Cu/Ba/bauxite: an Inexpensive and Efficient Alternative for Pt/Ba/Al$_2$O$_3$ in NO$_x$ Removal

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Cu/Ba/bauxite possesses superior NO$_x$ storage and reduction (NSR) performances, high thermal stability, strong resistance against SO$_2$ poisoning and outstanding regeneration ability in comparison with Pt/Ba/Al$_2$O$_3$. It can serve as a cheap and promising alternative for traditional Pt/Ba/Al$_2$O$_3$ in NO$_x$ removal from lean-burn engines.

With the rising concerns for energy crisis and global warming, lean-burn engines are widely used in vehicles to improve fuel economy and reduce greenhouse effect. However, the conventional three-way catalysts (TWCs) are incapable of effectively removing nitrogen oxides (NO$_x$) in lean-burn engine exhausts. One of the most feasible and attractive solution to reduce NO$_x$ emission is to use NO$_x$ storage and reduction (NSR) catalysts, which are usually composed of precious metals, alkali or alkaline earth metals, and high surface area supports. Pt/Ba/Al$_2$O$_3$ has been extensively investigated as a NSR model catalyst, but 1–2 wt% platinum loading in Pt/Ba/Al$_2$O$_3$ makes it quite expensive, the scarcity of platinum source in the earth also limits its further development and extensive application. Moreover, Pt/Ba/Al$_2$O$_3$ readily undergoes severe deactivation owing to sulfur poisoning and thermal aging, resulting in loss of catalytic activity and decline of the NO$_x$ removal efficiency. To meet the more stringent emission limit criteria, it is of great challenge to develop a practical NSR catalyst possessing high sulfur tolerance and low dependence on precious metals without sacrificing NSR capabilities.

Great efforts have been devoted to the improvement of sulfur-resistant ability and the replacement of platinum in the NSR catalysts. One effective strategy is to add the inexpensive transition metal oxides into Pt/Ba/Al$_2$O$_3$. The synergetic effect between the additives and catalysts can provide a positive structural and electrical environment for NSR. It was recently reported that the doping of CoO$_x$ or CuO$_x$ in Pt/Ba/Al$_2$O$_3$ increased NO$_x$ storage capacity and reduced the platinum loading. The addition of FeO$_x$ into Pt/Ba/Al$_2$O$_3$ was also discovered to enhance the tolerance against sulfur poisoning, but a strong deactivation is observed due to the interaction between Fe and Pt. Another strategy is the modification of supports. The blending of TiO$_2$ and Al$_2$O$_3$ was reported to help suppress sulfur deposition and enhance NO$_x$ storage capacity of the sulfur-aged catalysts. It was also observed that Ce$_{1-x}$Zr$_x$O$_2$ and Al$_2$O$_3$-Ce$_{1-x}$Zr$_x$O$_2$ nanopowders in the absence of platinum offered the comparable catalytic activity with Pt/Al$_2$O$_3$ for both NO$_x$ reduction and propane/propene oxidation. These studies have showed that introduction of the additional components into NSR catalysts could improve thermal resistance, redox property and interaction between active components and supports. Thus, it can be speculated that the combinational use of inexpensive transition metal oxides and titanium-containing composite supports would be a promising approach for the exploration of low-cost NSR catalysts.

Bauxite is an excellent support candidate in the catalytic reactions, which possesses many advantages, such as much cheap price, ready availability, rich reserve worldwide, good pore volume and air permeability, easy treatment and modification. It is composed of Al$_2$O$_3$, Fe$_2$O$_3$, TiO$_2$, CaO, SiO$_2$ and trace of platinum (Table S1), the synergetic interactions of the components will contribute to the partial or total replacement of platinum. Moreover, Fe and Ti are beneficial for the resistance against sulfur poisoning, but the investigation of bauxite in NO$_x$ removal is still unexplored. Herein, we report a sulfur-resistant and low-cost Cu/Ba/bauxite catalyst (1 wt% Cu and 16 wt% Ba), which shows superior NSR performances over Pt/Ba/Al$_2$O$_3$ (1 wt% Pt and 16 wt% Ba) without the addition of the extra platinum.
Results

The BET surface area of natural bauxite is 8 m²/g (Table S2), and it is increased to 153 m²/g after the hydrothermal treatment, which provides a possibility to serve as a support of NSR catalysts. After impregnating Cu and Ba, the specific surface of Cu/Ba/BAuxite (139 m²/g) is lower than that of Pt/Ba/Al₂O₃ (254 m²/g), but Cu/Ba/BAuxite possesses stronger sulfur resistance than Pt/Ba/Al₂O₃. A decrease of ca. 10% and 24% in surface area was found when Cu/Ba/BAuxite and Pt/Ba/Al₂O₃ were aged by SO₂, respectively. XRD patterns further confirmed superior sulfur tolerance of Cu/Ba/BAuxite over Pt/Ba/Al₂O₃. As shown in Figure S2, the XRD pattern of Pt/Ba/Al₂O₃–SO₂ exhibited the strong reflections typical for BaSO₄ at 2θ values of 26.0°, 28.7° and 42.6°, while only tiny characteristic peaks for BaSO₄ were observed in Cu/Ba/BAuxite–SO₂.

NSR performances of Cu/Ba/bauxite were initially evaluated using the NO-to-NO₂ oxidation, relative NOₓ storage capacity and NOₓ conversion. The NO-to-NO₂ oxidation is known to be an important step for NOₓ storage since the adsorption of NO₂ is more facile than that of NO in the Ba-based catalysts. Interestingly, Cu/Ba/bauxite exhibited higher NO-to-NO₂ oxidation ability than Pt/Ba/Al₂O₃ (Figure 1a). Moreover, along with the increment of temperature, the NO-to-NO₂ conversion was enhanced and reached the maximum at 300°C, subsequent decrease was observed in all of samples. The NO-to-NO₂ conversion in Cu/Ba/bauxite and Pt/Ba/Al₂O₃ at 300°C was 70% and 52%, respectively. As anticipated, the NO-to-NO₂ conversion in Cu/Ba/bauxite–SO₂ and Pt/Ba/Al₂O₃–SO₂ was decreased in comparison with their respective fresh samples, but the NO-to-NO₂ conversion in Cu/Ba/bauxite–SO₂ was still comparable with that of Pt/Ba/Al₂O₃. As shown in Figure 1b, both Cu/Ba/bauxite and Pt/Ba/Al₂O₃ possess excellent NOₓ storage performance, their relative NOₓ storage capacities are more than 95% and 90% in the range of test temperatures, respectively. The presence of SO₂ results in a slight decrement of NOₓ storage capacity in Cu/Ba/bauxite–SO₂, while a sharp decrement was observed in Pt/Ba/Al₂O₃–SO₂, which was mainly caused by the sulfation of storage component Ba, resulting in less Ba sites available to NOₓ storage. However, the presence of Fe, Ti and Cu in Cu/Ba/bauxite–SO₂ contributes to suppress the formation of BaSO₄ to a certain degree. As a result, Cu/Ba/bauxite shows much higher NOₓ storage capacity than Pt/Ba/Al₂O₃ in the presence of SO₂. The NOₓ conversion of the fresh and SO₂-aged catalysts was shown in Figure 1c. Surprisingly, although no extra platinum was added in Cu/Ba/bauxite, it still exhibits higher NOₓ conversion than Pt/Ba/Al₂O₃ at test temperatures. The NOₓ conversion of Cu/Ba/bauxite reaches 96% at 250°C with a promotion.

Figure 1 | NSR performances of fresh and SO₂-aged Cu/Ba/bauxite and Pt/Ba/Al₂O₃: (a) the NO-to-NO₂ conversion; (b) relative NOₓ storage capacity; (c) NOₓ conversion.
of 20% related to Pt/Ba/Al₂O₃. After aged by SO₂, the NOₓ conversion in Cu/Ba/ bauxite-SO₂ and Pt/Ba/Al₂O₃-SO₂ was decreased in comparison with fresh samples.

The thermal stability of Cu/Ba/ bauxite and Pt/Ba/Al₂O₃ was also examined. After Cu/Ba/ bauxite and Pt/Ba/Al₂O₃ were maintained at 350 °C for 200 h, an apparent thermal deactivation was observed in Pt/Ba/Al₂O₃ (Figure 2), the NOₓ conversion was decreased from 96 to 73%. However, only a 3% decrement was detected for Cu/Ba/ bauxite under the same conditions, which is much lower than that in Pt/Ba/Al₂O₃ and the reported NOₓ removal catalysts. Thermal deactivation was also shown by their BET surface area (Table S2), the surface area of Pt/Ba/Al₂O₃ and Cu/Ba/ bauxite was decreased ca. 38% and 11%, respectively. The average pore radius of Pt/Ba/ Al₂O₃ was increased from 2.390 to 5.886 nm, whereas no obvious change was observed in Cu/Ba/ bauxite.

To further identify NSR process of Cu/Ba/ bauxite, in situ DRIFTS experiments were carried out. As shown in Figure 3a, when Cu/Ba/ bauxite was exposed to 500 ppm NO and 8 vol% O₂ balanced by Ar at 200 °C, a strong band appears around 1249 cm⁻¹, suggesting the formation of a large number of free nitrite ions. A small band at 964 cm⁻¹ was also observed, which is assigned as the nitrite species. With lengthening of the exposure time, the band at 964 cm⁻¹ was
gradually weakened, while the characteristic peaks of nitrate around 1000 and 1365 cm⁻¹ gradually become stronger, indicating the transformation of nitrite to nitrate species. The negative peaks around 1575 and 1426 cm⁻¹ are probably due to the decomposition of carbonate. In situ DRIFTS spectra at different temperatures show the increase of temperature results in weakening of the strong band at 1245 cm⁻¹, while the nitrate peak at 1310 cm⁻¹ gets stronger and stronger, and the peak reaches the maximum at 300°C (Figure 3b), indicating that the increasing temperature favors the activation of oxygen species and subsequent oxidation of nitrites to nitrates. Thus, the NO₃⁻ storage process in Cu/Ba/bauxite should mainly proceed via nitrite route with small amount of nitrites to nitrate conversion below 300°C. When the temperature was increased above 300°C, the peak at 1245 cm⁻¹ totally disappears, implying the decomposition of nitrite species. Meanwhile, the nitrate peaks at 1310 and 1365 cm⁻¹ were greatly increased in intensity, indicating decomposition of nitrate species in part at high temperature.

When SO₂ was introduced into the system, two new peaks at 1172 and 1321 cm⁻¹ were detected (Figure 3c), which may be assigned to the chelating bidentate S-O vibration of bulk barium sulfate and the surface sulfate, respectively. It should be mentioned that the strong peak of nitrate around 1245 cm⁻¹ was not detected, while the peak of nitrate species around 1492 cm⁻¹ still exists, suggesting that NO₃⁻ storage probability is mainly via nitrite route in the presence of SO₂.

The XPS spectra of S2p for SO₂-aged and regenerated samples are shown in Figure 4. For Pt/Ba/Al₂O₃-SO₂, the strong peak around 169 eV indicates the formation of a large amount of sulfates species after exposure to SO₂, the peak intensity was decreased after the sample was reduced using 5 vol% H₂/Ar for 30 min at 400°C. However, the peak of S2p in Cu/Ba/bauxite-SO₂ is much weaker than that of Pt/Ba/Al₂O₃-SO₂, which means that the relative small amount of sulfate species was formed. Interestingly, the peak almost disappeared after Cu/Ba/bauxite-SO₂ was reduced using 5 vol% H₂/Ar for 30 min at 400°C. These results suggested Cu/Ba/bauxite-SO₂ possesses promising sulfur tolerance and regeneration ability.

Discussion

In our work, we have showed Cu/Ba/bauxite is a cheap and readily available NSR catalyst. The catalyst shows superior NSR performances over Pt/Ba/Al₂O₃ without the addition of extra platinum. It possesses outstanding resistance against sulfur poisoning and thermal stability. The good thermal stability of Cu/Ba/bauxite probably results from the natural source of the bauxite. The superior oxidation performance of Cu/Ba/bauxite was mainly ascribed to the oxidation ability of Fe, Cu and trace amounts of Pt in Cu/Ba/bauxite⁶,⁸, meantime, the presence of trace of Pt may inhibit the negative interaction of Pt and Fe in NSR. On the other hand, the Cu-based catalysts were well known for their better resistance against SO₂ deactivation than Pt-based catalysts⁹, the presence of Fe₃O₄ and Cu in Cu/Ba/bauxite-SO₂ may inhibit the formation of BaSO₄ to a certain degree. As a result, the synergetic effect of Cu, Fe, Ti, Ca and trace of Pt results in the promising performances of Cu/Ba/bauxite over Pt/Ba/Al₂O₃. The most important, the price of Cu/Ba/bauxite is much lower than that of Pt/Ba/Al₂O₃. The estimated starting material price of Cu/Ba/bauxite is less than 1% of Pt/Ba/Al₂O₃ owing to the lack of extra Pt addition. The application of Cu/Ba/bauxite will greatly cut down the catalysts cost owing to the replacement of platinum, which provides the incentive for industrial application in NOx removal of lean-burn engines.

Methods

Cu/Ba/bauxite and Pt/Ba/Al₂O₃ was prepared through conventional impregnation method. Natural bauxite was hydrothermally treated at 130°C for 96 h and followed by carbonation at 550°C for 4 h. The Ba (16 wt%) and Cu (1 wt%) were dispersed on pre-treated bauxite by the conventional impregnation method using aqueous Ba(NO₃)₂ and Cu(NO₃)₂·3H₂O solution, respectively. The resulting mixture was dried at 120°C overnight, calcined at 350°C for 2 h. The obtained sample was denoted as Cu/Ba/bauxite. The alumina support was synthesized by using the sol-gel method. The Ba (16 wt%) and Pt (1 wt%) were loaded on alumina by the conventional impregnation method using aqueous Ba(NO₃)₂ and H₂PtCl₆·6H₂O solution, respectively. The resulting mixture was dried at 120°C overnight, calcined at 350°C for 2 h. The obtained sample was denoted as Pt/Ba/Al₂O₃.

X-ray diffraction (XRD) patterns were recorded on a RIGAKU-Miniflex II X-ray diffractometer with Cu Kα radiation (λ = 1.5406 Å). N₂ physisorption measurement was performed on an ASAP 2020 apparatus, the sample was degassed in vacuo at 180°C at least 6 h before the measurement. The Brunauer-Emmett-Teller (BET) method was utilized to calculate the specific surface area in a relative pressure range of 0 ≤ P/P₀ ≤ 0.05~0.25. The pore size distribution was obtained from the adsorption branch using the Barrett Joyner Halenda (BJH) model. The SO₂-aged samples were firstly reduced by 10 vol% H₂/Ar from room temperature to 500°C and held for 2 h at 500°C, followed by the re-oxidation in 8% O₂/Ar flow at 500°C for 1 h. X-ray photoelectron spectroscopy (XPS) analysis was performed on a Physical Electronics Quantum 2000 Scanning ESCA Microprobe equipped with a monochromatic Al Kα source (Kα = 1.486.6 eV) and a charge neutralizer. The concentrations of bauxite sample were determined using a PANalytical Axios X-ray Fluorescence (XRF) spectrometer with a rhodium tube as the source of radiation. The results were analyzed by IQ Spectra and the concentrations were normalized to 100%. Inductively Coupled Plasma (ICP) was performed on a JY Ultima2 spectrometer. The NSR performances of the catalysts were estimated in a quartz flow reactor (8 mm i.d) with a fixed bed. A 0.5 g sample was pretreated under N₂ (30 mL/min) at 500°C for 1 h to remove surface-adsorbed species, and then cooled to test temperature. The feed gas was introduced using mass-flow controllers at a total flow rate of 600 mL/min. The gas compositions are listed in Table S3. The NOₓ-to-NO oxidation, relative NOₓ storage capacity (lean period) and NOx reduction (rich period) were evaluated after the catalysts were submitted to 10 lean/rich cycles and the catalytic system reached a steady cyclic state. The data were obtained based on the outlet NOx concentration under lean and rich conditions. The representative examples at 350°C were provided in Figures S3-S5. The NOx concentration was measured using an on-line chemiluminescence NO-NO₂-NOₓ analyzer (model 42i-HL, Thermo Scientific). After the measurements, the catalysts were sulfated by exposing to a feed gas containing 100 ppm SO₂ (Table S2), the catalysts were sulfated for 45 min each time. The sulfted Cu/Ba/bauxite and Pt/Ba/Al₂O₃ were denoted as Cu/Ba/bauxite-SO₂ and Pt/Ba/Al₂O₃-SO₂, respectively.

In situ DRIFTS was recorded on a Nicolet Nexus FT-IR spectrometer in the range of 650–4000 cm⁻¹ with 32 scans at a resolution of 4 cm⁻¹. In a DRIFTS cell, about 50 mg of powder sample was firstly pretreated with pure helium at test temperature for 30 min and then background spectrum was recorded. A mixture gas (500 ppm NO, 8 vol% O₂ and balanced by Ar) was introduced to the sample cell, and the spectra were immediately recorded at different exposure time up to 60 min. The DRIFTS experiments of the sulfated samples were conducted after the catalysts were exposed to NO/Ar at test temperature for 60 min, then additional 100 ppm SO₂ was introduced to the sample cell, and spectra were collected to follow the evolution of different surface species. The regeneration of Cu/Ba/bauxite and Pt/Ba/Al₂O₃ were performed with 5 vol% H₂/Ar, after the samples were exposed to 100 ppm SO₂ at 400°C for 60 min, and then DRIFTS spectra were recorded at different time. The final differential spectra were calculated by the Kubelka-Munk function.

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Author contributions
X.W. and Z.C. prepared the sample; X.W., Z.C., L.J. and R.W. designed the experiments and wrote the paper. X.W., Z.C. and Y.L. performed the measurements and analyzed the data. All authors discussed the results and commented on the manuscript.

Additional information
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