Ceria-based catalysts for the recovery of elemental sulfur from SO₂-laden gas streams

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Abstract

The catalytic reduction of SO₂ to elemental sulfur by CO or CH₄ over ceria-based catalysts under demanding operating conditions is reported in this paper. Cu- or Ni-containing ceria catalysts have been shown before to be highly active and selective for SO₂ reduction by CO in dry gas streams or in the presence of low amounts of H₂O. In this work, the activity/selectivity of 10 at.% La-doped ceria, Ce(La)Oₓ, and Cu- or Ni-containing Ce(La)Oₓ for the (SO₂ + CO) reaction was tested in gas streams containing 10–45 mol%H₂O at high space velocities (>80,000 h⁻¹). The addition of 5 at.% (~2.5 wt.%) copper or nickel significantly improved the low-temperature (<500°C) reactivity of Ce(La)Oₓ. This was correlated with the improved reducibility of ceria in the metal-ceria catalysts and the ensuing low-temperature activity for the water-gas-shift (WGS) reaction. The combined reduction of SO₂ and NO by CO was also studied in this work over the same catalysts. In dilute gas mixtures containing 0.1–1.0 mol%SO₂ and NO, stoichiometric amount of CO and in the presence of 40%H₂O, the NO presence in the feed gas enhances both the SO₂ conversion and the elemental sulfur yield. At 550°C, in the presence of NO, SO₂ conversion and sulfur yield over the 5%Ni-Ce(La)Oₓ catalyst were 0.94 and 0.77, respectively, the NO conversion to N₂ was complete, and the CO₂ produced was the sum of the SO₂ and NO reduction reactions by CO. Ceria-based materials are also active for SO₂ reduction by methane to elemental sulfur at temperatures higher than 550°C. The addition of Cu or Ni has a different effect on the sulfur selectivity of ceria under fuel-rich conditions. The Cu-ceria system is a complete oxidation catalyst to a much higher temperature than Ni-ceria. Over Ni-CeO₂, dissociation of methane begins at <550°C, and side reactions favor H₂S production over elemental sulfur. However, the 5 at.% Ni-Ce(La)Oₓ catalyst showed remarkable resistance to carbon deposition, both in the SO₂-methane reaction as well as in partial oxidation of methane by O₂ to synthesis gas with a 2:1 H₂:CO ratio. This is attributed to the high dispersion of nickel in this catalyst and the fast rate of oxygen supply from ceria to the nickel interface. Catalysts were characterized by temperature-programmed-reduction, XPS and STEM/EDS. © 2000 Elsevier Science B.V. All rights reserved.

Keywords: Copper oxide; Cerium oxide catalysts; Nickel; Oxygen storage; SO₂ reduction; Sulfur recovery; NO reduction; Elemental sulfur; Water-gas-shift; Methane

1. Introduction

Sulfur dioxide discharged into the atmosphere, primarily as a product of combustion processes is a major atmospheric pollutant causing various ecological damages and health problems. Flue gas desulfurization (FGD) of combustion exhaust gases is presently a costly process often involving complex flowsheets and “throw-away” sorbents [1,2]. Direct catalytic reduction of sulfur dioxide to elemental sulfur by various reductants such as carbon monoxide, methane or synthesis gas is emerging as a promising and cost-effective technological solution for a variety of...
desulfurization problems in chemical, refining, gas processing, metallurgical and power generation industries.

The direct reduction of sulfur dioxide by carbon monoxide to elemental sulfur under dry conditions has been known for a long time [3–5], but no substantial commercial experience exists. The overall reactions involved in this process are as follows:

\[
\begin{align*}
SO_2 + 2CO & \rightarrow 2CO_2 + (1/x)S_x \\
CO + (1/x)S_x & \rightarrow COS \\
2COS + SO_2 & \rightarrow 2CO_2 + (3/x)S_x
\end{align*}
\]

where \(x\) = 2–8 or higher. At high temperatures, predominantly gaseous elemental sulfur, \(S_2\), is produced through reaction 1 which can further react with CO to form carbonyl sulfide, COS. This, in turn may again reduce \(SO_2\) to elemental sulfur through reaction 3. However, COS is a more toxic compound than \(SO_2\) and its production should be minimized in a sulfur recovery process.

Various active catalyst systems have been reported for the \(SO_2\) reduction by CO. Most studied are alumina-supported transition metals and oxides (such as Cu, Fe, Mn, Cr, Ni, Pd, Ag and others) [6,7]. Production of COS usually proceeds to a substantial extent on these catalysts and water partially poisons the catalyst. Lower COS formation was realized on mixed oxides of elements from the lanthanide group and the groups IVA and VA [8] such as La-Ti-O. But this type of catalyst needs to be activated by reducing gases at temperatures above 600°C prior to reaction. ABO\(_3\) perovskite-type mixed oxides [9–13] were reported to have high selectivity to elemental sulfur over COS. The perovskite structure is lost after a short reaction period, but the resultant mixture of sulfides and oxysulfides remains active [14]. Recently, Ma et al. [15–17] reported on the activity of La\(_2\)O\(_2\)S for this reaction as well as for the combined reduction of \(SO_2\) and NO by CO. High selectivity to elemental sulfur was found in the absence of water. Mixed oxides of Co\(_3\)O\(_4\)-TiO\(_2\) were reported by Kim et al. [18] to be active and selective catalysts in dry gas streams.

Cerium oxide-based catalysts containing small amounts (1–5 wt.%) of a transition metal oxide such as Cu, Ni or Co appear to be the most active and selective catalysts disclosed to date for the reduction of \(SO_2\) by CO, both in dry or wet gas streams [19–24]. This type catalyst has excellent redox activity, produces sulfur as a primary product, can handle even trace amounts of \(SO_2\), and can operate at very high space velocities (>80,000 h\(^{-1}\)) at temperatures in the range of 400–500°C. The performance of these catalysts in gases containing excess H\(_2\)O and for the combined reduction of \(SO_2\) and NO is considered in this paper.

The application of ceria-based catalysts to reduction of \(SO_2\) by methane is presented in the second part of the paper. Methane is an attractive reductant due to abundant and cheap natural gas. The stoichiometric overall reaction between \(SO_2\) and CH\(_4\) is

\[
2SO_2 + CH_4 \rightarrow CO_2 + 2H_2O + 2[S]
\]

A number of side reactions and intermediate products are, in general, possible in the \(SO_2 + CH_4\) system. By-products such as H\(_2\)S, COS, CS\(_2\), CO and \(H_2\) can be formed under certain conditions. Catalysts previously studied for this reaction include bauxite [25], alumina [26], aluminum-chromium oxides [27], Fe-Cu-Cr and Cu-Cr-Mn oxides [28] and, more recently, transition metal sulfides: WS\(_2\), FeS\(_2\) and MoS\(_2\) [29–32], and CoO\(_4\)/Al\(_2\)O\(_3\) [33]. However, issues of low activity and/or low selectivity to elemental sulfur were reported with all these catalysts. The optimized ratio of \(SO_2\):CH\(_4\) used by various authors [27,28,32] is typically lower than the stoichiometric ratio of 2:1 (reaction 4) in order to shift the activity to lower temperature, albeit at the expense of selectivity to elemental sulfur. Additionally, under these fuel-rich conditions, undesired by-products such as CO as well as elemental carbon may be produced. Sarlis and Berk [26] reported that CO was formed with the Al\(_2\)O\(_3\)-based catalyst, the rate of CO formation becoming appreciable at \(T > 700°C\) and increasing with the CH\(_4\)/SO\(_2\) ratio. By adding MoO\(_3\), CO production was suppressed, but elemental carbon was deposited on the catalyst [30]. The rate of carbon formation increased with the CH\(_4\)/SO\(_2\) ratio, and became comparable to the rate of CO\(_2\) production at high temperature.

Cerium oxide is well known for its high oxygen storage capacity and oxygen reducibility which make it a most important additive in the three-way catalyst for a variety of redox reactions in the automotive exhaust gas [34]. Previous work in our lab has shown that ceria-based materials are highly active and stable.
catalysts for SO₂ reduction to elemental sulfur by CO at high space velocity [19–24]. Recently, we have found these catalysts to be very attractive also for the reduction of SO₂ by methane [35–37]. The operating temperature is about 200°C higher than that for the reduction of SO₂ by CO. Under fuel-rich conditions, we have reported a different sulfur selectivity between Cu- and Ni-containing ceria. This was attributed to the different activity of these two catalysts for side reactions that occur under fuel-rich conditions and in the presence of water vapor. The Ni-CeO₂ system catalyzes the partial oxidation of methane to synthesis gas [38] while the Cu-CeO₂ system remains a complete oxidation catalyst to a much higher temperature than the former. In this paper, these findings are related to the selectivity of each catalyst to elemental sulfur. The stability of Ni-CeO₂ against carbon deposition in CH₄-rich conditions is also discussed.

2. Experimental

Bulk ceria-based catalysts were prepared by the urea gelation/co-precipitation method using metal nitrates and urea by a procedure described in detail elsewhere [39]. Lanthanum (10 at.%) was typically used as a dopant of ceria because it suppresses the crystal growth and sintering of ceria during the calcination step [39,40]. The doped cerias are denoted as Ce(La)ₙOₙ. Samples were calcined in air for 3 h at 650°C or at 720°C for the (SO₂ + CO) or (SO₂ + CH₄) reaction tests, respectively. The resulting surface area was 90–100 m²/g for the 650°C samples and 45–70 m²/g for the 720°C samples, while the bulk density was 1.8–2 g/cm³.

Bulk elemental analysis was performed by inductively coupled plasma (ICP) atomic emission spectrometry (Perkin-Elmer Plasma 40). The BET surface area was measured in a Micromeritics Pulse Chemisorb 2705 apparatus. X-ray powder diffraction (XRD) analysis of the samples was performed on a Rigaku 300 instrument. Cu Kα radiation was used with a power setting of 60 kV and 300 mA. Tungsten was used as the internal d-spacing standard. The catalyst surface composition was determined by X-ray photo-electron spectroscopy (XPS) on a Perkin-Elmer 5100C system. All measurements were carried out at room temperature and without any sample pre-treatment. A Mg Kα X-ray source typically set at 15 kV and 20 mA was used in this work. All binding energies were adjusted relative to C 1s at 284.6 eV. The catalyst microstructure was studied in a vacuum generators HB 603 scanning transmission electron microscope (STEM) equipped with a X-ray microprobe of 0.14 nm optimum resolution for energy dispersive X-ray spectroscopy (EDS). For STEM analysis, the catalyst powder was dispersed on a nickel or copper grid coated with a carbon film and elemental maps were obtained on a 128 × 128 data matrix.

Catalyst activity tests were conducted in a laboratory-scale, packed-bed flow reactor made of quartz (I.D. = 10 mm) with a porous quartz frit supporting the catalyst. All samples were in powder form (<150 μm). The catalyst loading varied from 150 to 300 mg, and the typical flow rate was 100 cm³/min. The reactor was heated inside a Lindberg electric furnace. The temperature was measured by a quartz tube-sheathed K-type thermocouple placed at the top of the packed bed, and was controlled by a Wizard temperature controller. The reacting gases, all certified calibration gas mixtures with helium (Middlesex), were measured by mass flow controllers and mixed prior to the reactor inlet. The mole percentage of SO₂ in the feed gas was typically unity. Water was injected into a heated gas line with a calibrated syringe pump. The experiments were carried out under nearly atmospheric pressure. A cold trap connected at the outlet of the reactor was used to separate and collect the elemental sulfur and water from the product gas stream. The reactants and products were analyzed by an on-line HP5880A gas chromatograph (GC) equipped with a thermal conductivity detector (TCD). A 1/8 in. O.D. × 6 ft. long Teflon column packed with Porapak QS was employed to separate CH₄, CO, CO₂, COS, CS₂, H₂S and SO₂ while a 1/4 in. O.D. × 6 ft. long Teflon Chromosil 310 column was used to separate CO, CO₂, COS, CS₂, H₂S and SO₂ in the (SO₂ + CO) reaction tests. An additional column, Molecular Sieve 5 Å, was used to separate N₂, NO and CO in tests for combined NO/SO₂ reduction.

The results are shown in terms of sulfur dioxide conversion, X-SO₂, and elemental sulfur yield, Y-[S], or selectivity to elemental sulfur S, defined as follows:

\[ X-\text{SO}_2 = \frac{\text{SO}_2^{\text{in}} - \text{SO}_2^{\text{out}}}{\text{SO}_2^{\text{in}}} \]
\[ Y-[S] = \frac{S_{\text{out}}}{\text{SO}_2^{\text{in}}} \]

S = Y-[S]/X-SO₂

where SO₂ᵢₚ and SO₂ₑₒᵤₜ are mol percentages of SO₂ in the feed gas and effluent gas, respectively, while Sₒᵤₜ is the outlet elemental sulfur concentration. Typically, Sₒᵤₜ was determined by subtracting the mol percentages of H₂S and COS in the effluent gas from SO₂ᵢₚ-SO₂ₑₒᵤₜ.

Temperature-programmed reduction (TPR) of the as-prepared catalysts (720°C-calcined) in powder form (<150 μm) in a H₂/He gas mixture was carried out in a Cahn TG 121 thermogravimetric analyzer (TGA). Each sample was preheated to 500°C in He for 30 min to remove contaminants adsorbed on the sample. After cooling down to room temperature, the samples were heated in a 5%H₂/He mixture (600 cm³/min (STP)) at a heating rate of 10°C/min to 700°C, while monitoring the catalyst weight change in the TGA.

3. Results and discussion

3.1. SO₂ reduction by CO in the presence of excess water

Previous studies of the catalytic reduction of SO₂ by CO to elemental sulfur over ceria carried out in our lab [19–22], have shown that oxidizing agents such as carbon dioxide and water vapor inhibit the reaction of CO with SO₂. The activity is restored only at high temperatures upon desorption of water. An earlier FTIR study had shown that adsorption of CO was hindered on a hydroxylated ceria surface [41]. Liu and coworkers [19–21] first showed that incorporation of small amounts of transition metals, such as Cu, Ni, Co, Pt, and others into ceria enhances the catalytic activity of the latter, apparently by providing active surface sites for CO adsorption. The effect of water vapor content on sulfur yield was reported in an earlier publication on a catalyst with composition 15 at.%Cu-Ce(4.5 at.%La)Oₓ [19]. The sulfur yield was found to decrease from ca. 0.97 to ca. 0.53 as water vapor was raised from 0 to 4 mol%. However, further increasing the water vapor content did not cause a change in sulfur yield. Sulfur yield was maintained at the 0.45 level in the presence of 8 or 13%H₂O [19,21]. Because stoichiometric feed gas (CO:SO₂ = 2) was used, the produced H₂S relative to unreacted SO₂ was always kept at the stoichiometric ratio. Both the SO₂ conversion and sulfur yield were flat at high temperatures.

Ni and Cu are reported to have a similar enhancement effect on the activity of CeO₂ catalysts for a reacting gas mixture comprising no water or small amounts of water [19–21]. Higher levels of water vapor were examined recently [24]. Fig. 1 shows sulfur dioxide conversion and sulfur yield over low-content Cu- or Ni-Ce(La)Oₓ in the presence of 10%H₂O. Compared to Ce(La)Oₓ, addition of 5 at.% (~2.5 wt.%) copper or nickel significantly improved the low-temperature reactivity of the catalyst and its resistance to water vapor poisoning. Both metal-containing ceria catalysts showed similar performance with respect to SO₂ conversion and sulfur yield.

Next, a gas containing a large excess of water (>40%) was examined. In practice, this may be the composition of a regenerator offgas stream diluted with large amounts of process steam prior to SO₂ reduction. Ni-CeOₓ catalysts were found to retain a much higher activity than CeO₂ even in this type of feed gas [24]. Fig. 2 shows that in a reacting gas containing 42.8%H₂O, the SO₂ conversion over the Ni-CeOₓ catalyst is 0.54 at 450°C while it is only about 0.15 over the other ceria catalysts at the same temperature. The Ni-CeOₓ catalyst also gives a higher sulfur yield than the other catalysts which correlates with its higher SO₂ conversion. Such a large difference is believed to result from available CO adsorption sites on the metal that are not poisoned by H₂O. Reaction of CO with H₂O by the water-gas-shift (WGS) reaction may also assist the main reaction. In Fig. 2, all CO was consumed at 500°C over the Ni-ceria catalyst. As shown first by Liu [21], WGS takes place at a much lower temperature on Cu/ceria than on ceria. Metal/ceria catalysts may then be used with a variety of SO₂-laden gas streams, even when a large excess of process steam is added as a diluent.

Fig. 3 shows the CO conversion on ceria-based materials used in the WGS reaction. Cu or Ni-Ce(La)Oₓ are superior WGS catalysts to Ce(La)Oₓ, as indicated by a shift of the reaction light-off temperature to a lower value. This may be attributed to an enhancement of the redox activity of ceria by the presence of the metal [24,42]. We have examined this further by
comparing the reducibility of metal-modified ceria to that of ceria [24,39,40,42]. Similar to what is known for the noble metal/ceria systems [43], the reducibility of Cu- or Ni-CeO$_2$ is higher than that of ceria.

Table 1 lists reduction extents of ceria, defined here as the value of $x$ in Ce(La)$_{1-x}$O$_x$ and Cu- or Ni-Ce(La)$_{1-x}$O$_x$, obtained from H$_2$-TPR. For example, at 500°C, 15%Cu-Ce(La)$_{1-x}$O$_x$ and 5%Ni-Ce(La)$_{1-x}$O$_x$ showed a reduction extent of 1.80 and 1.84, respectively, while this was 1.95 for Ce(La)$_{1-x}$O$_x$. Reduction extents of metal-containing ceria are calculated after accounting for complete reduction of the metal oxide to the metal at each temperature. Enhanced reducibility of ceria in the system Cu-CeO$_2$ was also reported by Rodriguez et al. [44]. Thus, at low temperatures, both the WGS and SO$_2$ reduction reactions are favored on the metal
modified ceria, Figs. 1–3, while at high temperatures, the activities of all ceria-based catalysts converge.

3.1.1. Effect of NO on the SO$_2$ reduction by CO: combined SO$_2$/NO$_x$ reduction

There is ample scope for studying the simultaneous catalytic reduction of SO$_2$ and NO in exhaust gas streams, namely for flue gas cleanup application as well as in the automotive exhaust. In connection with the latter, SO$_2$ is detrimental to the three-way catalyst system [45], making it virtually impossible to attain the ultra-low levels of CO, NO and hydrocarbon emission specified in current regulations.

The effect of NO on the reduction of sulfur dioxide by carbon monoxide was examined in this work under wet conditions (with 42.8% H$_2$O) over the Ni-Ce(La)O$_x$ catalyst. At the same time, the conversion of NO and CO was monitored during these tests. Fig. 4a shows steady-state profiles of sulfur dioxide conversion and selectivity to product sulfur versus temperature over the 5%Ni-Ce(La)O$_x$ catalyst in a feed gas mixture consisting of 1%SO$_2$, 0.5%NO, 2.7%CO, 42.8%H$_2$O, bal. He at a contact time of 0.09 g s/cc (NTP). Results for the reacting gas mixture free of NO are included for comparison. For SO$_2$, reduction by CO to elemental sulfur, addition of NO significantly promotes both the SO$_2$ conversion and selectivity to sulfur. At 550°C, in the presence of 0.5%NO, SO$_2$ conversion and selectivity to sulfur are 0.94 and 0.82, respectively, which corresponds to a sulfur yield of 0.77. This is a very impressive value, considering the high content of water present. More interestingly, as shown in Fig. 4b, the NO conversion to N$_2$ is 100% at temperatures above 500°C. Due to higher conversion of SO$_2$ in the presence of NO, the measured CO$_2$ yield is higher.

In the presence of H$_2$O and NO, various side reactions may take place in the SO$_2$ + CO reaction system. Some of these are as follows:

\[
\begin{align*}
\text{CO} + \text{H}_2\text{O} & \rightarrow \text{CO}_2 + \text{H}_2 & (5) \\
\text{H}_2 + [\text{S}] & \rightarrow \text{H}_2\text{S} & (6) \\
2\text{CO} + 2\text{NO} & \rightarrow 2\text{CO}_2 + \text{N}_2 & (7) \\
\text{CO} + 2\text{NO} & \rightarrow \text{N}_2\text{O} + \text{CO}_2 & (8) \\
2\text{H}_2 + 2\text{NO} & \rightarrow \text{N}_2 + 2\text{H}_2\text{O} & (9) \\
\text{H}_2 + 2\text{NO} & \rightarrow \text{N}_2\text{O} + \text{H}_2\text{O} & (10)
\end{align*}
\]

In addition, CeO$_2$-based systems catalyze the reverse Claus reaction, and the hydrolysis of COS to H$_2$S which is complete above 200°C [21].

Table 2 shows the experimental data collected at the conditions of Fig. 4 over 5 at.%Ni-Ce(La)O$_x$ [24]. As can be seen from the value of $\Delta Y$-CO$_2$, the WGS reaction (5) which is responsible (along with reaction 6) for the reduced SO$_2$ conversion and sulfur yield under wet conditions (Fig. 2), seems to be suppressed in the presence of 0.5%NO in the gas stream. Alternatively, the data in Table 2 may be interpreted as the combination of the WGS and reaction 9, the latter consuming any hydrogen produced by WGS to react with NO...
Fig. 4. Combined SO₂/NO reduction by CO in excess H₂O-containing stream over 5%Ni-Ce(La)Oₓ. (a) SO₂ conversion and sulfur selectivity; (b) NO conversion and CO₂ production; ■ 1%SO₂–2%CO–42.8%H₂O-He, (●) 1%SO₂–0.5%NO–2.7%CO–42.8%H₂O-He, (0.09 g s/cm³ (NTP)).

rather than sulfur by reaction 6. In either case, the selectivity to elemental sulfur would be higher. The data shown in Table 2 are for a concentration of CO slightly above the stoichiometric corresponding to reactions 1 and 7. When excess CO was used in the feed gas, the conversion of SO₂ approached 100% at 550°C, but at a loss of sulfur selectivity [24].

On the basis of Fig. 4 and Table 2, Ni-CeO₂ appears to be a very effective catalyst for the simultaneous reduction of SO₂ and NO by CO to elemental sulfur and N₂, respectively. This catalyst was further investigated with a gas mixture comprising very low concentrations of reactants [24], the results being consistent with the above findings. The simultaneous reduction of SO₂ and NO by CO was also investigated recently by Ma et al. [15–17] over La₂O₃. Higher than 97%SO₂ and NO conversions were obtained over the temperature range from 450 to 600°C. Sulfur yield was above 93%. But, no information about the water effect under these conditions was reported. In sum-
Table 2
Combined SO$_2$/NO reduction over 5%Ni-Ce(La)O$_x$\textsuperscript{a}

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>X-SO$_2$\textsuperscript{b}</th>
<th>X-NO\textsuperscript{c}</th>
<th>Y-CO$_2$\textsuperscript{d} (%)</th>
<th>Y-CO$_2$\textsuperscript{e} (%)</th>
<th>ΔY-CO$_2$\textsuperscript{f} (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>400</td>
<td>0.404</td>
<td>0.62</td>
<td>1.672</td>
<td>1.677</td>
<td>–</td>
</tr>
<tr>
<td>450</td>
<td>0.561</td>
<td>0.8</td>
<td>2.290</td>
<td>2.294</td>
<td>–</td>
</tr>
<tr>
<td>500</td>
<td>0.831</td>
<td>1.0</td>
<td>3.378</td>
<td>3.375</td>
<td>–</td>
</tr>
<tr>
<td>550</td>
<td>0.949</td>
<td>1.0</td>
<td>3.757</td>
<td>3.76</td>
<td>–</td>
</tr>
<tr>
<td>600</td>
<td>0.977</td>
<td>1.0</td>
<td>3.823</td>
<td>3.94</td>
<td>0.117</td>
</tr>
<tr>
<td>650</td>
<td>0.979</td>
<td>1.0</td>
<td>3.827</td>
<td>4.13</td>
<td>0.303</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Reaction conditions: 1%SO$_2$–0.5%NO–2.7%CO–42.8%H$_2$O–He, 0.09 g s/cc.
\textsuperscript{b} Conversion of SO$_2$, measured.
\textsuperscript{c} Conversion of NO, measured.
\textsuperscript{d} Yield of CO$_2$, calculated from the stoichiometric ratio of CO:SO$_2$ = 2:1 and CO:NO = 1:1 in reactions 1 and 7.
\textsuperscript{e} Yield of CO$_2$, measured.
\textsuperscript{f} Difference of d–c is Y-CO$_2$ from water-gas-shift, reaction 5.

mary, Ni-CeO$_x$ appears to be the most active catalyst for reduction of both SO$_2$ and NO by CO. This is a very useful result for NO$_x$ reduction because most NO reduction catalysts are susceptible to sulfur poisoning.

It has been reported that reduced ceria is a good H$_2$S sorbent [46]. A separate experiment was conducted here to confirm that the product of the SO$_2$ reduction was indeed elemental sulfur and not H$_2$S absorbed by the bulk oxide. In this experiment, the reaction was carried out over 5%Ni-Ce(La)O$_x$ at 600°C for 6 h, monitoring the amount of H$_2$S. The feed gas contained 1%SO$_2$, 0.5%NO, 2.5%CO, 42.8%H$_2$O, bal. He, and a flow rate of 100 sccm was used. The catalyst load was 150 mg. Under the assumption that all available CeO$_2$ was fully sulfided to CeS$_2$, it would take 1.86 h to complete the sulfidation. Thus, under ideal absorption conditions where no diffusion limitations are present and for the above (high) sulfur capture by ceria (2S/Ce), H$_2$S breakthrough would occur after ~2 h. However, as shown in Fig. 5, the yield of H$_2$S was constant at ~14% over the 6 h-long test while X-SO$_2$ was 94%. This proves that the main reaction product was indeed elemental sulfur under these conditions. Additionally, this figure demonstrates the good activity/selectivity and stability of the 5%Ni-Ce(La)O$_x$ catalyst in this reaction system.

3.2. SO$_2$ reduction by CH$_4$

3.2.1. Catalyst activity and selectivity

Ceria-based materials are also active catalysts for SO$_2$ reduction to elemental sulfur by methane [35–37]. However, due to the refractory nature of methane, the reaction light-off temperature is about 600°C with a stoichiometric feed gas composition, i.e., ~200°C higher than that of SO$_2$ reduction by CO over the same materials. The CH$_4$:SO$_2$ feed ratio $R$, had a significant effect on the SO$_2$ conversion and sulfur selectivity. Fig. 6 shows the SO$_2$ conversion and sulfur selectivity profiles versus temperature for 5%Cu-Ce(La)O$_x$ at different $R$. The higher the ratio, the higher the SO$_2$ conversion and the lower the catalyst selectivity to elemental sulfur, even though no drop of selectivity was
observed below 650°C. By using excess methane, we can lower the reaction temperature, keeping both high SO₂ conversion and sulfur yield at an optimized level. Complete conversion of SO₂ and 83% sulfur yield were obtained at 675°C over a 5%Cu-Ce(La)Oₓ catalyst at a feed ratio of CH₄:SO₂ = 2 which is four times the stoichiometric ratio for reaction 4. However, undesirable CO as well as H₂S were detected in the product gases at temperatures above 650°C when the value of R exceeded unity.

The addition of 5 at.% Ni or Cu had a negligible effect on the activity of Ce(La)Oₓ as shown in Fig. 7, and reported elsewhere [35–37]. This is contrasted with the reduction of SO₂ by CO where addition of the same amount of these metals in Ce(La)Oₓ has a measurable effect [19–24,35]. This is because the light-off temperature of ceria-based materials for the SO₂ + CH₄ reaction depends on the thermal stability of surface sulfates whose decomposition temperature is ~550°C [23,35,36]. Cu-modified Ce(La)Oₓ...
showed the highest selectivity to elemental sulfur under fuel-rich conditions as shown in Fig. 7. On the other hand, the selectivity of nickel-modified Ce(La)O$_x$ was a complex function of feed ratio and temperature, as will be discussed below.

Under fuel-rich conditions ($R > 0.5$), the measured CO$_2$ product accounts for most of the CH$_4$ consumption while CO was also detected in the product gases. Fig. 8 shows the difference between the CH$_4$ consumed and CO$_2$ measured in the exit gas as a function of temperature for the three catalyst compositions under the conditions of Fig. 7. The measured amount of CO produced was close to the difference between the measured CH$_4$ consumption and CO$_2$ production. The carbon balance was better than 95%. Although we did not directly measure carbon formation, these results indicate that there was no or negligible carbon formation on the catalysts even under fuel-rich conditions. This was confirmed by XPS and other analysis of used samples [47].

Nekrich et al. [28] have proposed the following two parallel overall reactions for the SO$_2$ + CH$_4$ system:

$$\text{CH}_4 + 2\text{SO}_2 = 2\text{S} + \text{CO}_2 + 2\text{H}_2\text{O} \quad (4)$$

$$\text{CH}_4 + \text{SO}_2 = \text{H}_2\text{S} + \text{CO} + \text{H}_2\text{O} \quad (11)$$

The formation of CO and H$_2$S under fuel-rich conditions suggests that reaction 11 also occurs on ceria-based catalysts. Thus, methane is consumed both by complete oxidation by SO$_2$ to CO$_2$ and H$_2$O, and by partial oxidation to CO and H$_2$ products. The H$_2$ produced would further react with surface sulfur to form H$_2$S, resulting in low sulfur yield at temperatures higher than 650°C (Fig. 7).

Ceria-based materials such as Cu-CeO$_2$ and Ni-CeO$_2$ are active for total oxidation of methane at temperatures in the range 350–550°C [21,39,48,49]. Recently, Otsuka et al. [50–52] showed that ceria can also directly convert methane to syngas with H$_2$:CO = 2 at temperatures higher than 600°C. In this work, we examined the activity of Ce(La)O$_x$ in fuel-rich methane oxidation with a CH$_4$:O$_2$ molar feed ratio of 2 in the absence of SO$_2$. In agreement with Otsuka et al. [50–52], partial oxidation products, i.e. CO and H$_2$, were observed on Ce(La)O$_x$ at temperatures higher than 650°C as shown in Fig. 9. However, the activity and selectivity of Ce(La)O$_x$ for partial oxidation of methane (POM) were very low. Addition of 5 at% Ni into Ce(La)O$_x$ caused a dramatic improvement in POM activity (Fig. 9). Oxygen was completely converted at all temperatures tested. The POM light-off temperature is 550°C while only complete oxidation products — CO$_2$ and H$_2$O, were formed at $T < 550°C$. Methane conversion increased with temperature. At 650°C, ~94% methane conversion was measured and the CO yield was 82%, close to the equilibrium value [53] as shown in Fig. 9. The carbon balance was better than 95%. On the other hand, addition of Cu into Ce(La)O$_x$ suppressed the partial oxidation of methane up to 730°C as can be seen in Fig. 9.

The above results confirm that partial oxidation of methane can take place on ceria-based materials to produce synthesis gas, CO and H$_2$. While 5 at.% Ni-Ce(La)O$_x$ is much more active for POM than Ce(La)-O$_x$, the addition of Cu into Ce(La)O$_x$ suppresses the partial oxidation of methane. Thus, Cu-Ce(La)O$_x$ continues to be a complete oxidation catalyst to a much higher temperature (730°C) than Ni-Ce(La)O$_x$. These findings indicate that in the SO$_2$ + CH$_4$ reaction system, SO$_2$ acts as the oxygen source instead of O$_2$, except that the catalyst surface may be partially sulfated during the reaction [37]. Therefore, addition of copper catalyzes reaction 4 rather than reaction 11 which improves sulfur yield. On the other hand, Ni-Ce(La)O$_x$ materials appear to be more active for reaction 11 than Ce(La)O$_x$. 

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**Fig. 8.** Effect of catalyst composition on the difference between methane consumption and CO$_2$ production in 1%SO$_2$–2%CH$_4$–He, 0.18 g s/cm$^2$ (NTP) (■) Ce(La)O$_x$, (●) 5%Cu-Ce(La)O$_x$, (▲) 5%Ni-Ce(La)O$_x$. 

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Fig. 9. Partial oxidation of methane over (□) Ce(La)O\textsubscript{x}, (●) 5\%Cu-Ce(La)O\textsubscript{x}, (▲) 5\%Ni-Ce(La)O\textsubscript{x} catalysts (650°C-calcined) (3\%CH\textsubscript{4}–1.5\%O\textsubscript{2}–He, 0.36 g s/cm\textsuperscript{3} (NTP)); \(X\)-CH\textsubscript{4} = methane conversion, \(S\)-CO = selectivity to CO, \(Y\)-CO\textsubscript{2} = yield of CO/X-CH\textsubscript{4}.

Under methane-rich conditions, carbon formation is a potential problem which will deactivate the catalysts. Claridge et al. [54] observed that Ni had the highest rate of coke formation compared to other metals (Pd, Rh, Ru, Pt, Ir). Rapid carbon deposition on supported Ni catalysts has been a major consideration in studies of the POM reaction. However, the results of the present study suggest that carbon formation on Ce(La)O\textsubscript{x}, 5\%Cu-Ce(La)O\textsubscript{x} and 5\%Ni-Ce(La)O\textsubscript{x} catalysts was negligible under fuel-rich conditions. Since Ni-containing ceria may be more prone to coke formation, the 5\%Ni-Ce(La)O\textsubscript{x} material was further tested in POM at 650°C for 100 h with a CH\textsubscript{4}:O\textsubscript{2} feed ratio of 2 at a contact time of 0.18 g s/cc. No deactivation was observed during 100 h-on-stream as shown in Fig. 10. Methane conversion and CO yield were maintained at \(\sim\)90\% and 82\%, respectively. This catalyst was characterized further as described below.

3.2.2. Catalyst characterization

XPS was used to analyze the surface composition of 5\%Ni-Ce(La)O\textsubscript{x} samples after use in SO\textsubscript{2} + CH\textsubscript{4} reaction with a feed ratio CH\textsubscript{4}:SO\textsubscript{2} = 2 or after 100 h-on-stream at 650°C in POM (CH\textsubscript{4}:O\textsubscript{2} = 2). Fig. 11 shows the O1s and C1s spectra of these samples and as-prepared 5\%Ni-CeO\textsubscript{2} for comparison. Two O1s peaks were found in the XP spectra of the used catalysts. The high binding energy peak at \(\sim\)531.8 eV can be assigned to sulfate or carbonate while the low binding energy peak at \(\sim\)529.4 eV is assigned to metal oxide [55]. The C1s spectra show a prominent peak at a binding energy of 284.6 eV which is assigned to residual hydrocarbons. A small peak at 288.5 eV which was also observed on the fresh materials, may be due to residual surface CO\textsubscript{3}\textsuperscript{2–}. There was no peak due to graphitic or carbide surface carbon species whose binding energy is at \(\sim\)283 eV [55]. The deconvolution of the C 1s signal of used 5\%Ni-Ce(La)O\textsubscript{x} catalysts gave a dominant peak of full width at half height.
Fig. 11. XPS of (1) as-prepared 5%Ni-CeO$_2$, (2) used 5%Ni-Ce(La)O$_x$ in 1%SO$_2$–2%CH$_4$–He, (3) used 5%Ni-Ce(La)O$_x$ at 650°C for 100 h for POM.

Fig. 12. STEM/EDS elemental mapping of 5%Ni-Ce(La)O$_x$, as prepared; calcined at 720°C.
FWHH/D: 8 eV at 284.6 eV and a component at 283.03 eV with 3–8% of the total carbon signal, close to that on the fresh material. Therefore, XPS found no carbon deposited on the surface of 5%Ni-Ce(La)Oₓ after use in fuel-rich conditions. Furthermore, for 5%Ni-Ce(La)Oₓ after 100 h-on-stream (see Fig. 10), the weight loss in temperature-programmed oxidation, carried out in a thermogravimetric analyzer over the temperature range of 25–800°C was negligible (0.13 wt.%/100 h) [38,47]. Therefore, the low-content Ni-ceria catalyst, 5 at.%Ni-Ce(La)Oₓ, displays a high resistance to carbon deposition.

The high carbon resistance of 5 at.%Ni-Ce(La)Oₓ might be attributed to its high dispersion of nickel and high oxygen mobility. In general, a high degree of metal dispersion may reduce coke formation [56]. Fig. 12 shows the STEM/EDS elemental mapping of the as prepared 5%Ni-Ce(La)Oₓ. Nickel was highly dispersed in the ceria matrix. Also, XPS analysis found no surface enrichment of nickel in Ni-Ce(La)Oₓ with a nickel loading up to 10 at.%, suggesting a homogeneous distribution of nickel throughout ceria. A similar conclusion has been reached for the perovskite-supported Ni catalysts [57,58] and rare earth metal oxide (La, Ce, Y) modified Ni/Al₂O₃ catalysts [59–61]. The enhanced resistance to carbon deposition was attributed to improved dispersion of the active component nickel.

The high oxygen storage capacity and oxygen ion mobility of ceria may also contribute to the high carbon resistance of 5 at.%Ni-Ce(La)Oₓ. A high rate of oxygen supply to the nickel interface may speed up the surface oxidation of carbon species, thus inhibiting the deposition of carbon on the catalyst surface. A similar reason may be applied to perovskite-supported nickel [57,58] as perovskite-type oxides are well known for their high oxygen mobility. For the modified Ni/Al₂O₃ catalyst [59–61], addition of cerium can modify the surface oxygen properties of aluminum oxide, thus, providing a higher rate of surface oxidation of carbon species and improving this material’s resistance to carbon deposition.

4. Conclusions

Cu- or Ni-modified ceria are active and selective catalysts for SO₂ reduction by CO to elemental sulfur under a variety of operating conditions. The addition of 5 at.% (~2.5 wt.%) copper or nickel significantly improves the low-temperature reactivity of ceria and its resistance to water vapor poisoning up to very high levels of water (>40%). Such difference is believed to result from available CO adsorption sites on the metal that are not poisoned by H₂O. The metal-ceria catalysts also show high activity for low-temperature WGS reaction. Both the CO-SO₂ and CO-H₂O reactions are favored over Ni- or Cu-containing ceria due to the enhanced redox activity of ceria in the presence of the metal.

The combined reduction of SO₂ and NO by CO was studied in this work over the ceria-based catalysts. In dilute gas mixtures containing 0.1–1.0 mol%SO₂ and NO, stoichiometric amount of CO and in the presence of 40%H₂O, the NO presence in the feed gas enhances both the SO₂ conversion and the elemental sulfur yield. The Ni-CeO₂ appears to be the most active catalyst for the combined reduction of SO₂ and NO by CO. This is a very useful result for NO reduction because most NO reduction catalysts are susceptible to sulfur poisoning.

Ceria-based materials are also active for SO₂ reduction by methane to elemental sulfur at temperatures higher than 550°C. The addition of Cu or Ni into ceria has different effect on the sulfur selectivity of the catalyst under fuel-rich conditions. The Cu-ceria system is a complete oxidation catalyst to a much higher temperature than Ni-ceria. Over Ni-CeO₂, dissociation of methane begins at <550°C, and side reactions favor H₂S production over elemental sulfur. However, the 5 at.%Ni-Ce(La)Oₓ catalyst showed a remarkable resistance to carbon deposition, both in the SO₂-methane reaction as well as in partial oxidation of methane by O₂ to synthesis gas with a 2:1 H₂:CO ratio. This is attributed to high dispersion of nickel and the fast rate of oxygen supply from ceria to the nickel interface.

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