction. Prior to each activity measurement, the catalysts were pretreated at 500 °C for 30 min with 21% vol O₂/N₂ flow. Catalytic activities were measured in the temperature range of 120 to 210 °C. The typical time to achieve the steady state was 2.5 h. NO conversion, promotion effect, and (*E*_p) and N₂ selectivity were calculated at steady state using the equation below:

$$X_{\text{NO}} = \frac{C_{\text{NO}_{\text{x,in}}} - C_{\text{NO}_{\text{x,out}}}}{C_{\text{NO}_{\text{in}}}} \times 100\% \quad (1)$$

$$E_p = (X_{x\text{Mn-2Cu/SAPO}} - X_{2\text{Cu/SAPO}}) \times 100\% \quad (2)$$

$$\text{N}_2 \text{ selectivity} = \left(1 - \frac{2[\text{N}_2\text{O}]_{\text{outlet}}}{[\text{NO}_x]_{\text{inlet}} - [\text{NO}_x]_{\text{outlet}}} \right) \times 100\% \quad (3)$$

where *C*_{NO_x in} and *C*_{NO_x out} represent the inlet and outlet NO_x concentration. *X*_{xMn-2Cu/SAPO}, *X*_{2Cu/SAPO} and *X*_{xMn/SAPO} respectively represent the NO_x conversion of *x*Mn-2Cu/SAPO, 2Cu/SAPO and *x*Mn/SAPO at 210 °C, and [N₂O]_{outlet} represents the outlet N₂O concentration.

2.3 Characterization

X-ray diffraction patterns were collected on a bruker D8 Advance X-ray diffractometer with a Ni-Filtered Cu K α with a step size of 0.02 in the 2 θ range of 5° to 40°.

Ammonia temperature programmed desorption (NH₃-TPD) experiments were carried out as follows. 150 mg sample was first pretreated in 21% vol O₂/N₂ at 500 °C for 30 min and then cooled to 100 °C in N₂ and a total flow of 100 ml min⁻¹ containing 2500 ppm NH₃ in N₂ was injected into the reactor for 2 hours to achieve a steady state. Once the catalyst was saturated, NH₃ was switched off and the catalyst was swept by N₂ overnight. Finally, the catalyst was heated in N₂ at a temperature ramp to 700 °C with a heating rate of 10 °C min⁻¹.

NO oxidation experiments were carried out as follows. 150 mg sample was first pretreated in 21% vol O₂/N₂ at 500 °C for 30 min and then cooled to 210 °C in N₂. Then a total flow of 100 ml min⁻¹ containing 500 ppm NO and 3% vol O₂ was injected into the reactor for 30 min to achieve the steady state.

The contents of the elements were determined by ion coupled plasma (ICP) optical emission spectroscopy (Thermo IRIS Intrepid II) after microwave digestion.

The BET surface area, pore volume and pore size of the catalyst samples were measured by N₂ adsorption using the MICROMERITICS ASAP 2020 surface area and porosity analyzer.

Diffuse reflectance UV-vis spectra were recorded in the range of 200–800 nm against a BaSO₄ as a reference standard on a HitachiU-3900 UV-vis spectrophotometer equipped with an integration sphere.

H₂-TPR experiments were performed using 20 mg sample as follows. The catalysts were first pretreated at 500 °C for 30 min in a highly pure O₂ (40 ml min⁻¹) stream. Then the furnace temperature was decreased to room temperature, and feed containing 5% vol H₂ in N₂ was fed at a flow rate of 40 ml min⁻¹. H₂-TPR runs were performed as the temperature increased from room temperature to 800 °C at a linear heating rate of 10 °C min⁻¹ and then the temperature was kept constant for 30 min at 800 °C to ensure complete reduction. Hydrogen was measured by TCD.

X-ray photoelectron spectroscopy (XPS) analyses were performed using PHI quantera SXM Scanning ESCA Microprobe (Physical Electronics) with a hemispherical detector operating at a constant pass energy (PE = 55 eV). An X-ray source at 210 W (*I* = 15 mA, *U* = 14 kV) and Al K α radiation (1486.6 eV) were used. Intensities were estimated from the integration of each peak, after smoothening, subtracting the L-shaped background,



and fitting the experimental curve to a combination of Lorentzian and Gaussian lines of variable proportions. All binding energies were referenced to the C 1s line at 284.8 eV.

Diffuse reflectance infrared Fourier transform spectra (DRIFTS) were measured on an FT-IR spectrometer (Thermo Nicolet NEXUS870) with an MCT detector and high temperature reaction chamber (Harrick Scientific Praying Mantis) with ZnSe Windows, which was connected to a gas-dosing system. The powder sample was placed in a sample cup and heated by a cartridge heater underneath the sample. The temperature was adjusted using a K-type thermocouple connected to a Harrick temperature controller. Before each measurement, oxidation pretreatments were executed at 500 °C for 1 h. Background spectra were collected before adsorption for 32 scans with a resolution of 4 cm⁻¹ in N₂. DRIFTS spectra were recorded in the range of 4000–650 cm⁻¹ for 32 scans with a resolution of 4 cm⁻¹.

3 Results and discussion

3.1 Low temperature NH₃-SCR activity and N₂ selectivity

Fig. 1a shows the NO conversion of xMn–2Cu/SAPO. There was not much difference in the catalytic activity between 2Cu/SAPO and 1Mn–2Cu/SAPO, 2Mn–2Cu/SAPO and 4Mn–2Cu/SAPO. The activity of 8Mn–2Cu/SAPO was slightly lower than the activity of 2Mn–2Cu/SAPO. As manganese loading increased from 1% to 2%, the activity increased obviously and the SCR activity of 2Mn–2Cu/SAPO, 4Mn–2Cu/SAPO and 8Mn–2Cu/SAPO was much higher than that of 2Cu/SAPO and 1Mn–2Cu/SAPO.

For 2Mn–2Cu/SAPO and 4Mn–2Cu/SAPO, the conversion of NO at 120 °C was about 60%; as the temperature increased to 210 °C, the conversion reached 95%.

Fig. 1b shows the N₂ selectivity of xMn–2Cu/SAPO. Good N₂ selectivity was observed, being above 95% for all catalysts. Other researchers^{10,11} also found good N₂ selectivity for the supported SAPO-34 catalysts.

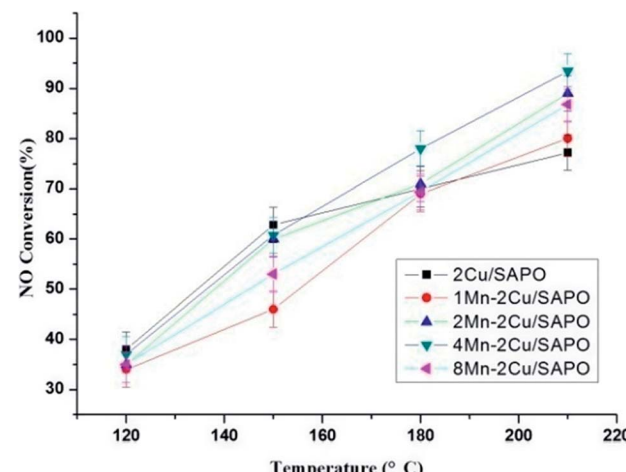


Fig. 2 Water resistance test of xMn–2Cu/SAPO for low-temperature SCR. Conditions: 500 ppm NO + 500 ppm NH₃ + 16.3% H₂O + 3% vol O₂ + N₂ balance, GHSV = 65 000 h⁻¹.

Compared with xMn/SAPO ($x = 1, 2, 4, 8$), the previously prepared,¹⁴ xMn–2Cu/SAPO showed better SCR activities, especially for 2Mn–2Cu/SAPO, indicating that the interactions between copper and manganese might increase its SCR activity.

3.2 Water resistance test

Fig. 2 shows the water resistance test of xMn–2Cu/SAPO. The poison effect of water might come from the reversible adsorption of H₂O¹⁵ or irreversible dealumination process of zeolite.¹⁶

In Fig. 2, all catalysts showed strong resistance to water at high temperature. It could be caused by the lower adsorption capability of water at high temperature, since SAPO-34 was highly hydrothermally stable¹⁶ and the poison effect might originate from the competitive adsorption of water with ammonia.

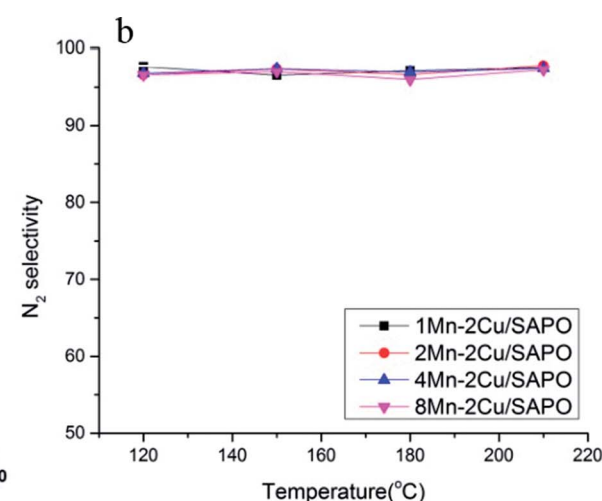
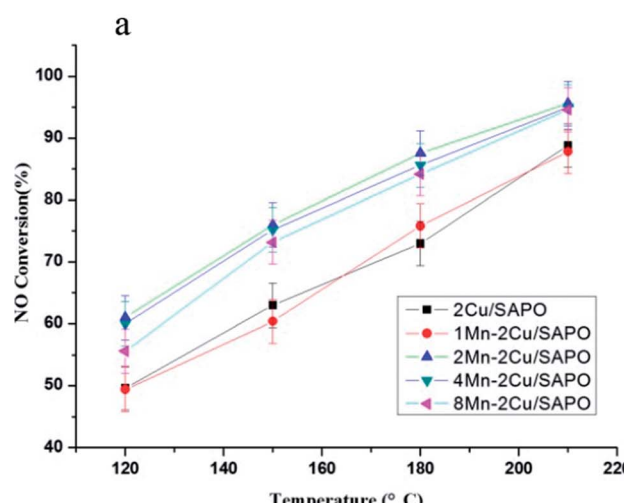


Fig. 1 (a) Catalytic activity and (b) N₂ selectivity of xMn–2Cu/SAPO. Conditions: 500 ppm NO + 500 ppm NH₃ + 3% vol O₂ + N₂ balance, GHSV = 65 000 h⁻¹.



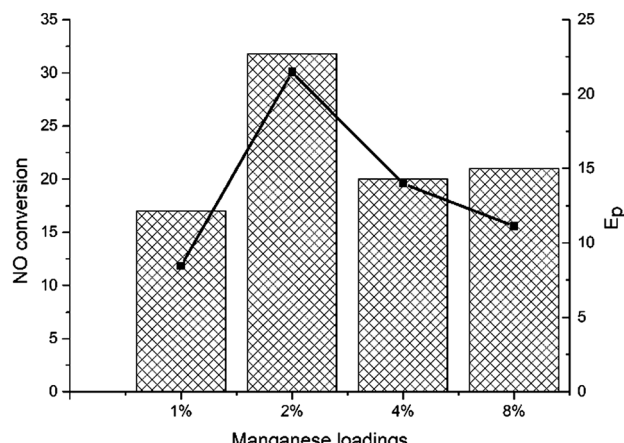


Fig. 3 NO oxidation activity and promotion effect of manganese.

3.3 NO oxidation activity and promotion effect

Fig. 3 showed the NO oxidation activity of x Mn–2Cu/SAPO and the promotion effect. 2Mn–2Cu/SAPO had the best NO oxidation activity, indicating the interaction between copper and manganese elements, which accelerated the NO oxidation process. Good oxidation activity would certainly help the oxidation of Cu^+ to Cu^{2+} in SCR reaction. The E_p profile was almost the same as the NO conversion profile, indicating that the increase in NO oxidation activity would promote the low-temperature SCR activity. Compared with the x Mn/SAPO ($x = 1, 2, 4, 8$) previously prepared,¹⁴ the NO oxidation activities of x Mn–2Cu/SAPO were much higher, which could be one reason for the increased SCR activity.

3.4 Physicochemical properties

As showed in Table 1, the BET surface area and micro pore volume decreased gradually as manganese loading increased; the surface areas of 2Mn–2Cu/SAPO, 4Mn–2Cu/SAPO and 8Mn–2Cu/SAPO were similar. x Mn/SAPO ($x = 1, 2, 4, 8$) previously prepared¹⁴ showed the same trend: the surface area and pore volume decreased as manganese loading increased.

The manganese loading, copper loading and Mn/Cu molar ratio from ICP result were almost the same as the set values. Meanwhile the Mn/Cu molar ratio from the XPS results was quite different from that of ICP results. Since XPS and ICP reflect the elemental information of the surface phase and the

bulk phase respectively, the difference might come from the enrichment of manganese on surface.

3.5 XRD

XRD patterns of x Mn–2Cu/SAPO are depicted in Fig. 4. The diffraction peaks of chabazite phase with the space group of $R\bar{3}m$ were identified, indicating that the crystalline structure of SAPO-34 remained unchanged after catalyst preparation. When manganese loading increased from 0 to 4%, only the chabazite phase was detectable, indicating that the manganese species were well dispersed. As manganese loading reached 8%, the diffraction peaks of the Mn_2O_3 phase appeared, indicating the aggregation of manganese species and decreased manganese dispersity. On x Mn/SAPO ($x = 1, 2, 4, 8$), which was previously prepared,¹⁴ there too were no peaks attributed to manganese oxides until manganese loading increased to 8%.

3.6 NH₃-TPD

Fig. 5 showed the effluent NH₃ profiles during the NH₃-TPD process. The amount of NH₃ desorption was closely related to the amount of acid sites. As expected, when manganese loading increased, the amount of NH₃ desorption decreased, which was caused by the replacement of H⁺ in the framework of SAPO by Mn⁴⁺ in the ion exchange process.

In Fig. 5 there are three NH₃ desorption peaks, *i.e.* a low temperature peak at 250 °C, a middle temperature peak at 400 °C and a high temperature peak at 500 °C, which could be assigned to the weak acid site, the middle strong acid site and the strong acid site, respectively.¹⁷

The shapes of 1Mn–2Cu/SAPO, 2Mn–2Cu/SAPO and 4Mn–2Cu/SAPO were similar while the activity differed a lot, indicating that the related acid sites might not have significant influence on the SCR activity for x Mn–2Cu/SAPO. For x Mn/SAPO ($x = 1, 2, 4, 8$),¹⁴ the amount of ammonia desorption also decreased as manganese loading increased.

3.7 H₂-TPR

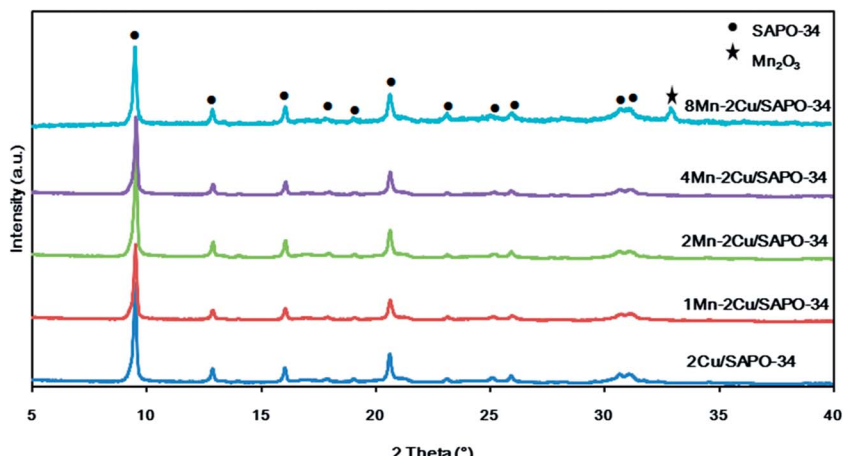
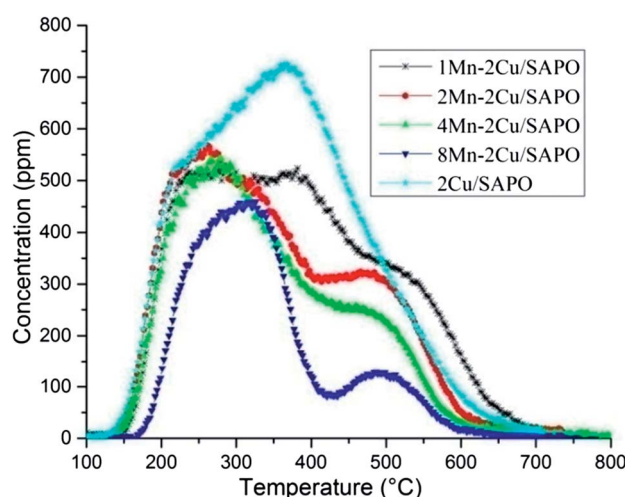
The H₂-TPR profiles of the four catalysts are depicted in Fig. 6. For 8Mn–2Cu/SAPO, there were two sharp peaks at 278 °C and 327 °C, respectively. The former indicated the reduction of MnO₂ to Mn₂O₃, while the latter indicated the reduction of Mn₂O₃ to Mn₃O₄.³ Compared with Mn/TiO₂,^{18,19} the reduction peaks moved to a lower temperature, indicating its good redox capacity.

Table 1 Physico-chemical properties

Sample	BET surface area (m ² g ⁻¹)	Micro pore volume (cm ³ g ⁻¹)	Micro pore width (nm)	Mn loading ^a (wt%)	Cu loading ^a (wt%)	Mn/Cu molar ratio ^a	Mn/Cu molar ratio ^b
2Cu/SAPO	467	0.231	2.36		1.951		
1Mn–2Cu/SAPO	459	0.230	1.90	1.054	2.016	0.51	1.587
2Mn–2Cu/SAPO	430	0.228	2.53	2.108	1.998	1.055	2.594
4Mn–2Cu/SAPO	426	0.206	1.96	3.846	2.048	1.878	3.424
8Mn–2Cu/SAPO	424	0.196	2.54	8.306	1.848	4.495	9.104

^a Determined by ICP. ^b Determined by XPS.



Fig. 4 XRD patterns of x Mn-2Cu/SAPO.Fig. 5 NH_3 -TPD of x Mn-2Cu/SAPO.

In the TPR patterns, the reduction process of copper in SAPO-34 zeolite could be separated into two steps, *i.e.* the reduction of Cu^{2+} to Cu^+ at about $300\text{ }^\circ\text{C}$ and the reduction of Cu^+ to Cu^0 at about $440\text{ }^\circ\text{C}$,¹⁰ while in Fig. 6 the peak strength of 1Mn-2Cu/SAPO was relatively low, indicating that the peak strength of 2Cu/SAPO was even lower, making the assignation of copper species difficult and unreasonable, so the assignation of copper was not discussed. The hydrogen consumption amount of copper reduction was relatively low, so the ignorance of copper species would not influence the discussion of manganese species.

As manganese loading increased, the peaks at $278\text{ }^\circ\text{C}$ and $327\text{ }^\circ\text{C}$ increased, indicating that the proportion of Mn^{3+} and Mn^{4+} species increased. For 1Mn-2Cu/SAPO catalysts, only one broad peak between 250 and $400\text{ }^\circ\text{C}$ was detectable, indicating that most of the manganese species were Mn^{2+} since Mn^{2+} cannot be reduced below $700\text{ }^\circ\text{C}$. Since Mn^{2+} species were inactive for SCR reaction, addition of manganese did not help increase the activity for 1Mn-2Cu/SAPO, which was consistent

with E_p trends and NO oxidation results. For 2Mn-2Cu/SAPO the peaks belonging to Mn^{3+} and Mn^{4+} species started to generate, indicating its good redox capacity. For 4Mn-2Cu/SAPO and 8Mn-2Cu/SAPO, the peaks at $278\text{ }^\circ\text{C}$ and $327\text{ }^\circ\text{C}$ increased greatly, indicating that the ratio of Mn^{3+} and Mn^{4+} increased. When manganese loading was above 2%, Mn^{3+} and Mn^{4+} species began to generate, which is consistent with the E_p trends and NO oxidation results.

Compared with x Mn/SAPO ($x = 1, 2, 4, 8$) previously prepared,¹⁴ for which the temperature peak of H_2 -TPR was about $353\text{ }^\circ\text{C}$ and $442\text{ }^\circ\text{C}$, the peak temperature moved to a lower value, indicating that the interactions between copper and manganese species might accelerate oxidation reaction and increase the catalytic oxidation activity, which would promote low temperature SCR reaction.

Table 2 represents the hydrogen consumption of x Mn-2Cu/SAPO during TPR processes. As manganese loading increased, hydrogen consumption increased gradually, while the actual hydrogen consumption was much lower than the theoretical value. The ratio of experimental hydrogen consumption to

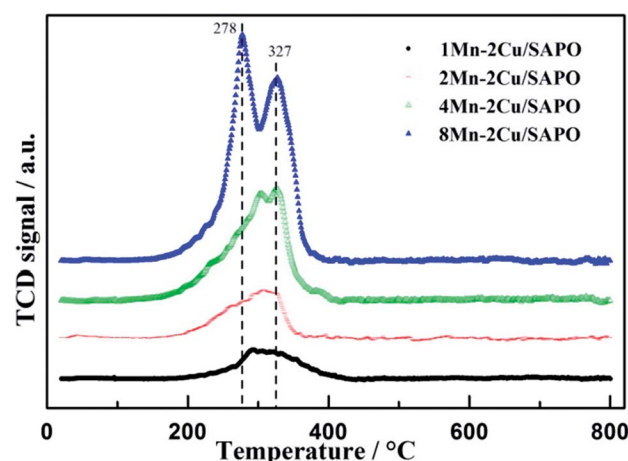
Fig. 6 H_2 -TPR of x Mn-2Cu/SAPO.

Table 2 Hydrogen consumption

	8Mn-2Cu/SAPO	4Mn-2Cu/SAPO	2Mn-2Cu/SAPO	1Mn-2Cu/SAPO
Hydrogen consumption ^a (10^{-2} mmol)	2.04	1.11	0.468	0.327
Hydrogen consumption ^b (10^{-2} mmol)	3.53	2.08	1.35	0.989
Hydrogen consumption ^a /hydrogen consumption ^b (%)	57.8	53.4	35.7	33.1

^a The actual hydrogen consumption calculated from TPR profiles. ^b The theoretical hydrogen consumption, assuming all of the Cu species are Cu^{2+} and reduced to Cu^0 , and all of the Mn species are Mn^{4+} and reduced to Mn^{2+} in TPR process.

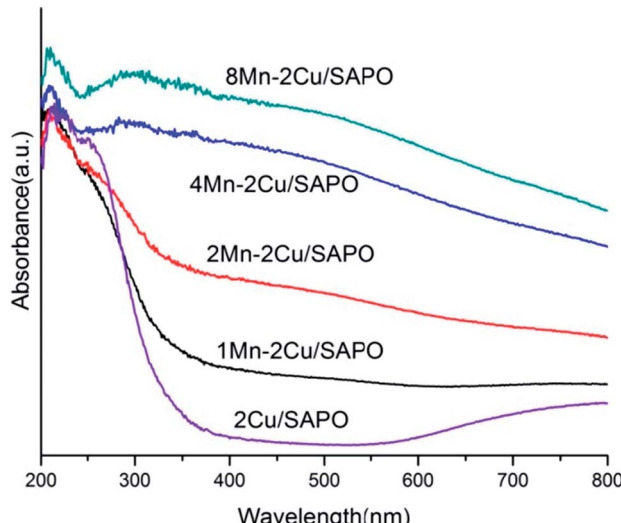


Fig. 7 DR UV-vis spectra recorded under ambient atmosphere of $x\text{Mn-2Cu/SAPO}$.

theoretical hydrogen consumption of 8Mn-2Cu/SAPO and 4Mn-2Cu/SAPO was similar, as well as that of 1Mn-2Cu/SAPO and 2Mn-2Cu/SAPO. Since the ratio of experimental hydrogen consumption to theoretical hydrogen consumption was highly related to the proportion of Mn^{3+} and Mn^{4+} , the proportion of Mn^{3+} and Mn^{4+} for 1Mn-2Cu/SAPO and 2Mn-2Cu/SAPO were similar, as well as that of 8Mn-2Cu/SAPO and 4Mn-2Cu/SAPO.

3.8 DR UV-vis

The UV-vis spectra of the samples are given in Fig. 7. In Fig. 7, the absorption bands at 240 nm were related to the charge transfer processes between the framework aluminum and oxygen atoms of aluminophosphate.²⁰ The band at 220 nm was attributed to the oxygen-to-metal charge-transfer of the isolated $\text{Cu}^+/\text{Cu}^{2+}$ bound on the framework of zeolites.^{21,22} The bands at 280–300 nm were assigned to copper oxide clusters.^{23,24} The bands at 700–800 nm were assigned to the d-d transition of isolated Cu^{2+} in weak disordered octahedral coordination of O-containing ligands.^{23,25}

The broad bands in the range 320–380 nm were attributed to the $\text{Mn}^{3+} \leftarrow \text{O}^{2-}$ charge transfer transition superimposed on the ${}^5\text{B}_{1g} \rightarrow {}^5\text{B}_{2g}$ crystal field d-d transition.²⁶ The band at 322 nm was tentatively assigned to the $\text{Mn}^{3+} \leftarrow \text{O}^{2-}$ charge transfer in Mn_3O_4 in which manganese was octahedrally coordinated with oxygen.^{27–29} The band at 255–276 nm could be assigned to the $\text{Mn}^{2+} \leftarrow \text{O}^{2-}$ charge transfer transition in tetrahedral oxygen coordination.^{27,28} In the $\alpha\text{-Mn}_2\text{O}_3$ structure, Mn^{3+} ions occupied octahedral sites and, if highly symmetric, a single spin-allowed absorption band in the d-d transition region was expected similarly to $[\text{Mn}(\text{H}_2\text{O})_6]^{3+}$ at 500 nm.

As manganese loading increased, the band at 320–380 nm increased gradually especially when the manganese loading changed from 2% to 4%, indicating that the percentage of Mn^{3+} increased.

Meanwhile the bands at 255–276 nm were generated, indicating that the percentage of Mn^{2+} decreased. Consistent with

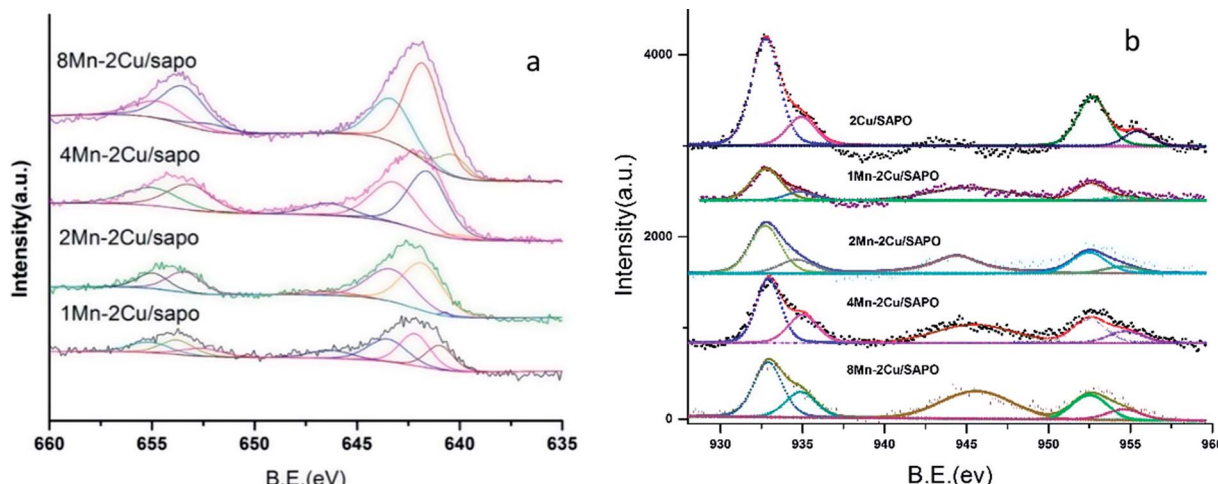


Fig. 8 XPS spectra of (a) Mn 2P and (b) Cu 2P.



Table 3 Binding energies (eV) of Mn^{x+} 2P_{3/2} and the percentage of Mn^{x+}

Compound	Mn ²⁺		Mn ³⁺		Mn ⁴⁺	
	Peak (eV)	%	Peak (eV)	%	Peak (eV)	%
1Mn-2Cu/SAPO	640.97	26.3	642.19	38.8	643.51	34.9
2Mn-2Cu/SAPO	640.69	0.5	641.83	55.8	643.36	44.2
4Mn-2Cu/SAPO	639.99	0.1	641.58	56.2	643.18	43.8
8Mn-2Cu/SAPO	640.42	9.0	641.83	60.7	643.44	30.3

Table 4 Binding energies (eV) of Cu^{x+} 2P_{3/2} and the percentage of Cu^{x+}

Compound	Cu ⁺		Cu ²⁺	
	Peak (eV)	%	Peak (eV)	%
2Cu/SAPO	932.75	78.3	934.93	21.7
1Mn-2Cu/SAPO	932.75	79.9	934.81	20.1
2Mn-2Cu/SAPO	932.64	75.6	934.56	24.4
4Mn-2Cu/SAPO	932.84	65.7	934.93	34.3
8Mn-2Cu/SAPO	932.82	67.2	934.80	32.7

the H₂-TPR result, the UV-vis results showed the partial oxidation of Mn²⁺ to Mn³⁺ or Mn⁴⁺, when manganese loading increased.

The UV-vis spectra of xMn-2Cu/SAPO were similar to that of xMn/SAPO ($x = 1, 2, 4, 8$)¹⁴ in that the percentage of Mn³⁺ increased suddenly as manganese loading increased from 2% to 4%, indicating that the distribution of manganese species might not have been influenced by the copper species.

3.9 XPS

Fig. 8a shows the XPS spectra of Mn 2P of xMn-2Cu/SAPO. Two main peaks assigned respectively to Mn 2P_{3/2} at 642.5 eV and Mn 2P_{1/2} at 654 eV were observed.

By performing peak-fitting deconvolution, the Mn 2P_{3/2} peak could be separated into three peaks, *i.e.* 640.8–640.9 eV, 642.1–642.3 eV and 643.8–644.1 eV, which correspond to the Mn²⁺ species, Mn³⁺ species and Mn⁴⁺ species,^{3,30} respectively. The atomic percentages of manganese species in different valence states are shown in Table 3.

When manganese loading increased from 1% to 2%, the percentage of Mn³⁺ and Mn⁴⁺ increased and the percentage of

Mn²⁺ decreased a lot, while when manganese loading kept increasing, the percentage of manganese species did not change much. This was consistent with the NO oxidation, H₂-TPR and UV-vis results.

As manganese loading increased, the percentage of Mn³⁺ species increased, and the percentage of Mn⁴⁺ species also increased, except for 8Mn-2Cu/SAPO. This was consistent with the UV-vis and H₂-TPR results. Though H₂-TPR, UV-vis and XPS all showed that there were many Mn⁴⁺ and Mn³⁺ species on 8Mn-2Cu/SAPO, the XRD patterns showed the Mn₂O₃ phase, which indicated the aggregation of manganese species and decreased manganese dispersity, and the NH₃-TPD results showed that there were much less acid sites. The decreased manganese dispersity and the less acid sites could lead to the lower SCR activity. The XPS spectra of xMn/SAPO ($x = 1, 2, 4, 8$)¹⁴ also showed that most of the manganese species were Mn³⁺ and Mn⁴⁺.

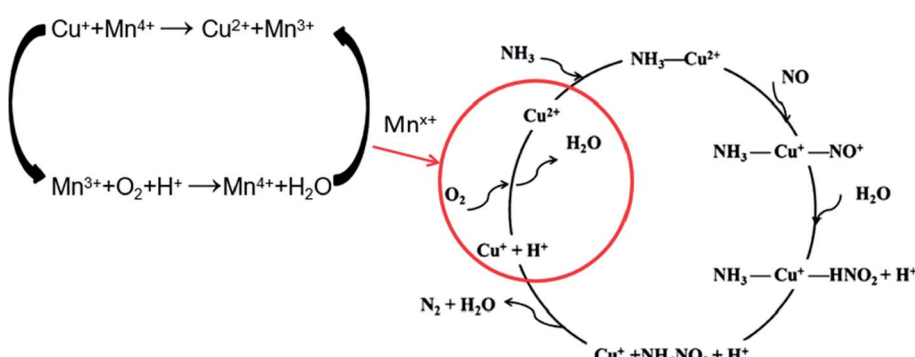
Fig. 8b shows the XPS spectra of Cu 2P; two peaks respectively attributed to Cu 2P_{3/2} at 932.6 eV and Cu 2P_{1/2} at 952.7 eV were observed. The shake-up satellite was found at about 945 eV, indicating the presence of Cu²⁺ species. As manganese loading increased, the shake-up satellite increased, indicating that the amount of Cu²⁺ species increased.

By performing peak-fitting deconvolution, the Cu 2P_{3/2} peak could be separated into two peaks, at about 932.7 eV and 934.8 eV respectively, which was ascribed to the Cu⁺ and Cu²⁺ species.³¹ The atom percentage of Cu⁺ and Cu²⁺ is shown in Table 4. As manganese loading increased, the ratio of Cu²⁺ increased, indicating that the addition of manganese might oxidize Cu⁺ into Cu²⁺, which was strong evidence that there was strong interaction between the copper and manganese species.

According to the results above and referring to the mechanism of SCR on Cu/SAPO Feng Gao⁹ had put forward, the mechanism of SCR on 2Mn-2Cu/SAPO could be proposed. Fig. 9 showed the proposed low temperature SCR mechanism on 2Mn-2Cu/SAPO. In the catalytic cycle, Mn^{x+} acts as a catalyst for the oxidation step of Cu⁺ to Cu²⁺ in low temperature SCR reaction.

3.10 DRIFTS

In situ DRIFTS was performed to identify the adsorbed NO species. Fig. 10 shows the DRIFTS spectra of 4Mn-2Cu/SAPO at 150 °C exposed to different gas phases. In Fig. 10a, there was only one positive band at 1360 cm⁻¹ in the range of 1200–

**Fig. 9** The proposed low temperature SCR mechanism on 2Mn-2Cu/SAPO.

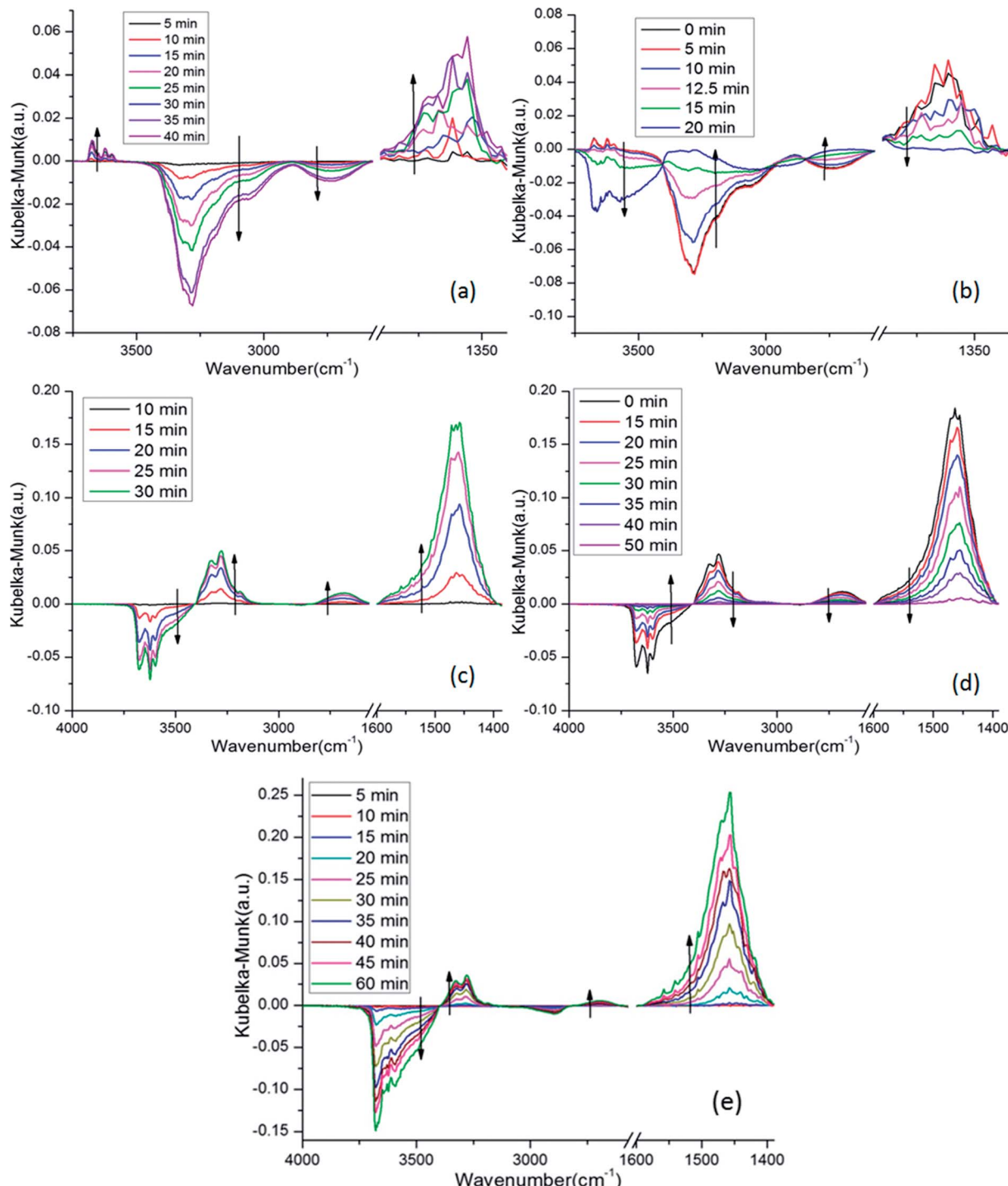


Fig. 10 DRIFTS spectra of 4Mn–2Cu/SAPO at 150 °C exposed to gas phase of (a) 500 ppm NO + 3% vol O₂ + N₂ (balance); (b) 500 ppm NH₃ + 3% vol O₂ + N₂ (balance) after saturation with NO; (c) 500 ppm NH₃ + 3% vol O₂ + N₂ (balance); (d) 500 ppm NO + 3% vol O₂ + N₂ (balance) after saturation with NH₃; (e) 500 ppm NO + 500 ppm NH₃ + 3% vol O₂ + N₂ (balance).

1600 nm, which belongs to nitrite groups,^{32,33} suggesting that those NO species were nitrite groups. The negative bands at 3280 cm⁻¹ and 2730 cm⁻¹ might be attributed to the depletion of Brønsted base sites by the nitrite groups. Three bands at 3675 cm⁻¹, 3623 cm⁻¹ and 3598 cm⁻¹ could be attributed to the appearance of O–H bond of nitrite groups (–N–O–H) in different

chemical environments. When NH₃ was introduced into the system, as shown in Fig. 10b, the bands associated with nitrite groups decreased rapidly, indicating reaction between NH₃ and NO related species.

In Fig. 10c, when NH₃ was introduced into the system, the bands at 1463 cm⁻¹ increased greatly, indicating the generation

of NH_3 species on the Brønsted acid sites,²² while the bands at 1210 cm^{-1} increased slightly, which was consistent with the fact that most of the acid sites on SAPO-34 were Brønsted acid sites. The negative bands at 3681 cm^{-1} , 3625 cm^{-1} and 3600 cm^{-1} were attributed to the depletion of Brønsted acid sites, which was caused by the adsorption of NH_3 . Specifically, the negative bands at 3681 cm^{-1} were assigned to the occupation of P-OH sites by NH_3 . The other two bands were related to the depletion of Si-OH-Al acid sites. The band at $3147\text{--}3396\text{ cm}^{-1}$ was attributed to the N-H stretching vibrations of NH_4^+ .²² When NO was introduced into the system, as shown in Fig. 10d, the bands related to the NH_3 species decreased gradually, which was caused by the reaction between NO and NH_3 .

Fig. 10e and c are similar in that only bands related to NH_3 are visible, while no other bands appeared, especially the negative bands at 2700 cm^{-1} and $3000\text{--}3400\text{ cm}^{-1}$ were invisible.

This was strong evidence that only the Elay-Rideal mechanism was effective on the 4Mn-2Cu/SAPO at $150\text{ }^\circ\text{C}$. Since if Langmuir-Hinshelwood mechanism was effective, the adsorbed and well activated NO and NH_3 related species would react with each other, while the NO species not well activated would accumulated and be detectable for DRIFTS, leading to the appearance of negative bands at 2700 and $3000\text{--}3400\text{ nm}^{-1}$, as Fig. 10a and b show. Due to the fact that those peaks were not observed in Fig. 10e, all these observations lead to the exclusion of the Langmuir-Hinshelwood mechanism and the assurance of the Elay-Rideal mechanism. In the Elay-Rideal mechanism, NO molecules from gas phase directly react with the well activated NH_3 species adsorbed on the surface and leave as products to gas phase with the active site left.

4 Conclusion

In this work, the SCR activity and physicochemical properties of $x\text{Mn-2Cu/SAPO}$ were investigated. The $\text{H}_2\text{-TPR}$, UV-vis and XPS results showed that Mn^{4+} and Mn^{3+} species began to appear when manganese loading reached 2%. The NO oxidation results showed that 2Mn-2Cu/SAPO had the best NO oxidation activity, indicating that good interaction existed between copper and manganese, which was consistent with the XPS results of Cu 2p binding energy.

The graph of the promotion effect of manganese showed that for 2Mn-2Cu/SAPO , manganese had a good promotion effect on 2Cu/SAPO . According to $\text{H}_2\text{-TPR}$, UV-vis, XPS and NO oxidation results, the promotion effect of manganese on 2Cu/SAPO comes from the generation of Mn^{3+} and Mn^{4+} species. The addition of manganese increased the ratio of Cu^{2+} . The Mn^{3+} and Mn^{4+} species might take part in the catalytic cycle and accelerate the oxidation of Cu^+ to Cu^{2+} .

From the *in situ* DRIFTS results, the acid sites on the surface were mainly Brønsted acid sites, the main mechanism on 4Mn-2Cu/SAPO-34 at low temperature was the Elay-Rideal mechanism.

Conflicts of interest

There are no conflicts to declare.

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