Influence of pretreatment conditions on low-temperature carbon monoxide oxidation over CeO2/Co3O4 catalysts

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Received 19 September 2005; received in revised form 8 April 2006; accepted 12 April 2006
Available online 12 June 2006

Abstract

The modification of ceria (CeO2) on high surface area cobaltic oxide (Co3O4) and the promotion effect of CO oxidation via different pretreatment conditions were studied. The high-valence cobalt oxide was prepared first by precipitation-oxidation in ice bath, followed by reduction at 230 °C to get pure and high surface area cobaltic oxide (assigned as R230, SBET = 100 m2 g−1). Further, the Ce20/R230 catalyst was prepared by impregnation with cerium nitrate (20% cerium loading). Pretreatment of Ce20/R230 catalyst was divided into two methods: reduction (under 200 and 400 °C) and calcination (under 350 and 550 °C). The catalysts were characterized by X-ray diffraction (XRD), infrared (IR), and diffuse reflectance spectroscopy (DRS), and temperature-programmed reduction (TPR) and by nitrogen adsorption at 196 °C. The results showed that pretreatment under low-temperatures obtained both larger surface area and better activity. The Ce20/R230-H200 catalyst exhibited the highest surface area (SBET = 109 m2 g−1) and the best catalytic activity in CO oxidation with T50 (50% conversion) at 88 °C among series of catalysts due to the combined effect of cobaltic oxide and ceria. The optimized pretreatment of CeO2/Co3O4 catalysts can clearly enhance the catalytic activity.

Keywords: Cobaltic oxide; Ceria; CO oxidation

1. Introduction

Low-temperature abatement of carbon monoxide is important in environmental pollution control [1]. There are several reports describing the use of noble metals for CO oxidation at ambient temperatures [2–6]. However, the high cost of noble metals and their sensitivity to sulfur poisoning have stimulated the search for substitute catalysts. Because of the price and the limited availability of previous metals, considerable attention has been paid to various transition metal oxides and mixed metal oxides [7–12]. Among these metal oxides, cobaltic oxide (Co3O4) is very attractive for the preparation of oxidation catalysts because of the presence of mobile oxygen in Co3O4 [8–13]. The high activity of Co3O4 on CO oxidation is likely to be due to the relatively low ΔH of vaporization of O2 [14,15]. This means that the Co–O bond strength of Co3O4 is relatively weak, so easily reactive oxygen can be obtained from the lattice oxygen. Thus, many researchers have measured the catalytic activity of Co3O4 for CO oxidation [8–13,16].

Ceria has a high oxygen storage capacity and well-known catalytic and redox properties (couples of Ce4+/Ce3+), making more oxygen available for the oxidation process [17]. The most important property is the oxygen storage and releasing capacity of CeO2 [18–22]. It improves CO oxidation and NOx reduction. As is also well known, CeO2 is a promoter additive [23–25]. The rapid growth in the applications and characterization of CeO2-containing catalysts has been documented [26–32]. In particular, the CuO/CeO2 catalyst was reported to be very active for the complete CO oxidation and even comparable to noble metals [26]. Good catalytic activity is also observed for CoO/CeO2 composite catalysts [30], for which the T50 (50% conversion) approaches 125 °C. However, finding strategies to improve and control redox properties in preparation process to raise the catalytic activity is still a fundamental challenge.
In a previous work, we have demonstrated that the activity of Co$_3$O$_4$ toward CO oxidation is enhanced significantly by increasing the surface area ($S_{BET}$) [12]. For this reason the present work has been devoted to study the effect of ceria on the high surface area cobaltic oxide. The study object of this work is to understand the correlations between the effects of pretreatment conditions on the CeO$_2$/Co$_3$O$_4$ catalysts and the catalytic properties for CO oxidation. The characteristics of catalysts are determined by means of X-ray diffraction (XRD), infrared (IR), and diffuse reflectance spectroscopy (DRS), and by nitrogen adsorption at $-196$ °C and temperature-programmed reduction (TPR). The catalytic activities for CO oxidation of CeO$_2$/Co$_3$O$_4$ catalysts are also compared.

2. Experimental

2.1. Catalyst preparation

The high-valence cobalt oxide (marked as CoO$_3$) was synthesized first by the precipitation-oxidation method in an aqueous solution. The detailed preparation procedure was described in a previous paper [10]. Furthermore, the pure cobaltic oxide (Co$_3$O$_4$) was refined from CoO by a controlled hydrogen reduction in a temperature-programmed reduction system to $230$ °C (assigned as R230).

The CeO$_2$/Co$_3$O$_4$ catalyst was prepared by impregnation of the Co$_3$O$_4$ support with an aqueous solution of the desired concentration of Ce(NO$_3$)$_3$. The catalyst loading is 20 wt.% and assigned as Ce20/R230. The catalyst was dried at $110$ °C for 24 h, followed by two pretreatment methods: reduction in an H$_2$ stream (under 200 and 400 °C, assigned as Ce20/R230-H200 and Ce20/R230-H400) for 2 h and calcination in an air stream (under 350 and 550 °C, assigned as Ce20/R230-C350 and Ce20/R230-C550) for 4 h.

2.2. Catalyst characterization

X-ray diffraction (XRD) measurements were performed using a Siemens D5000 diffractometer with Cu K$_{α1}$ radiation ($λ = 1.5405$ Å) at 40 kV and 30 mA with a scanning speed in 2θ of 2 ° min$^{-1}$. Diffraction peaks of crystalline phase were compared with those of standard compounds reported in the JCPS data file (Co$_3$O$_4$: 34-0394; CeO$_2$: 09-0418). The crystallite sizes of cobaltic oxide and ceria were estimated using the Scherrer equation.

Nitrogen adsorption isotherms at $-196$ °C were determined volumetrically with Micrometics ASAP 2010. The catalysts were pre-outgassed at $5 \times 10^{-5}$ Torr for 3 h at $110$ °C. The surface area was determined from the nitrogen adsorption isotherm. The infrared spectra were obtained with a Bomen DA-8 spectrometer in the range of 500–800 cm$^{-1}$. One milligram of each powder sample was diluted with 200 mg of vacuum-dried IR-grade KBr and subjected to a pressure of 8 tonnes.

The DRS spectra were measured at room temperature on a Hitachi U3410 spectrophotometer equipped with an integration sphere. The spectra were recorded against a BaSO$_4$ standard in the region 200–800 nm.

Reduction behavior of CeO$_2$/Co$_3$O$_4$ catalysts was studied by temperature-programmed reduction (TPR). About 25 mg of the sample was heated in a flow of 10% H$_2$/He gas mixture at a flow rate of 10 ml min$^{-1}$. During TPR, the temperature was programmed to rise at 10 °C min$^{-1}$ to 550 °C.

2.3. Catalytic activity measurement

The catalytic activity of CeO$_2$/Co$_3$O$_4$ catalysts towards CO oxidation was carried out in a continuous flow micro-reactor. The reaction gas, a mixture of 10% O$_2$/He with 4% CO/He, was fed to a 0.5 g catalyst at a rate of 20 ml min$^{-1}$. Steady-state catalytic activity was measured at each temperature with the reaction temperature raised from room temperature to 200 °C in steps of 25 °C. The effluent gas was analyzed on-line using a Varian 3700 gas chromatograph with a carbosphere column. Before reaction, the catalyst was pretreated in flowing 10% O$_2$/He at $110$ °C for 1 h to drive away molecules pre-adsorbed from the atmosphere.

3. Results and discussion

3.1. Characteristic analysis

The phase composition and the purity are characterized by XRD. Fig. 1 shows the XRD patterns of ceria, cobaltic oxide and the Ce20/R230 catalysts. All samples except R230 (Fig. 1(b)) show a well-crystallized CeO$_2$ phase with a fluorite-type structure. Besides the CeO$_2$ phase, a separate crystalline Co$_3$O$_4$ (Fig. 1(c)–(e)), CoO (Fig. 1(e) and (f)) and Co (Fig. 1(f)) species are found for the Ce20/R230 catalysts. The degree of crystallinity of both phases (Co$_3$O$_4$ and CeO$_2$) increases with increasing calcined temperature (Fig. 1(c) and (d)), while the particle size is similar for samples calcined under 350 and 550 °C (~7.7 nm for Co$_3$O$_4$ and ~6.0 nm for CeO$_2$). Comparison with the results in our previous paper [12], as the cobaltic oxide calcined at 400 °C, the particle size for Co$_3$O$_4$ increases from 8.2 to 15.4 nm. Apparently, the addition of ceria shows good resistance to the sintering of cobaltic oxide under high-temperature due to the strong interaction of cobaltic oxide and ceria (the combined effect can be further characterized by TPR). Also, the degree of crystallinity of mixed phases and CeO$_2$ increases with increasing reduction temperature (Fig. 1(e) and (f)). The particle size of CeO$_2$ is not affected under low or high reduced temperature (~6.0 nm), while the effect of treatment temperature under reduced atmosphere is apparent for both composition and particle size of CoO$_x$. According to the diffraction patterns and the width of the (2 0 0) diffraction pattern of crystalline CoO, the composition of CoO$_x$ in Ce20/R230-H200 (Fig. 1(e)) is a mixed phase [Co$_3$O$_4$ and CoO, although the (3 1 1) peak of Co$_3$O$_4$ overlaps with the (1 1 1) peak of CoO, the faint peak of (3 1 1) diffraction pattern can be further characterized by TPR] and the particle size of CoO is 6.0 nm. As the reducing temperature increases, the composition of Ce20/R230-H400 shows CoO and Co mixed phases and the particle
size of CoO grows to 8.3 nm. The compositions of Ce20/R230 catalysts under different treatments are summarized in the third column of Table 1. Inspection of these results reveals that: (i) the higher the treating temperature, the sharper the diffraction peaks, indicating the degree of crystallinity is progressively growing with temperature; (ii) the slight change of particle size for CoO means that the introducing of ceria can resist the sintering of CoOx.

The surface area (SBET, m² g⁻¹) of Ce20/R230 catalysts is determined from nitrogen adsorption isotherms measured at −196 °C. The data of SBET are given in the second column of Table 1. The results show that the optimized pretreatment can obtain high surface area of CeO2/Co3O4 catalysts. The increase in treating temperature induces a decrease in its surface area (i.e., Ce20/R230-C550 < Ce20/R230-C350; Ce20/R230-H200 < R230 < Ce20/R230-H400). The induced decrease

Table 1
Characterization and catalytic activity for CO oxidation over Ce20/R230 catalysts

<table>
<thead>
<tr>
<th>Catalysts</th>
<th>SBET (m² g⁻¹)</th>
<th>Compositionᵃ</th>
<th>TPR (°C)</th>
<th>CO oxidation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>CeO₂</td>
<td>Ce₂O₃</td>
<td>Co₃O₄</td>
</tr>
<tr>
<td>R230</td>
<td>100</td>
<td></td>
<td>–</td>
<td>280</td>
</tr>
<tr>
<td>CeO₂</td>
<td>64</td>
<td></td>
<td>–</td>
<td>280</td>
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<td>333</td>
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<tr>
<td>Ce₂O₄</td>
<td>40</td>
<td>CeO₂, Co₂O₃, CoO</td>
<td>–</td>
<td>260</td>
</tr>
<tr>
<td>Ce₂O₅</td>
<td>109</td>
<td>CeO₂, Co₂O₃, CoO</td>
<td>–</td>
<td>312</td>
</tr>
<tr>
<td>Ce₂O₆</td>
<td>60</td>
<td>CeO₂</td>
<td>–</td>
<td>333</td>
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<tr>
<td>Ce₂O₇</td>
<td>31</td>
<td>CeO₂, Ce₂O₃</td>
<td>–</td>
<td>260</td>
</tr>
<tr>
<td>Ce₂O₈</td>
<td>40</td>
<td>CeO₂, Co₂O₃, CoO</td>
<td>–</td>
<td>312</td>
</tr>
</tbody>
</table>

ᵃ Characterized from XRD data.
ᵇ Temperature for 50% CO conversion.
due to the thermal treatment might be attributed to grain growth of the particles or collapse of pores [33]. The higher surface area for Ce20/R230-H200 (or Ce20/R230-C350) compared with that for Ce20/R230-H400 (or Ce20/R230-C550) can be understood from the pore size distribution and volume. The results are shown in Fig. 2. The pore size of R230 (Fig. 2(a)) distributes from 2 to 4 nm and from 5 to 18 nm. The pore size of Ce20/R230-H200 (Fig. 2(d)) ranges from 2 to 15 nm and that of Ce20/R230-C350 (Fig. 2(b)) ranges from 5 to 20 nm. The Ce20/R230-H200 possesses larger pore volume than the Ce20/R230-H400. Also, the pore volume of Ce20/R230-C350 is larger than Ce20/R230-C550. The addition of ceria can promote the more uniform pore distribution. Apparently, pores in Ce20/R230-H400 and Ce20/R230-C550 have collapsed. The higher pore volumes of the composites are consistent with the higher surface areas. From the increasing of crystalline and decreasing of surface area of mixed oxides under high-temperature treating conditions, a possible explanation is the collapse of pores.

Fig. 3 shows the IR absorption spectra of ceria, cobaltic oxide and the Ce20/R230 catalysts. All samples except CeO2 and Ce20/R230-H400 (Fig. 3(a) and (f)) show two distinct bands at 575–578 ($\nu_1$) and 660–663 ($\nu_2$) cm$^{-1}$ that are due to the stretching vibrations of the Co–O bond of cobaltic oxide [34,35]. The $\nu_1$ band is characteristic of OB$_{3}^{\alpha}$ (where B$^{\alpha}$ denotes the Co$^{3+}$ in octahedral holes) vibration and the $\nu_2$ band is attributed to the A$^{\alpha}$B$^{\beta}$O$_3$ (where A$^{\alpha}$ denotes the Co$^{2+}$ in tetrahedral holes) vibration in the spinel lattice [36,37]. At the same time, the two bands shift apparently to a lower wavenumber for Ce20/R230-H200 (Fig. 3(e)) due to the mixed phases [11] which indicate a larger number of defects at the surface of smaller nanoparticle (higher surface area) crystals to weaken the Co–O bond strength. The displacement of the bands is in accordance with the structure of CoO$_{x}$ (spinel for Co$_3$O$_4$ and face-centered cubic for CoO).

The DRS spectral features are sensitive to nano-sized oxide particles [38]. The existence of Co$_3$O$_4$ is further confirmed by the DRS spectroscopy. The DRS spectra of ceria, cobaltic oxide and the Ce20/R230 catalysts are shown in Fig. 4. Both maxima characteristic bands at 720 and 420 nm (see Fig. 4(b)–(e)) are due to the formation of Co$_3$O$_4$ [39,40] where a migration to octahedral sites occurs. Herein it is worth noting that these bands become more intense as the surface area of catalyst increases. Pure CeO$_2$ (Fig. 4(a)) shows a broad absorption feature with an
absorption edge around 400 nm characteristic of the semiconducting nature [41,42]. The maximum characteristic band for pure CeO$_2$ or CeO$_2$/Co$_3$O$_4$ mixed oxides at 305 nm (see Fig. 4(a), (c)–(f)) corresponds to the localized O → Ce charge transfer transition involving a number of surface Ce$^{4+}$ ions. Therefore the DRS results are in good agreement with XRD and IR analysis.

In order to understand the relationship of a Co–O bond strength of Ce20/R230 catalysts, we further recognized with TPR technique to understand the reduction behavior of CoO$_x$. Fig. 5 shows the TPR profiles for ceria, cobaltic oxide and the Ce20/R230 catalysts. All the samples except CeO$_2$ and Ce20/R230-H200 (Fig. 5(a) and (g)) exhibit a similar TPR profile, consisting of two well-resolved reduction peaks (assigned as α-peak and β-peak). These profiles point to a two-step reduction process: the first one (α-peak) of low intensity, starts at low-temperature and overlaps with the more intense second one (β-peak). According to the literature [10–12,30,43,44], the low-temperature α-peak can be ascribed to the reduction of Co$^{3+}$ ions, present in the spinel structure, into Co$^{2+}$ (Eq. (1)), with the subsequent structural change to CoO, which followed the higher temperature β-peak and is due to the reduction of CoO to metallic cobalt (Eq. (2)):

$$\text{Co}_3\text{O}_4 + \text{H}_2 \rightarrow 3\text{CoO} + \text{H}_2\text{O}$$

$$\text{CoO} + \text{H}_2 \rightarrow \text{Co} + \text{H}_2\text{O}$$

As can be seen in Fig. 5(b)–(f) and the fifth and sixth columns of Table 1, an apparent shift of the α-peak and β-peak appears under different treatments which are related to the surface areas of composites. Compared to the $S_{\text{BET}}$ (the second column of Table 1), both the α-peak and β-peak shift to lower temperatures as the $S_{\text{BET}}$ increases, i.e., the α-peak and β-peak of Ce20/R230-H200 sample ($S_{\text{BET}} = 109 \text{ m}^2 \text{ g}^{-1}$) are at 260 and 360 °C (Fig. 5(f)), respectively. While, the α-peak and β-peak of Ce20/R230-C550 sample ($S_{\text{BET}} = 31 \text{ m}^2 \text{ g}^{-1}$) are at 333 and 380 °C (Fig. 5(e)), respectively. These results indicate that the increasing of $S_{\text{BET}}$ of Ce20/R230 catalysts can weaken the bond strength of Co–O and can promote more lattice oxygen desorption from Co$_3$O$_4$ to decrease the reduction temperature. Aside from the consecutive reduction of Co$_3$O$_4$, a reduction peak with lower intensity at 503 °C (Fig. 5(a)) is presented for pure CeO$_2$. This peak is attributed to the removal of surface capping oxygen ions during the reduction reaction [45,46]. Moreover, the reduction behaviors of β-peak in Ce20/R230 catalysts (Fig. 5(d)–(f)) and in pure Co$_3$O$_4$ (Fig. 5(b)) are somewhat different. The tailing β-peak in Ce20/R230 catalysts is attributed to the spillover of hydrogen from Co$_3$O$_4$ to further reduce the capping oxygen of ceria, while the physically mixed sample [assigned as Ce20/R230-P and shown in Fig. 5(c)] does not show the tailing β-peak. According to the shift of reduction temperatures and tailing β-peak behavior, the combined effect between the Ce$_2$O$_3$ and Co$_3$O$_4$ is more evidence for the Ce20/R230-H200 sample. Also, it is interesting to compare the reduction behavior of Ce20/R230-H400 and Ce20/R230-H200 catalysts. These are in agreement with the results of XRD analysis: different mixed phases obtained under reducing conditions give various diffraction patterns and reduction behavior. A consecutive reduction for Co$_3$O$_4$ species [that exist in Ce20/R230-H200 sample, see Fig. 5(f) and Fig. 1(e)] and a single peak for CoO at 290 °C [that exist in Ce20/R230-H400 sample, see Fig. 5(g) and Fig. 1(f)] are observed separately. The pre-treatment at 400 °C reduction has removed the capping oxygen of ceria, no tailing peak occurs in Ce20/R230-H400 sample.

3.2. Catalytic activity

In order to assess the catalytic activity of CeO$_2$ on CeO$_2$/Co$_3$O$_4$ catalyst, we studied light-off curves for CO oxidation in a continuous flow micro-reactor. Fig. 6 presents the catalytic activities of ceria, cobaltic oxide and the Ce20/R230 catalysts as a function of reaction temperature. The catalytic activities of pure CeO$_2$ and Co$_3$O$_4$ are also included for comparison. In the reaction temperature ranges (room temperature to 200 °C), except for pure CeO$_2$, the CO conversion over each sample generally increased with the reaction temperature. The activity of pure CeO$_2$ is quite low at temperatures above 200 °C, while the pure Co$_3$O$_4$ and CeO$_2$/Co$_3$O$_4$ mixed oxides possess higher
activities than pure CeO$_2$. A series of different thermal treatments procedure are used in the attempt to determine the optimum pretreatment condition for the catalysts. The $T_{50}$ (the conversion of CO reached 50%) is chosen to judge the catalytic activity for each sample as shown in the last column of Table 1. Clearly, the activity toward CO oxidation is enhanced significantly by increasing the $S_{\text{BET}}$ of catalyst. The higher the $S_{\text{BET}}$ of catalyst, the lower the $T_{50}$ value appears, meaning the higher activity.

Compared to the $S_{\text{BET}}$ (the second column of Table 1) of catalysts, except for pure CeO$_2$, the activity increases significantly by increasing the $S_{\text{BET}}$, i.e., Ce20/R230-H200 ($T_{50} = 88$ °C and $S_{\text{BET}} = 109 \text{ m}^2 \text{ g}^{-1}$) > R230 ($T_{50} = 98$ °C and $S_{\text{BET}} = 100 \text{ m}^2 \text{ g}^{-1}$) > Ce20/R230-C350 ($T_{50} = 112$ °C and $S_{\text{BET}} = 60 \text{ m}^2 \text{ g}^{-1}$) > Ce20/R230-C550 ($T_{50} = 140$ °C and $S_{\text{BET}} = 31 \text{ m}^2 \text{ g}^{-1}$) ~ Ce20/R230-H400 ($T_{50} = 142$ °C and $S_{\text{BET}} = 40 \text{ m}^2 \text{ g}^{-1}$). The best active sample (Ce20/R230-H200) is achieved over the low-temperature reducing pretreatment (reduction under 200 °C), where $T_{50}$ is reached at temperatures as low as 90 °C and 100% CO conversion is reached at about 200 °C. The increase in $T_{50}$ for the Ce20/R230-H400 may mean that cobalt metal covering the surface of CoO decreases the $S_{\text{BET}}$ to depress the CO oxidation activity. Also, in the calcined pretreatment, the low-temperature treating condition can get higher activity.

On the basis of the above results, we suggest that the catalytic activity of Ce20/R230 catalysts is closely related to their surface area. Increasing of the $S_{\text{BET}}$ of CeO$_2$/Co$_3$O$_4$ catalysts weakens the Co–O bond strength and promotes more lattice oxygen desorbed from Co$_3$O$_4$ to increase the CO oxidation activity. In summary, the addition of CeO$_2$ into Co$_3$O$_4$ can sustain a high catalytic activity as long as an optimized pretreatment will maintain high surface area.

4. Conclusion

CeO$_2$/Co$_3$O$_4$ catalysts of different pretreatment conditions have been characterized and tested for CO oxidation. The following conclusions are drawn:

(1) Addition of ceria shows good resistance to the sintering of cobaltic oxide under high-temperature due to the combined effects of cobaltic oxide and ceria.
(2) It is necessary to prepare CeO$_2$/Co$_3$O$_4$ catalysts with larger surface area for a good catalytic performance.
(3) The optimized pretreatment under low-temperature is more suitable to prepare CeO$_2$/Co$_3$O$_4$ catalysts for CO abatement application.
(4) The most active sample is Ce20/R230-H200. The $T_{50}$ for CO oxidation is 88 °C.

Acknowledgement

We are pleased to acknowledge the financial support for this study by the National Science Council of the Republic of China under contract number NSC 93-2113-M-014-003.

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