Abstract

We studied the mechanism of NO reduction as well as its selectivity and reactivity in the presence of excess O₂. Results show that fuel injection and/or pretreatment are important for ceria catalyst reduction and carbon deposition on the catalyst surface. Oxygen defects of reduced ceria are the key sites for the reduction of NO into N₂. The deposited carbon acts as a buffer reductant, i.e., the oxidation of carbon by lattice oxygen recreates oxygen defects to extend the NO reduction time interval. A small amount of NO showed a full conversion into only N₂ both on the reduced Zr-La doped ceria and reduced Pt-Zr-La doped ceria. Only when the catalyst is oxidised NO is converted into NO₂.

1. Introduction

A mandatory introduction of a CO₂ emission target of 95 g/km by the year 2020 in the EU for automotive manufactures drives the development of increasingly fuel efficient cars. The requirement of the current Euro 6 emission standard has led to the development of highly efficient lean burn turbo-charged engines and catalytic deNOₓ systems (Lean NOₓ Traps (NSR) for gasoline engines and Urea-Selective Catalytic reduction (SCR) and or a combination of SCR and NSR for diesel engines). Euro 7 (or EURO 6B and 6C) requires a further reduction of the NOₓ emissions from 0.18 g/km (Euro 5) via 0.08 g/km (Euro 6) to 0.04 g/km, while particulate matter emissions remain at 0.0045 g/km. Currently, the amount of NO emission, tested in the laboratory, does not reflect the amount emitted during the real driving conditions [1-3]. As of September 2017, the European Commission will proclaim that the real driving emission (RDE) will partially replace the current laboratory test in order to introduce new car models into the market [4]. These new regulation means that the tested cars will be driven outside on a real road under conditions of acceleration and deceleration that are beyond the current laboratory testing procedures. The pollutant emissions will be measured by portable emission measuring systems (PEMS). The RDE testing is expected to reduce the current difference of the pollutant emissions between laboratory and road.

Up to 2.1 times NOₓ emission (0.168 g/km), relatively to the current Euro 6 NOₓ emission standard (0.08 g/km), is allowed under the RDE testing by September 2017 [4]. This highly indicated that current available technologies: Three-way catalyst (TWC) [5-7], Urea-SCR (Selective Catalytic Reduction) [8-10], Lean NOₓ Traps (NSR) [11-13] and combination thereof still need to be significantly improved since almost all Euro 6 certified diesel cars in the RDE testing emitted on average 8 times higher than the current NOx emission standard [1-3]. In future, the NOₓ emission will become more and more stringent. Therefore, effective exhaust emissions after-treatment technologies will be needed.

Recently, Bisaiji et al. (Toyota Motor Company) developed the Di-Air system in which short fuel rich and long fuel lean periods are created by the directly injecting hydrocarbons (HC’s) at a high frequency downstream of the engine in the exhaust system upstream of a NSR catalyst (Pt/Rh/Ba/K/Ce/Al₂O₃) [14]. The Di-Air system is promising to meet the future NOx emission standards under real driving test conditions. Compared to the current NSR system’s narrow operating temperature window, caused by the requirement to store the NOₓ as nitrite/nitrate on the Ba component, the Di-Air system retains a high NO conversion (above 80%) up to 800 °C at a high space velocity up to 1.2·10⁵ 1/l/h. They found that HC’s injection was more effective in the NO conversion in comparison to that of CO and H₂ [14, 15]. The unique performance of the Di-Air system was attributed to the formation of stable isocyanate (CNO) and isocyanide (CN) intermediates on the catalyst surface as being evidenced by FTIR observations [16]. However, CN and CNO species were observed at a low temperature (200 °C) and their intensities decreased as the temperature increased. The CN and CNO species were hardly detected at 600 °C. The high NO reduction efficiency at high temperatures in Di-Air system as reported can be difficult to relate to these CN and CNO species.

A different explanation can be conceived to explain the performance of the Di-Air system. In the 1990’s Illán-Gómez et al. [17, 18] reported that activated carbon is active in NOx reduction in a temperature window between 400 and 500 °C. A large fuel injection upstream of a catalyst bed might promote the carbon formation and this in turn might enhance the NOₓ reduction. Another explanation can be that the reduced catalyst sites reduce NOₓ directly, while the formed adsorbed hydrocarbon fragments and carbon deposits act as an oxygen scavenger under lean conditions.

The working principle of the Di-Air system has not been clearly illustrated by the inventor (Toyota), academia, or industry, making it difficult to further optimise the system for real application due to the lack of fundamental understanding and reasoning. In order to test the aforementioned alternative explanations, we initiated a detailed investigation into the operating principle of the Di-Air. The TAP (Temporal Analysis of Products) technique (a vacuum pulse-response technique) was mainly utilised to investigate the mechanism of NO reduction and the role of each Di-Air catalyst component (Pt/Rh/Ba/K/Ce/Al₂O₃) [19]. In addition, a flow reactor (under an
5% of O2 (with He for balance) was used with a space velocity of NO in N2, flow rate of 200 mL/min) was admitted to the cell. In order to mimic fuel injection in the Di-Air system, C3H6 is used as a model fuel to pre-treat the catalyst. Ceria was pre-treated (reduced) by C3H6 (1000 ppm in N2, flow rate 200 mL/min) for 2 h. N2 was used to flush the cell for 20 min. Subsequently, NO (1000 ppm NO in N2, flow rate of 200 mL/min) was admitted until they leave the packed bed. The reactant and product(s) are recorded versus time (response) by a mass spectrometer, details of this set-up can be found elsewhere [19]. In the TAP experiments with 15NO and 18O2 pulses: 1) the catalyst was firstly oxidised using an O2 pulse train; 2) secondly, a pre-reduction step with C3H6 pulses was performed until the effluent of the reactor remained constant.

In-situ Raman spectra (Renishaw, 2000) were recorded using a temperature controlled in-situ Raman cell (Linkam, THMS 600). Ten scans were collected for each spectrum in the 100-4000 cm\(^{-1}\) range using a continuous grating mode with a resolution of 4 cm\(^{-1}\) and scan time of 10 s. The spectrometer was calibrated daily using a silicon standard with a strong band at 520 cm\(^{-1}\). The ceria was firstly pre-treated by C3H6 (1000 ppm in N2, flow rate 200 mL/min) for 2 h. N2 was used to flush the cell for 20 min. Subsequently, NO (1000 ppm NO in N2, flow rate of 200 mL/min) was admitted to the cell.

A flow reactor was used to explore the reactivity and selectivity of NO in O2. 200 mg catalyst was placed in a 6 mm inner-diameter quartz reactor tube. A feed composition of 0.2% or 0.05% of NO and 5% of O2 (with He for balance) was used with a space velocity of 6.7·10\(^4\) 1/h. The reactor effluent was online analysed by mass spectrometry (MS, Hiden Analytical, HPR-20 QIC) and infrared (IR) spectroscopy (Perkin–Elmer, Spectrum One). For IR analysis a gas cell with KBr windows with a path length of ∼5 cm was used. The spectra were measured in a continuous mode using the Perkin-Elmer ‘Time-Base’ software between 4000-700 cm\(^{-1}\) wavenumbers with a spectral resolution of 8 cm\(^{-1}\) and an acquisition of 8 scans per spectrum, resulting in a time interval of 23 s between each displayed spectrum.

3. Results

3.1 Mechanism study

3.1.1 15NO and 18O2 pulses over C3H6 reduced ceria in TAP

In order to mimic fuel injection in the Di-Air system, C3H6 is used as a model fuel to pre-treat the catalyst. Ceria was pre-treated (reduced) by C3H6 at 560 °C before 15NO pulse experiment.

Figure 1. Reactant and product evolution upon pulsing (A) 15NO at 560 °C and (B) 18O2 at 530 °C over by C3H6 reduced ceria (21.2 mg) in TAP.

The pre-treatment of C3H6 resulted in 5.2·10\(^{17}\) oxygen atoms / mg Cat extraction from the ceria lattice and 1.6·10\(^{18}\) carbon atoms / mg Cat deposition on the ceria surface. Figure 1A showed the result of 15NO pulses over C3H6 reduced ceria at 560 °C. The discontinuities in Figure 1A were attributed to the refilling of the 15NO buffer vessel in the TAP. From 0 to 5340 pulses (Figure 1A), a full NO conversion was observed with 15N2 as the main product (hardly any CO was formed). The evolution of CO2 was observed from pulse number 5340 onwards. The evolution of CO2 was closely followed by a temporary decrease in NO conversion. The NO conversion increased again from pulse number 8900. From pulse number 20000 to 35000, a full NO conversion was once again observed, while 15N2 and CO2 were the exclusive products. From pulse number 35000 till the end of the experiment, a progressive decrease to zero in NO conversion was observed. The N2 and CO2 production showed the same trend as the NO conversion. No or hardly any 15N2O or 15NO2 were formed and no traces of -CN or -CNO were detected. In order to identify the oxygen species participating in the oxidation of deposited carbon after C3H6 pre-treatment, an 15O2 pulse experiment at over 530 °C C3H6 reduced ceria was carried out. The pre-treatment of C3H6 led to approximately 2 hypothetical reduced ceria layers and 2-3 wt.% carbon deposition. Figure 1B showed the product evolution during the 15O2 pulse experiment. The discontinuities in Figure 1B were at points where the 15O2/He buffer vessel in the TAP was refilled. All introduced 15O2 were completely converted until a steep oxygen breakthrough profile was observed from pulse number 35000. Exclusive C15O2 was evolved until pulse number 4000, where C16O2 started to form, and gradually became the dominant product. After
14000 pulses C\textsuperscript{18}O\textsubscript{16}O and C\textsuperscript{18}O\textsubscript{2} were observed. After 35000 pulses C\textsuperscript{16}O, C\textsuperscript{16}O\textsubscript{2}, C\textsuperscript{12}O\textsuperscript{16}O, C\textsuperscript{18}O, and C\textsuperscript{18}O\textsubscript{2} decreased to zero, where \textsuperscript{18}O\textsubscript{2}, \textsuperscript{16}O\textsubscript{2}, and \textsuperscript{18}O\textsubscript{16}O started to breakthrough in a kind of exchange equilibrium. The observed C\textsuperscript{18}O (m/e=30) was due to the fragmentation from C\textsuperscript{18}O\textsubscript{2} and C\textsuperscript{18}O\textsubscript{16}O and showed the same trend as the C\textsuperscript{18}O\textsubscript{2} and C\textsuperscript{18}O\textsubscript{16}O products. \textsuperscript{16}O\textsubscript{16}O was the main product after oxygen breakthrough.

3.1.2 \textit{In-situ} Raman

![In-situ Raman response for NO over at 560 °C C\textsubscript{3}H\textsubscript{6} reduced ceria. Raman spectra of band at (A) 460 cm\textsuperscript{-1} and (B) 1575 and 1350 cm\textsuperscript{-1}; (C) Raman bands intensity of 460, 1575, and 1350 cm\textsuperscript{-1} responses versus time.](image)

Figure 2 showed \textit{in-situ} Raman spectra for the NO reduction over at 560 °C C\textsubscript{3}H\textsubscript{6} reduced ceria. The band at 460 cm\textsuperscript{-1} was attributed to the symmetric stretch mode of Ce-O\textsubscript{6} crystal unit, which was characteristic for the fluorite ceria structure [20]. The bands at 1575 and 1350 cm\textsuperscript{-1} were assigned to G band and D band of carbon in the form of graphene or graphite, respectively [21]. The G band was usually assigned to zone centre phonons of E\textsubscript{2g} symmetry of the perfect graphite structure, and the D peak was a breathing mode of A\textsubscript{1g} symmetry which was forbidden in a perfect graphite structure and only became active in the presence of structural defects and disorders. The band at 460 cm\textsuperscript{-1} was initially hardly visible and its intensity increased during the NO flow (Figure 2A and C). In contrast to the band at 460 cm\textsuperscript{-1} (Figure 2A and C), the band at 1575 and 1350 cm\textsuperscript{-1} had the same intensities during the NO flow and started to decrease after 200 min (Figure 2B and C).

3.1.3 \textsuperscript{15}NO over C\textsubscript{3}H\textsubscript{6} reduced Pt/ceria in TAP

Similarly to ceria, Pt/ceria was also pre-treated by C\textsubscript{3}H\textsubscript{6}. Instead of 560 °C, Pt/ceria pre-treatment with C\textsubscript{3}H\textsubscript{6} was carried out at 450 °C, which was 110 °C lower than those of the experiments performed over bare ceria (Figure 1 and 2). The pre-treatment of C\textsubscript{3}H\textsubscript{6} resulted in 6.3·10\textsuperscript{17} oxygen atoms / mg\textsubscript{Cat} extraction from the ceria lattice and 2.8·10\textsuperscript{17} carbon atoms / mg\textsubscript{Cat} deposition on the catalyst surface. Figure 3 showed the result of \textsuperscript{15}NO pulse over at 450 °C C\textsubscript{3}H\textsubscript{6} reduced Pt/ceria. From pulse number 0 to 1000 (Figure 3), full NO conversion was observed with \textsuperscript{15}N\textsubscript{2} as the main product (small amount of CO formed). The evolution of CO\textsubscript{2} was observed from pulse number 1000. Instead of a temporary decrease in NO conversion, NO showed full conversion until pulse number 4000, where \textsuperscript{15}N\textsubscript{2} and CO\textsubscript{2} were the only products. From pulse number 4000 onwards, a progressive decrease to zero for the NO conversion was observed. The N\textsubscript{2} and CO\textsubscript{2} production followed the same trend as the NO conversion. No or hardly any \textsuperscript{15}N\textsubscript{2}O or \textsuperscript{15}NO\textsubscript{2} were formed and no traces of -CN or -CNO were detected.

![Reactant and product evolution upon pulsing \textsuperscript{15}NO over at 450 °C C\textsubscript{3}H\textsubscript{6} reduced Pt/ceria (10 mg).](image)

Figure 3. Reactant and product evolution upon pulsing \textsuperscript{15}NO over at 450 °C C\textsubscript{3}H\textsubscript{6} reduced Pt/ceria (10 mg).

3.2 NO reduction in O\textsubscript{2} over reduced ceria

Similarly to TAP experiments, C\textsubscript{3}H\textsubscript{6} was also used as a model fuel in the flow reactor. The ceria was pre-treated with C\textsubscript{3}H\textsubscript{6} (1.25% in N\textsubscript{2}) for 2 h at 600 °C and subsequently (0.2% or 0.05%) NO and 5% O\textsubscript{2} (balance He) was admitted with a flow rate of 200 mL/min. The pre-
treatment of C\textsubscript{3}H\textsubscript{6} resulted in \(7.9 \times 10^{17}\) oxygen atoms / mg \textit{Cat} extraction from the ceria lattice and \(2 \times 10^{18}\) carbon atoms / mg \textit{Cat} deposition on the ceria surface. Figure 4A showed the results of 0.2\% NO + 5\% O\textsubscript{2} over the reduced ceria. m/e= 28 and 44 were observed during the first 90 s. m/e=28 could be attributed to the CO and N\textsubscript{2}. The formation of CO was confirmed from the IR spectra (Figure 5). N\textsubscript{2}, however, was not able to be detected due to its IR inactivity. To prove the formation of N\textsubscript{2}, 0.2\% \textsuperscript{15}NO + 5\% O\textsubscript{2} over the reduced ceria were performed (not shown). The observation of \textsuperscript{15}N\textsubscript{2} (m/e=30) proved that the reduction of NO into N\textsubscript{2}. m/e of 44 could be attributed to N\textsubscript{2}O and CO\textsubscript{2}. From the IR spectra (Figure 5), no N\textsubscript{2}O was detected, indicating that the m/e=44 was originated from CO\textsubscript{2}. Initially full NO and O\textsubscript{2} conversion and both started to breakthrough at 75 s and became stable after 100 s.

\[\text{Figure 4. Reactant and product evolution in flow reactor for 0.2\% NO + 5\% O\textsubscript{2} over reduced ceria; inset is NO evolution.}\]

Similarly to 0.2\% NO + 5\% O\textsubscript{2} (Figure 4A), 0.05\% NO + 5\% O\textsubscript{2} over the reduced ceria \textit{C\textsubscript{3}H\textsubscript{6}} showed the formation of N\textsubscript{2}, CO, and CO\textsubscript{2}. Both NO and O\textsubscript{2} started to breakthrough at 75 s. Figure 5 showed the IR spectra responses for 0.2\% NO + 5\% O\textsubscript{2} in helium over reduced ceria. The interval of IR spectra is 23 s. Initially peaks at 2174 and 2116 cm\textsuperscript{-1}, assigned to CO, were observed and these peaks vanished after 138 s. The broad band at 2350 cm\textsuperscript{-1} was assigned to CO\textsubscript{2} and followed the same trend in time as CO response. The peaks at 1601 and 1628 cm\textsuperscript{-1}, appearing from 116 s onwards, were attributed to the formation of NO\textsubscript{2}. During the whole experiment, no N\textsubscript{2}O was detected (IR peak intensity 2235 cm\textsuperscript{-1}, detection limit 1 ppm). Same IR responses were found for 0.05\% NO + 5\% O\textsubscript{2} experiment, both NO and NO\textsubscript{2} were observed from 93 s.

\[\text{Figure 5. IR spectra of 0.2\% NO + 5\% O\textsubscript{2} over reduced ceria in flow reactor.}\]

\[\text{Figure 6. Reactant and product evolution in flow reactor for 0.2\% NO + 5\% O\textsubscript{2} over reduced Pt/ceria; inset is NO evolution.}\]

\[\text{Figure 6 showed the results of 0.2\% NO + 5\% O\textsubscript{2} over the reduced Pt/ceria. m/e= 28 and 44 were observed during the first 150 s. m/e=28 could be attributed to the CO and N\textsubscript{2}. The formation of CO was confirmed from the IR spectra (Figure 7). N\textsubscript{2}, however, was not able to be detected due to its IR inactivity. To prove the formation of N\textsubscript{2}, 0.2\% \textsuperscript{15}NO + 5\% O\textsubscript{2} over the reduced ceria were performed (not shown). The observation of \textsuperscript{15}N\textsubscript{2} (m/e=30) proved that the reduction of NO into N\textsubscript{2}. m/e of 44 could be attributed to N\textsubscript{2}O and CO\textsubscript{2}. From the IR spectra (Figure 5), no N\textsubscript{2}O was detected, indicating that the m/e=44 originated from CO\textsubscript{2}. NO showed full conversion until 55 s and reached a stable signal from 150 s.}\]

\[\text{Figure 7. IR spectra of 0.2\% NO + 5\% O\textsubscript{2} over reduced Pt/ceria in flow reactor.}\]

\[\text{3.3 NO reduction in O\textsubscript{2} over reduced Pt/ceria}\]

Similarly to ceria, the Pt/ceria was pre-treated with \textit{C\textsubscript{3}H\textsubscript{6}} (1.25\% in N\textsubscript{2}) for 2 h at 600 °C and subsequently (0.2\% or 0.05\%) NO and 5\% O\textsubscript{2} (balance He) was admitted with a flow rate of 200 mL/min. The pre-treatment of \textit{C\textsubscript{3}H\textsubscript{6}} resulted in \(9.8 \times 10^{17}\) oxygen atoms / mg \textit{Cat} extraction from the ceria lattice and \(1.7 \times 10^{19}\) carbon atoms / mg \textit{Cat} deposition on the catalyst surface.
Similarly to 0.2% NO+ 5% O₂ (Figure 4), 0.05% NO + 5% O₂ over the reduced Pt/ceria C₃H₆ showed the formation of N₂, CO, and CO₂. Both NO and O₂ started to breakthrough at 55 s and reached stable signals from 150 s and 100 s, respectively.

Figure 7 showed the IR spectra responses for 0.2% NO + 5% O₂ in helium over reduced Pt/ceria. The interval of IR spectra is 23 s. Initially peaks at 2174 and 2116 cm⁻¹, assigned to CO₂ broad band at 2350 cm⁻¹, assigned to CO₂ were observed and these peaks vanished after 105 s. The peaks at 1601 and 1628 cm⁻¹, appearing from 85 s onwards, were attributed to the formation of NO₂. During the whole experiment, no N₂O was detected (IR peak intensity 2235 cm⁻¹, detection limit 1 ppm). Same IR responses were found for 0.05% NO + 5% O₂ experiment, both NO and N₂O were observed from 93 s. Same IR responses were found for 0.05% NO + 5% O₂ experiment, both NO and N₂O were observed from 105 s.

4. Discussion

4.1 Mechanism study

High frequency fuel injection was applied in the Di-Air system. To mimic this Di-Air system in the present study, the catalyst was pre-treated by C₃H₆ before NO reduction experiments. In order to distinguish N₂ and CO (m/e= 28) as well as CO₂ and N₂O (m/e=44), ¹⁵NO was used to replace ¹⁵NO.

Starting with the ceria, full NO conversion was observed from pulse number 0 to 5000 with exclusive formation of ¹⁵N₂ as shown in Figure 1A. The absence of oxidation products CO and CO₂ before pulse number 5000 indicated that the carbonaceous residues, left on the surface after C₃H₆ pre-reduction, did not directly participate in the reduction of NO to N₂. A mechanism of direct dissociation of adsorbed NO on oxygen anion defect centres, in which the O side of NO filled the oxygen anion vacancy, and the N species recombined to form N₂ can explain the observation.

NO dissociated on oxygen anion defect centers of ceria, which was also found for the H₂ and CO reduction pre-treatment [19, 22]. NO dissociation resulted in a progressive re-oxidation of the catalyst with increasing NO pulse number, thereby decreasing the number of oxygen anion defect sites. When ceria is partially re-oxidised, CO₂ formation is observed, indicating that ceria became active in the oxidation of these deposited carbonaceous residues at a specific ceria oxidation state. The oxidation of carbonaceous deposits to CO₂ molecule created two oxygen anion defect centres, forming one N₂ molecule from 2 NO molecules. This delayed CO₂ formation will inhibited the refilling of oxygen anion defect centers and cause a temporary decline in the NO conversion.

To support the role of lattice oxygen in the oxidation of deposited carbon, i.e. gas-phase NO did not directly participate in the carbon deposit oxidation reaction, ¹⁶O₂ was pulsed over at 530 °C C₃H₆ reduced ceria, as shown in Figure 1B. The initial exclusive formation of C¹⁶O and C¹⁶O₂ indicated that gas-phase oxygen was not directly involved in the oxidation of the carbon deposits, which was in agreement with previous findings in diesel soot on ceria based catalysts [23]. In summary, NO re-oxidised reduced ceria and was not directly involved in carbonaceous deposits oxidation. The oxidation of carbonaceous deposits to CO₂ by oxygen species originating from the lattice oxygen, will recreate oxygen anion defect centers, which were accountable for additional NO dissociation. Therefore, these carbonaceous deposits could be seen as a delayed or stored reductant.

Since TAP is an ultra-vacuum technique, in-situ Raman (at atmospheric pressure) was applied to confirm the results obtained from TAP. NO reduction into N₂ was performed over at 560 °C C₃H₆ reduced ceria, as shown in Figure 2. The band at 460 cm⁻¹ (Figure 2A) was attributed to the symmetric stretch mode of the Ce-O₈ crystal unit, which was characteristic for reduced fluorite ceria structure [20]. This peak disappeared during the C₃H₆ pre-treatment, while it re-appeared under NO. This indicated that the reduced ceria was re-oxidised during NO flow. The bands at 1575 and 1350 cm⁻¹ were assigned to G band and D band of carbon in the form of graphene or graphite (Figure 2B) and they remained constant during the first 200 min of NO in N₂ flow (Figure 2C). The change of the bands’ intensity during NO flow indicated that the oxidation of carbon commenced later than the re-oxidation of ceria. This confirmed the TAP’s finding: NO firstly re-oxidised oxygen defects in the reduced ceria and subsequently the oxidation of carbon started when ceria was largely oxidised.

Ceria is a vital ingredient in the Di-air catalyst and oxidises hydrocarbons and reduces NO into N₂. However, high temperatures (above 500 °C) were necessary to oxidise C₁₆O/C₁₆O₂. To successfully use ceria based catalysts, promoters might be beneficial to convert hydrocarbons at lower temperatures, thereby, a (deeper and faster) reduction of the ceria with a larger carbon deposits pool.

To lower the hydrocarbon activation, Pt was loaded on ceria and it significantly increased the C₃H₆ reactivity. The reduction of Pt/ceria by C₃H₆ at 450 °C led to 6.3-10¹⁷ oxygen atoms/mg Cat of extraction from ceria lattice, which was around 1-10¹⁷ oxygen atoms/mg Cat more than the reduction of ceria at 560 °C. However, the amount of deposited carbon was around 2.8-10¹⁸ carbon atoms/mg Cat, which was around 10 times less than that over ceria pre-treated by C₃H₆ at 560 °C. Although less carbon atoms were deposited over the Pt/ceria at 450 °C, Pt/ceria was still active in C₃H₆ oxidation and ceria reduction, since bare ceria was hardly active in C₃H₆ oxidation at 450 °C.

Figure 3 showed the ¹⁵NO pulses over Pt/ceria pre-treated by C₃H₆ at 450 °C. The absence of oxidation products CO and CO₂ before pulse number 1000, indicated that the carbonaceous residues, left on the surface after C₃H₆ pre-reduction, did not directly participate in the reduction of ¹⁵NO into ¹⁵N₂. Carbon oxidation started from pulse number 1000. This indicated that the presence of Pt did not change the principle of NO reduction into N₂. Oxygen defects were still the key sites for NO reduction into N₂ and the deposited carbon acted as a buffer reductant.

4.2 Competition of NO with O₂

For a typical diesel exhaust composition, approximately 200 ppm NO has to be reduced to 10 ppm NO in competition with an excess of 5% O₂, 5% CO₂, and 5-10% H₂O in order to meet the future automotive legislation emission standards. NO has to compete especially with O₂. O₂ is a strong oxidant and is also able to fill the oxygen defects of reduced ceria and oxidise deposited carbon. Although the results from TAP experiment showed that NO was able to be reduced on oxygen defect sites into N₂, it was still a question whether NO was able to reduce into N₂ in the presence of excess O₂. Therefore, 500 and 2000 ppm of NO was used to compete with 5% O₂ using reduced ceria and Pt/ceria.

As shown in the Figure 4A, the co-feeding 0.2% NO with 5% O₂ over reduced ceria led to the formation of CO, N₂, and CO₂ during
the first 75 s. Both NO and O₂ re-oxidised the oxygen defects. The formation of CO and CO₂ led to a consumption of the deposited carbons. NO and O₂ broke through simultaneously after 75 s and no conversion was found after 100 s. This indicated that NO was able to compete with O₂ for the oxygen defects although the NO concentration was 25 times lower than that of O₂. For 0.05% NO + 5% O₂, NO concentration was 100 times lower than that of O₂ and it was still able to compete with O₂ for oxygen defects.

The same experiments were also performed over the Pt/ceria. As shown in the Figure 6, the co-feeding 0.2% NO with 5% O₂ resulted in the formation of N₂, CO, and CO₂. Compared the intensity of CO in Figure 4 and 6, more CO was formed over Pt/ceria than that in the case of ceria. The presence of Pt accelerated carbon oxidation reaction. In other words, the oxidation of carbon was more favorable to CO over the Pt/ceria than over ceria. This observation indicated that Pt accelerated the carbon oxidation and the diffusion of O via the ceria lattice became a rate limiting step. NO and O₂ broke through simultaneously after 55 s and no conversion observed after 100 s. This further indicated that NO could compete with O₂, since no NO was observed/detected (detection limit 1 ppm) in the process of ceria catalyst re-oxidation and deposited carbon oxidation.

The results in Figure 4 and 6 indicated that small amount of NO was able to compete with excess of O₂ over both Zr-La doped ceria and Pt/Zr-La doped ceria. The presence of Pt accelerated carbon oxidation to CO, but did not alter the competition of NO with O₂.

4.3 Selectivity of NO reduction in O₂

Ceria can promote the oxidation of NO to NO₂ and, therefore, it is commonly used in the field of SCR-NH₃ and soot oxidation [24, 25]. Although NO showed full conversion during the 0.2% (and 0.05%) NO + 5% O₂ co-feeding over reduced ceria the first 75 s (Figure 4), the selectivity of NO reduction was more important than the reactivity. Undesired by-products, i.e., N₂O and NO₂, were frequently reported in NOx abatement technologies.

The result of NO reduction in O₂ (Figure 4 and 6) showed that N₂O was not detected during the whole experiment. NO₂ was only formed when NO broke through in the gas phase. From the isotopic 0.2% ¹⁵NO + 5% O₂, the formation of only ¹⁵N₂ in the first 75 s proved that N₂ was the exclusive product in the NO reduction into N₂ in O₂.

Pt is the preferred noble metal (catalyst) for the oxidation of NO into NO₂ for example in diesel soot oxidation [26]. Pt is commonly used in the NSR catalyst system and HC-SCR, which is regarded as the active site for the NO decomposition into N₂. Therefore, it will be a challenge to control the selectivity of NO conversion in the presence of O₂ to undesired by-products: N₂O and NO₂. To ascertain whether the addition of Pt to ceria could change the selectivity of NO into N₂, 0.2 % NO with 5% O₂ was fed to reduce Pt/ceria. Similarly to reduced ceria, NO in the presence of O₂ was reduced to N₂ over reduced Pt/ceria (Figure 4 and 6). No N₂O was observed in the whole experiment. NO₂ was only detected when NO broke through. From the 0.2% ¹⁵NO + 5% O₂, the formation of ¹⁵N₂ in the first 55 s proved that N₂ was the exclusive product in the NO reduction to N₂ in the presence O₂. The results of Figure 5 and 7 indicated that the addition of Pt to ceria did not change the selectivity of NO to N₂.

The results in Figure 4-7 clearly demonstrated that the oxygen defects in reduced ceria were the active sites for the NO reduction. As long as ceria was reduced, NO was selectively converted to N₂. NO₂ was only formed when (Pt) ceria was completely re-oxidised.

5. Conclusions

Ceria is a promising starting material in the development of a Di-Air NOx abatement technology. Fuel injection directly into the catalyst will result in ceria reduction and carbon deposition. Oxygen defects on reduced ceria are responsible for NO decomposition into N₂, while carbon deposits will maintain a reduced ceria surface state, and, thereby, extend the effectiveness of the hydrocarbon injections (limit fuel penalty). Platinum lowers the reduction temperature of ceria and carbon deposition. Small amount of NO can completely compete with the excess of O₂ that no NO is detected during ceria re-oxidation and deposited carbon oxidation process. Moreover, NO is selectively reduced into N₂ in the presence of O₂ over reduced ceria and Pt/ceria. Only when (Pt) ceria is fully re-oxidised, NO will be converted into NO₂.

6. References


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