Preparation and Characterization of CeO$_2$-MoO$_3$/TiO$_2$ Catalysts for Selective Catalytic Reduction of NO with NH$_3$

Ye Jiang*, Xuechong Wang, Zhimin Xing, Changzhong Bao, Guitao Liang

College of Pipeline and Civil Engineering, China University of Petroleum, Qingdao 266580, China

ABSTRACT

CeO$_2$-MoO$_3$/TiO$_2$ catalysts were prepared by three methods: a single step sol-gel (SG), impregnation (IM) and coprecipitation (CP) method. These catalysts were investigated for selective catalytic reduction (SCR) of NO with NH$_3$. The results indicate that the catalyst prepared by the SG method possessed the widest temperature window (about 250–475°C), best SCR activity below 450°C and resistance against 10% H$_2$O and 1000 ppm SO$_2$ at a gas hourly space velocity (GHSV) of 90,000 h$^{-1}$. According to the results of BET, XRD, XPS, H$_2$-TPR and TEM, it is found that larger BET surface area, more highly dispersed active species of Ce and Mo, presence of more Ce$^{3+}$ and chemisorbed oxygen, synergistic effect among ceria, molybdenum and titania, and better redox ability can contribute to the better deNO$_x$ performance of the CeO$_2$-MoO$_3$/TiO$_2$ catalyst prepared by the SG method, compared with those by the IM and CP methods.

Keywords: Selective catalytic reduction (SCR); CeO$_2$-MoO$_3$/TiO$_2$; NO removal; NH$_3$; Preparation methods.

INTRODUCTION

Nitrogen oxides (NO$_x$), emitted from mobile and stationary sources, are to blame for the worsening environment because of their contribution to acid rain, photochemical smog, haze formation and ozone depletion. The selective catalytic reduction (SCR) with NH$_3$ is currently the most efficient technology for the abatement of NO$_x$ in the flue gas from stationary sources such as coal-fired power plants and municipal solid waste (MSW) incineration plants (Busca et al., 1998; Roy et al., 2009). V$_2$O$_5$-WO$_3$(MoO$_3$)/TiO$_2$ has been put into industrial applications for the past few decades (Pârvulescu et al., 1998; Fu et al., 2014). However, some unavoidable issues have to be raised, such as a relatively narrow operating temperature range (300–400°C), high activity for the oxidation of SO$_2$ to SO$_3$ with increasing vanadium loadings (Dunn et al., 1998), formation of N$_2$O at high temperatures above 400°C (Yates et al., 1996) and toxicity of vanadium species. Owing to the before-mentioned disadvantages of vanadium-based catalysts, an increasing number of researchers have recently turned their attention to ceria for the NH$_3$-SCR reaction due to their high oxygen storage capacity and outstanding redox property (Luo et al., 2001; Guo et al., 2014; Zhu et al., 2015). Ceria have been reported to be mixed with other metal oxides such as TiO$_2$ (Xu et al., 2008; Gao et al., 2010a; Shan et al., 2012), Al$_2$O$_3$ (Shen et al., 2009, Guo et al., 2013), WO$_3$ (Peng et al., 2013), MoO$_3$ (Li and Li, 2014), etc. In our previous study, we developed a CeO$_2$-MoO$_3$/TiO$_2$ catalyst, which exhibited high SCR activity and excellent resistance against 10% H$_2$O and 1000 ppm SO$_2$ (Jiang et al., 2015). It is well accepted that catalyst preparation method has a substantial impact on the loading, dispersion and interaction of the active components (Lee et al., 2014). In addition, because of the presence of H$_2$O and SO$_2$ in flue gas, the resistance against them is important for any potential SCR catalysts from a practical point view. Therefore, it is indispensable to investigate the relation between preparation method and the SCR activity of CeO$_2$-MoO$_3$/TiO$_2$ catalysts in the absence and presence of H$_2$O and SO$_2$.

In this work, CeO$_2$-MoO$_3$/TiO$_2$ catalysts were prepared by three methods: a single step sol-gel (SG), impregnation (IM) and coprecipitation (CP) method. The main purpose of this investigation was to determine an optimal method for preparing CeO$_2$-MoO$_3$/TiO$_2$ catalysts with the best deNO$_x$ performance.

METHODS

Catalyst Preparation

In this work, the mass ratio of CeO$_2$-MoO$_3$/TiO$_2$ was 2:1:10 in the prepared CeO$_2$-MoO$_3$/TiO$_2$ catalysts. The CeO$_2$-MoO$_3$/TiO$_2$ catalysts were denoted as CMT$(y)$, where $y$ represented the preparation method of CeO$_2$-MoO$_3$/TiO$_2$ catalyst. Nitric acid, anhydrous ethanol and aqueous ammonia
(3 mol L⁻¹) were purchased from Xilong Chemical Industry Corp. (Guangzhou, China), and other reagents (analytically pure) used in the catalyst preparation were purchased from Sinopharm Chemical Reagent Corp. (Shanghai, China).

**Single step Sol-Gel Method (SG)**

One solution of butyl titanate (0.1 mol) and anhydrous ethanol (2.3 mol) was added dropwise to the other solution of deionized water (1.9 mol), anhydrous ethanol (0.6 mol), nitric acid (0.1 mol) and cerium nitrate (0.01 mol) at room temperature under continuous and vigorous stirring for 3 h to yield a transparent colloidal solution. The solution was dried at 80°C for 24 h to form the xerogel, and then calcined in air at 500°C for 5 h.

**Impregnation Method (IM)**

The catalyst was prepared by impregnation of TiO₂ (0.125 mol) with an aqueous solution containing cerium nitrate (0.012 mol) and ammonium molybdate (0.001 mol). The sample was dried at 110°C for 12 h, followed by calcination in air at 500°C for 5 h.

**Coprecipitation Method (CP)**

Titanium sulfate (0.125 mol), cerium nitrate (0.012 mol) and ammonium molybdate (0.001 mol) were dissolved in deionized water (11.1 mol). Aqueous ammonia was slowly added to the solution under continuous and vigorous stirring at room temperature until pH reached 11. After aged for 3 h, the sediment was filtered and washed with deionized water for six times to eliminate sulfate. The obtained sample was dried at 110°C for 12 h, and then calcined in air at 500°C for 5 h.

**SCR Activity Measurements**

The SCR activity measurements were carried out in a fixed-bed quartz reactor of 8 mm inner diameter using 0.35 g catalyst (150–250 µm). The feed gas composition was as follows: 1000 ppm NO, 1000 ppm NH₃, 3 vol.% O₂, 10 vol.% H₂O (when used), 1000 ppm SO₂ (when used) and N₂ as balance gas. H₂O was injected with a micro-syringe and 1000 ppm SO₂ (when used) using Barrett-Joyner-Halenda (BJH) method. The pore size distribution was measured by N₂ adsorption/desorption at −196°C using a ASAP2020-M system (Micromeritics Instrument Corp.). The specific surface area were evaluated by the Brunauer-Emmette-Teller (BET) method. The pore size distribution was measured using Barrett-Joyner-Halenda (BJH) method.

The crystallite structure was identified by the X-ray diffractometer (Panalytical Corp.) with a X’Pert PRO MPD system by using Cu Kα radiation.

X-ray photoelectron spectroscopy (XPS) was used to determine the atomic concentration and the chemical state of the elements on the catalyst surface with a Thermo ESCALAB 250 spectrometer using monochromated Al Kα X-rays (hv = 1486.6 eV) as a radiation source at 150 W. The binding energies were referenced to the C 1s line at 284.6 eV.

To evaluate the redox ability of the catalysts, temperature programmed reduction (H₂-TPR) was performed on an Autosorb-iQ-C chemisorption analyzer (Quantachrome Instrument Corp.) with a thermal conductivity detector (TCD).

The catalysts were analyzed with a JOEL JEM-2010 electron microscope to observe their microstructures.

**RESULTS AND DISCUSSION**

**Catalytic Activity Evaluation**

**NH₃-SCR Activity**

The activities of various samples for the SCR of NO with NH₃ were measured as a function of temperatures and the results are shown in Fig. 1. It was clear that CMT(SG) possessed the best catalytic performance at the temperature below 450°C and the widest activity temperature window (about 250–475°C) for > 90% NO conversion. When the reaction temperature exceeded 450°C, the activity was shown as follows: CMT(CP) > CMT(SG) > CMT(IM). These indicated that catalyst preparation methods had a massive impact on the SCR performance of CeO₂-MoO₃/TiO₂ catalysts.

**Effect of H₂O and SO₂**

H₂O and SO₂ are the main components in flue gas from stationary sources, so it is important to investigate their effect on the SCR activities of various catalysts and the results are shown in Figs. 2–4. As shown in Fig. 2, when H₂O was added into the feed gas, the activity of CMT(SG) decreased a little in the temperature range of 150–300°C and was not nearly inhibited at 300–450°C. As for CMT(IM), the presence of H₂O caused a marked decrease in its activity at the temperatures below 400°C and had almost no inhibition at 400–450°C. It was noted that NO conversions over CMT(SG) and CMT(IM) were both enhanced at 500°C after the addition of H₂O, and yet the activity of CMT(SG) was higher than that of CMT(IM). With regard to CMT(CP), a nearly complete loss of its activity was observed. The H₂O adsorbed on catalyst surface can block the active sites, leading to the decrease of the NH₃-SCR activity (Liu et al., 2014b). It could be seen that CMT(SG) had the best tolerance to H₂O in these three samples.

Effect of SO₂ on NO conversion over CeO₂-MoO₃/TiO₂ catalysts prepared by different methods is shown in Fig. 3. Before adding SO₂, the SCR reaction had been stabilized for 1 h, and nearly 100% of NO conversions were achieved over CMT(SG) and CMT(IM). The addition of 1000 ppm SO₂ to the feed gas had little effect on the catalytic activities of CMT(SG). On the contrary, the catalytic activity of CMT(IM) decreased with time and about 80% NO conversion was
Fig. 1. NO conversion as a function of reaction temperature for various catalysts. Reaction conditions: [NO] = [NH₃] = 1000 ppm, [O2] = 3 vol.%, balance N₂ and GHSV = 90,000 h⁻¹.

Fig. 2. Effect of H₂O on NO conversion over various catalysts. (■) CMT(SG), (●) CMT(IM), (▲) CMT(CP). Reaction conditions: [NO] = [NH₃] = 1000 ppm, [O2] = 3 vol.%, [H₂O] = 10% (□), balance N₂ and GHSV = 90,000 h⁻¹.

obtained after 7.5 h. As for CMT(CP), the original activity was substantially lower than those of the other two catalysts. After the addition of SO₂, it still remained the original low activity. From Fig. 3, it can be also seen that there were almost no changes in the activities of CMT(IM) and CMT(CP) when SO₂ was cut off. According to Xu et al. (2009), the presence of SO₂ in the feed gas led to the formation of NH₄HSO₄, Ce(SO₄)₂ and Ce₂(SO₄)₃ on Ce/TiO₂ catalyst. NH₄HSO₄ could deposit on the surface of the catalyst and block the active sites. Ce(SO₄)₂ and Ce₂(SO₄)₃ disrupted the Ce³⁺/Ce⁴⁺ redox cycle and inhibited the formation and adsorption of nitrate species (Xu et al., 2009). Compared with CMT(IM) and CMT(CP), CMT(SG) might inhibit the formation of these surface sulfate species, thereby having the best tolerance to SO₂.

Fig. 4 illustrates the co-effect of H₂O and SO₂ on NO conversion over various catalysts. CMT(SG) exhibited the highest resistance to H₂O and SO₂ and a relatively high NO conversion (about 80%) was still maintained after 7.5 h. Upon switching off H₂O and SO₂, the activity was nearly restored to its original level. For CMT(IM), the NO conversion decreased sharply to about 10% in 3.5 h and subsequently the low activity was retained. As to CMT(CP), the addition of H₂O and SO₂ led to the rapid decrease in its activity and less than 10% of NO conversion was obtained after 3.5 h. After removing H₂O and SO₂, only a very small part of the SCR activity was restored over CMT(IM) and CMT(CP). It was obvious that the preparation methods had a different impact on the resistance of CeO₂-MoO₃/TiO₂ catalysts to H₂O and SO₂. It is known that Ce(SO₄)₂ and Ce₂(SO₄)₃ have high thermal stability (Xu et al., 2009). Considering that the removal of H₂O and SO₂ resulted in regaining the activity of CMT(SG), the deactivation of CMT(SG) by H₂O and SO₂ might be mainly attributed to the deposition of NH₄HSO₄ blocking the active sites on the catalyst surface. Furthermore, SO₂ hardly reacted with...
Fig. 3. Effect of SO₂ on NO conversion over various catalysts at 350°C. Reaction conditions: [NO] = [NH₃] = 1000 ppm, [O₂] = 3 vol.%, [SO₂] = 1000 ppm, balance N₂ and GHSV = 90,000 h⁻¹.

CMT(SG) to form Ce(SO₄)₂ and Ce₂(SO₄)₃. It might be ascribed to the competitive adsorption of H₂O at the catalyst surface and the strong interaction among Ce, Mo and Ti in CMT(SG). By contrast, the great loss of the activity of CMT(IM) and CMT(CP) might be attributed to the formation of NH₄HSO₄, Ce(SO₄)₂ and Ce₂(SO₄)₃.

Characterization of Catalysts

BET Results

The physical properties of the CeO₂-MoO₃/TiO₂ catalysts prepared by different methods are compared in Table 1. The BET surface area of CMT(SG) was the largest among the three samples. The total pore volume is in the following sequence: CMT(SG) < CMT(IM) < CMT(CP). The order of the average pore diameter was: CMT(SG) < CMT(CP) < CMT(IM). It was clear that different preparation methods could result in the different microstructure of CeO₂-MoO₃/TiO₂ catalysts. The similar phenomena were found over CeO₂/TiO₂ (Gao et al., 2010b), CeO₂/Al₂O₃ (Guo et al., 2013) and CeO₂-ZrO₂-WO₃ (Ning et al., 2015) prepared by different methods.

XRD Results

XRD patterns of various samples are presented in Fig. 5. CMT(SG) exhibited only the characteristic lines of anatase TiO₂. The absence of the characteristic diffraction peaks due to CeO₂ and MoO₃ indicated that CeO₂ and MoO₃ were highly dispersed and existed as amorphous or highly dispersed species on the surface of anatase TiO₂. This also suggested that there existed a strong synergistic effect among Ce, Mo and Ti in CMT(SG). By contrast, an extremely weak diffraction peak of Ce₂(MoO₄)₃ (PDF30-0303) was
Table 1. Physical properties of different samples.

<table>
<thead>
<tr>
<th>Samples</th>
<th>BET surface area (m² g⁻¹)</th>
<th>Total pore volume (cm³ g⁻¹)</th>
<th>Average pore diameter (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMT(IM)</td>
<td>65.30</td>
<td>0.2464</td>
<td>12.749</td>
</tr>
<tr>
<td>CMT(SG)</td>
<td>103.90</td>
<td>0.1779</td>
<td>5.664</td>
</tr>
<tr>
<td>CMT(CP)</td>
<td>101.53</td>
<td>0.3480</td>
<td>11.882</td>
</tr>
</tbody>
</table>

Fig. 5. XRD patterns of (a) CMT(IM), (b) CMT(CP) and (c) CMT(SG).

observed on CMT(CP) in addition to the characteristic lines of anatase TiO₂. As for CMT(IM), quite a few diffraction peaks ascribed to cubic CeO₂ were detected, as reported by Liu et al. (2014a). This implied that ceria crystallites were present in CMT(IM). Furthermore, it can be seen from Fig. 5 that the sharpness and relative intensity of the main anatase peak (101) for CMT(IM) was obviously larger than the two other samples. This indicated the interaction was weakest among Ce, Mo and Ti in CMT(IM), thereby giving rise to the segregation of CeO₂ crystallites from TiO₂ carrier. It seems reasonable to suppose that, over the CeO₂-MoO₃/TiO₂ catalyst prepared by a single step sol-gel method, ceria could be well dispersed and kept as an amorphous phase, which would help to improve its catalytic activity.

XPS Results

Complicated XPS spectra of Ce 3d for various samples are illustrated in Fig. 6. The sub-bands labeled u, u′, u″ and v, v′, v″ are assigned to Ce⁴⁺, while u′ and v′ belong to Ce³⁺ (Gao et al., 2010c; Chen et al., 2010; Shan et al., 2012). It is generally acknowledged that ceria-based catalysts possess the excellent reducibility via the redox shift between Ce⁴⁺ and Ce³⁺, which enhances the oxygen store/release capacity. Accordingly, the SCR activity of CeO₂-MoO₃/TiO₂ catalyst should have something to do with the ratio of Ce³⁺/(Ce⁴⁺ + Ce³⁺). From Fig. 6, it can be seen that the redox couple of Ce³⁺/Ce⁴⁺ was observed in all of the three samples. The ratios of Ce³⁺/(Ce⁴⁺ + Ce³⁺) for CeO₂-MoO₃/TiO₂ catalysts were shown as follows: 26.3% for CMT(IM) < 35.4% for CMT(CP) < 41.7% for CMT(SG). This indicated that preparation methods could lead to the difference in the Ce valance state over CeO₂-MoO₃/TiO₂ catalysts while more Ce³⁺ were present in CMT(SG). The existence of Ce³⁺ could create a charge imbalance, vacancies and unsaturated chemical bonds on the catalyst surface, leading to the increase in chemisorbed oxygen or/and weakly bonded oxygen species on the catalyst surface (Yang et al., 2006). As a result, it could be inferred that CMT(SG) would possess superior deNOₓ performance, which had been exactly confirmed by the results of the activity tests.

The O 1s XPS spectra of various samples can be fitted into three overlapping peaks, as shown in Fig. 7. The sub-band at 529.0–530.1 eV can be assigned to the lattice oxygen (denoted by Oα) (Chen et al., 2010; Gao et al., 2010c). Two shoulder peaks at 530.6–531.0 eV and 532.4–533.0 eV are ascribed to the surface adsorbed oxygen (denoted by Oβ) and the surface oxygen by hydroxyl species and/or adsorbed water species as contaminants at the surface (denoted by Oγ), respectively (Yang et al., 2006; Chen et al., 2010; Gao et al., 2010c). The calculated Oβ ratio increased according to the following sequence: 31.2% for CMT(IM) < 32.7% for CMT(CP) < 34.1% for CMT(SG). Surface adsorbed oxygen (Oβ) has been reported to play a substantial role in SCR reaction due to its higher mobility than lattice oxygen (Oα) (Shan et al., 2012). High Oβ ratio facilitated the oxidation of NO to NO₂ in the SCR reaction (Adamowska et al., 2008; Chen et al., 2012), thereby improving the activity of the catalysts. This was in good agreement with the results of the activity tests.
**H₂-TPR Results**

H₂-TPR profiles of various samples are compared in Fig. 8. For CMT(SG), a single broad reduction peak centered at 548°C was observed and the onset temperature ($T_{\text{onset}}$) was about 350°C. The reduction process of CMT(SG) could be interpreted as the coreduction of surface oxygen of ceria and well dispersed Mo species. It should be noted that the reduction of bulk ceria did not take place in the temperature range of 300–700°C, since it occurred only above 750°C (Devaiah et al., 2014; Li et al., 2014). CMT(IM) possessed two reduction peaks entered at 478°C and 747°C, which was similar to that reported in the study of Liu et al. (2014a). The first peak could be associated with the coreduction of surface oxygen of ceria and octahedral well dispersed Mo species, while the second peak could be assigned to the coreduction of bulk ceria and tetrahedral Mo species (Liu et al., 2014a). As for CMT(CP), the reduction temperature of the main peak shifted to higher temperature (612°C). From Fig. 8, it could be also found that $T_{\text{onset}}$ of CMT(SG) was the lowest among the three samples. This suggested that CMT(SG) could be reduced most easily. From the BET and XRD results, it could be inferred that the easier reduction for CMT(SG) might be attributed to its higher surface area and better dispersion of Ce and Mo on TiO₂ carrier, which were highly beneficial in improving the activity of the catalysts. In addition, it should be noted that CMT(SG) and CMT(CP) showed distinct H₂-TPR behaviors though displayed similar crystallinity of anatase TiO₂ and dispersion of ceria and MoO₃ (Fig. 5). This implied that the interaction among Ce, Mo and Ti might have a significant impact on...
the reducibility of the samples. According to the XPS results, the ratios of Ce$^{3+}$/Ce$^{4+}$ were different in various samples. It has been reported that Mo$^{6+}$ species can be present in both tetrahedral and octahedral coordination (Park et al., 1997). Therefore, it can be inferred that the difference in H$_2$ consumption for various samples was likely to correlate with the formation of subtly different surface structure of Ce and Mo oxide species, resulting from different catalyst preparation methods. H$_2$-TPR results demonstrated that CMT(SG) possessed the excellent redox ability.

**TEM Results**

The HR-TEM micrographs of various samples are shown in Fig. 9. As shown in Fig. 9(a), the primary particle size of CMT(IM) was about 20 nm, and the crystals of TiO$_2$ and CeO$_2$ could be observed distinctly. This implied that crystalline phase of TiO$_2$ and CeO$_2$ existed in CMT(IM), which was consistent with XRD results. There were three kinds of lattice fringes: 0.35 nm, 0.32 nm and 0.31 nm of lattice fringes matched anatase (101) (Jiang et al., 2015), CeO$_2$ (111) (Vantomme et al., 2005) and MoO$_3$(021) (Jiang et al., 2017).
and Song, 2007), respectively. It was noted that MoO3 was not determined by XRD. It might be because MoO3 was well dispersed and existed as amorphous or highly dispersed species in CMT(IM). For CMT(SG) and CMT(CP), the primary particle size was less than 10 nm, and the crystals of TiO2, CeO2 and MoO3 could not be discriminated. These results indicated that less crystalline phase of TiO2 formed in these two samples, which was in good agreement with XRD results. In CMT(SG), only lattice fringes ascribed to anatase TiO2 were determined. This meant that CeO2 and MoO3 were well dispersed and existed as amorphous or highly dispersed species. As to CMT(CP), besides the lattice fringes matching anatase (101), 0.34 nm and 0.23 nm of lattice fringes were identified, which were ascribed to MoO3 (040) (Du et al., 2014) and MoO3 (041) (Zhong et al., 2014). According to the XRD results, Ce2(MoO4)3 was present in CMT(CP). Therefore, it could be concluded that a single step sol-gel method might improve the dispersion of CeO2 and MoO3 in CMT(SG), which was conducive to enhance its catalytic activity.

CONCLUSIONS

CeO2-MoO3/TiO2 catalysts were prepared by three methods. The catalyst prepared by a single step sol-gel method showed the best SCR activity and resistance to 10% H2O and 1000 ppm SO2. From the results of BET, XRD, XPS, H2-TPR and TEM, it could be inferred that the superior deNOX performance should be attributed to large surface area, highly dispersed active species of Ce and Mo, presence of more Ce3+ and chemisorbed oxygen, strong interaction among ceria, molybdenum and titania, and good redox ability. As a result, a single step sol-gel method is an effective one for the preparation of CeO2-MoO3/TiO2 catalyst for NH3-SCR of NO.

ACKNOWLEDGMENTS

The authors wish to express their sincere appreciation for the financial supports from the National Science Foundation of China (No. 51506226), Shandong Provincial Natural Science Foundation (No. ZR2015EM010) and "the Fundamental Research Funds for the Central Universities" (No. 15CX05005A).

REFERENCES

Devaiah, D., Jampaiah, D., Saikia, P. and Reddy B.M. (2014). Structure dependent catalytic activity of Ce0.8Th0.2O2−δ and TiO2 supported Ce0.8Th0.2O2−δ solid solutions for CO oxidation. J. Ind. Eng. Chem. 20: 444–453.


Received for review, August 30, 2016
Revised, January 1, 2017
Accepted, January 5, 2017