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Distinct structure-activity relationship and reaction mechanism over BaCoO$_3$/CeO$_2$ catalysts for NO direct decomposition

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ABSTRACT

The dependency on morphology is crucial for achieving highly efficient direct decomposition of NO. Herein, a BaCoO$_3$/CeO$_2$ catalyst is synthesized using CeO$_2$ small particles (p), spheres (s) and rods (r) as supports. The NO conversion to N$_2$ (NTN2) at 800 °C follows the order BaCoO$_3$/CeO$_2$-r (78.8 %) > BaCoO$_3$/CeO$_2$-s (75.9 %) > BaCoO$_3$/CeO$_2$-p (56.9 %) > BaCoO$_3$ (8.6 %) at a space velocity 1 g s/cm$^3$. BaCoO$_3$/CeO$_2$-r exhibits high tolerance to O$_2$ and stability with conversion decreasing from 78.8 % to 74.6 %, 60.0 % and 50.0 % at 800 °C with 1, 5 and 10 vol% O$_2$, respectively. The high redox activity, higher active oxygen mobility and NO adsorption capability ensures its superior performance, while the high surface area (31.29 m$^2$/g) and uniform distribution of active sites on the surface further promote the activity. The mechanism of NO direct decomposition is elucidated by in situ Diffuse reflectance infrared Fourier transform spectroscopy, $^{18}$O$_2$ isotopic transient exchange experiments and density functional theory (DFT) calculation.

1. Introduction

Nitrogen oxides (NOx), emitted in the flue gas of the industrial furnace such as new-type dry cement production process (800 °C) and gasification cooling stack in steelmaking (750–800 °C), are extremely toxic to human health and environment via forming acid rain, photochemical smog, and ozone layer depletion [1]. Currently, several technologies have been proposed for NOx abatement, including selective catalytic reduction (SCR) with ammonia, selective non-catalytic reduction and NOx storage and reduction, etc. [2–8]. Among these technologies, NO direct decomposition (2NO → N$_2$ + O$_2$) has been regarded as the most appealing pathway for NO removal due to its eco-friendly and no requirement of reductants [9]. The NO direct decomposition reaction possesses thermodynamic feasibility but belongs to a slow kinetic reaction with high activation energy (~335 kJ/mol) [10]. Therefore, the development of high-performance catalyst for NO elimination and the elucidation of the reaction mechanism are imperative.

Transition metal oxides, rare earth metal oxides, zeolites, and perovskite catalysts have been explored for NO direct decomposition [11–14]. Recently perovskite structured oxides have been investigated in many works [15]. Xu et al. reported that BaCoO$_3$, BaMnO$_3$ and BaFeO$_3$ perovskite catalysts achieve NO conversion to N$_2$ (NTN2) of 29.2 %, 28.7 % and 24.9 % at 650 °C (1 g s/cm$^3$), respectively [16]. CeO$_2$, owing to its outstanding redox activity, oxygen storage-release capability and resistance to sintering, has been widely used as one catalyst component in the three-way catalyst, etc. [17,18]. The effect of BaCoO$_3$ content on the activity and the interaction between BaCoO$_3$ and CeO$_2$ in a sample xBaCoO$_3$-CeO$_2$ synthesized with a citric acid-nitrate one-pot method was examined in our previous work. The NTN2 of the 5 %BaCoO$_3$-CeO$_2$ sample is 70.1 % while that of BaCoO$_3$ is 27.2 % at 750 °C (1.5 g s/cm$^3$) [19]. The 5 %BaCoO$_3$-CeO$_2$ sample remains NTN2 71.1 % at 800 °C in the presence of 1 vol% O$_2$. In BaCoO$_3$-CeO$_2$ sample,
BaCoO$_3$ is proved as the primary active sites to adsorb NO, while CeO$_2$ improves the reducibility, oxygen mobility and sintering resistance [20]. The morphology of CeO$_2$ was demonstrated to influence the metal active site dispersion and induce a strong metal-support interaction (MSI) with an appropriate morphology [18,21]. But the intrinsic structure-activity relationship (morphology, reaction interface, oxygen vacancy, etc.) over BaCoO$_3$ loading on different morphology of CeO$_2$ are still lacking for NO direct decomposition reaction. For the mechanism of NO direct decomposition on the perovskite-type catalyst, Shin et al. [22] proposed that two NO molecules attack active sites and oxygen vacancies simultaneously and nitrosyl species (NO$^-$) form and adsorb on iron on form N$_2$ and O$_2$ over SrFeO$_{3-x}$. catalyst. Teraoka et al. assumed that the attacking of two NO molecules in the adjacent oxide ion vacancies happens successively on a La$_{0.6}$Sr$_{0.4}$MnO$_{3-δ}$O$_2$ catalyst and the reaction follows the first-order law [23]. Chen et al. guessed that two adsorbed NO$^-$ dissociate to produce N$_2$ firstly, and NO$^-$ reacts with surface oxygen to form a NO$_2$ intermediate, which decomposes rapidly to produce NO and O$_2$ over a La$_{1-x}$Ba$_x$CoO$_3$ catalyst [24]. Recently, Xie et al. thought that NO adsorption on the metal site near oxygen vacancy to form NO$_2^0$ and O$^+$ is the rate determining step (RDS) on a BaCoO$_3$-CeO$_2$ catalyst [21]. However, so far, there is no direct evidence to elucidate its reaction mechanism for NO direct decomposition, particularly the roles of different intermediates on the perovskite-type catalysts.

Herein, a series of CeO$_2$ with varying morphologies, i.e., irregular CeO$_2$ particles (CeO$_2$-p), CeO$_2$ spheres (CeO$_2$-s) and CeO$_2$ rods (CeO$_2$-r), were prepared and employed as the supports in a BaCoO$_3$/CeO$_2$ catalyst. The distinct structure-activity relationships of the catalysts were identified. The nature of reaction mechanisms with/without O$_2$ was investigated in situ DRIFT, isotopic (H$_2$O) transient exchange experiment and DFT calculation. The results obtained comprehensively elucidate the effect of CeO$_2$ morphology on the catalytic activity.

2. Experimental

2.1. Sample preparation

The BaCoO$_3$/CeO$_2$ catalyst samples were prepared with an excess impregnation method. The irregular CeO$_2$ particles (CeO$_2$-p) was prepared with direct thermal decomposition of Ce(NO$_3$)$_3$·6H$_2$O at 600 °C. Both the nano-particle (CeO$_2$-s) and nano-rod (CeO$_2$-r) ceria samples were synthesized with a hydrothermal technique and supplied by Liaoning KeLong Fine Chemical Co., LTD, China. The prescribed amounts of Ba(NO$_3$)$_2$ and Co(NO$_3$)$_2$·6H$_2$O were dissolved in distilled water, and suitable amount of citric acid was added to the solution. The mole ratio of Ba: Co: citric acid is 1:1:1.5. The solution was stirred continuously at 80 °C and the CeO$_2$ support was immersed to achieve a loading of 5 wt% BaCoO$_3$. The reason for choosing BaCoO$_3$ with a mass ratio of 5% in all samples is that the 5% BaCoO$_3$/CeO$_2$ catalyst exhibit the best NO decomposition activity of all catalysts with the loading content of BaCoO$_3$ from 3–100 wt% in our previous work [19]. Then it was dried at 110 °C for 12 h and calcined at 400 °C for 1 h and 700 °C for 5 h. The samples are referred to as BaCoO$_3$/CeO$_2$-p (BCP), BaCoO$_3$/CeO$_2$-s (BCS), and BaCoO$_3$/CeO$_2$-r (BCR). A BaCoO$_3$ sample was also prepared with the dissolution of Ba(NO$_3$)$_2$, Co(NO$_3$)$_2$·6H$_2$O and suitable amount of citric acid in distilled water. The solution was stirred and evaporated at 80 °C to form a gel, followed by drying at 110 °C for 12 h and calcined at 400 °C for 1 h and then at 700 °C for 5 h.

2.2. Characterisation

The X-ray diffraction (XRD) patterns were recorded with a X’PERT PRO MPD Alpha1 diffractometer at 40 mA and 45 kV with a Cu K$_\alpha$ radiation source ($\lambda = 0.154056$ nm). Raman spectrum was carried out to obtain oxygen vacancy information on DXR (American Thermo Electron) with a 532 nm laser. The specific surface area (SSA) was measured using a BEL sorp Mini II instrument. N$_2$ physisorption isotherms were acquired at –196 °C after outgassing at 250 °C for 3 h under vacuum. The morphology and elemental distribution were observed with a high-resolution transmission electron microscope (HRTEM, JEM 2100F, Oxford) equipped with energy dispersive X-ray detector (EDX) mapping (Tecnai G$^2$ F20, Oxford) and a scanning electron microscope (SEM, Hitachi S-4800). X-ray photoelectron spectroscopy measurement (XPS) was carried out with a Kratos Axis Ultra DLD spectrometer. The binding energy was calibrated with the Cl$_1$s peak at 284.8 eV.

H$_2$ temperature-programmed reduction (H$_2$-TPR) was performed on an Ichem700 apparatus. The sample (50 mg) was pretreated at 500 °C for 1 h under Ar flow (30 mL/min) to remove adsorbed impurities, and then cooled down to room temperature (RT). The sample was then heated from RT to 900 °C at 10 °C/min with a flow of 5 vol%H$_2$/Ar (30 mL/min). Temperature-programmed desorption of O$_2$ (O$_2$-TPD, Ichem700), and NO (NO-TPD-Mass spectrometer, MS, Pfeiffer Omnistar$^{TM}$) were done to examine the chemisorption of O$_2$ and NO, respectively. For O$_2$-TPD, the sample was pretreated at 500 °C for 1 h under Ar flow (30 mL/min) and exposed to O$_2$ (30 mL/min) for 1 h at RT, followed by operating from RT to 900 °C at 10 °C/min with pure Ar (30 mL/min) after purging with Ar (30 mL/min) for 0.5 h. For NO-TPD-MS, after being pretreated at 700 °C under Ar (50 mL/min) for 1 h, the sample (200 mg) was cooled to 80 °C and then adsorbed to saturation with 2 vol% NO/Ar for 1 h. The desorption gases were recorded with the MS from 80 °C to 700 °C at 10 °C/min.

In-situ Diffuse reflectance infrared Fourier transform (DRIFT) spectra were obtained with a Thermo Fisher Scientific Nicolet iS50 spectrometer. The catalyst was pretreated in helium atmosphere with a flow rate of 50 mL/min at 600 °C for 1 h and then cooled to RT. The background spectra were collected at desired temperatures during the cooling process. Afterwards, the reaction gases of 1 vol% NO/He, and 1 vol% NO+2 vol% O$_2$/He were switched into the in-situ reaction cell with 30 mL/min for 0.5 h until stability, respectively. The steady state spectra were recorded with accumulating 32 scans at 30, 200, 300, 400, 500, 550 and 600 °C, respectively. The isotopic (H$_2$O) exchange experiments were carried out in an in-situ fixed-bed reactor (i.d. = 3 mm), and the products were monitored with an online mass spectrometer (MS, Pfeiffer Omnistar$^{TM}$). A four-way switch valve was firstly used to switch the gas path instantaneously and retain steady continuous flow of reactant gases, as shown in Fig. S1 in the supporting information. The sample (65 mg, 0.09–0.15 mm) was pretreated at 800 °C for about 20 min at the flow rate of 30 mL/min in Ar. Afterwards, the pre-mixed reactant gas (30 mL/min, 1 vol% NO/Ar) was constantly introduced into the reactor at 800 °C for 20 min and the products were monitored with the online MS. Pure Ar was used to purge at 30 mL/min for 10 min to remove the gases. Finally, the mixed reactant gas (30 mL/min, 1 vol% NO + 1 vol% $^{18}$O$_2$/Ar) was pulsed into the reactor at 800 °C for about 25 min to detect the isotopic products adequately. The isotopic gas ($^{18}$O$_2$) was supplied by WUHAN NEWRADAR SPECIAL GAS Co., LTD.

The density functional theory (DFT) simulation was further performed to clarify the perovskite-support interaction and reaction pathways, and the detailed calculation method was introduced in the supporting information.

2.3. Catalytic activity

The catalyst samples (0.5 g, 35–45 mesh) were placed in a fix-bed quartz tube reactor (i.d. = 6 mm). The sample was pretreated to remove the gaseous impurities on the surface at 500 °C for 0.5 h in Ar (30 mL/min) and cool down to RT. 8000 ppm NO/Ar was introduced into the reactor at atmospheric pressure with a flow rate 30 mL/min (1 g s/cm$^2$, GHSV=5600 h$^{-1}$). Each temperature step was maintained for 20 min to reach steady state from 500 to 800 °C with an interval of 50 °C. The N$_2$ concentration was quantified online with the same MS calibrated with an internal standard method. The concentrations of NO, O$_2$, CO$_2$ and SO$_2$ were measured by online multicomponent analyzers (Gasboard-
300UV for NO/O₂, Gasboard-300plus for SO₂, Hubei Cubic-Ruiyi Instrument Co., Ltd. And QGS-08 for CO₂, Maihak). The oxygen tolerance of the catalyst was measured with feeding 1, 5, and 10 vol% O₂ in the 8000 ppm NO/Ar flow with a total flow rate 30 mL/min. The 5 vol% H₂O was introduced into the gas mixture by passing the gas stream through a gas saturator at 35 °C to study the effect of H₂O.

The activity is evaluated with NTN2 according to equation (1),

\[
\text{NO Conversion to N}_2 = \frac{2[N_2]_{\text{out}}}{[NO]_{\text{in}}} \times 100\%
\]

where \([NO]_{\text{in}}\) and \([N_2]_{\text{out}}\) are the inlet and outlet concentrations of NO and N₂, respectively.

The reaction rate \(r\) of NTN2 at 800 °C is calculated with equation (2),

\[
r = \frac{F \cdot c \cdot X}{W_{\text{cat}}}
\]

where \(F\) is the total feed flow (mol/s), \(c\) is NO inlet concentration (%), \(X\) is NTN2 (%), \(W_{\text{cat}}\) is the weight of catalyst (g).

3. Results

3.1. Catalytic activity

The NO direct decomposition activity of the catalyst samples at 500–850 °C is depicted in Fig. 1a. Pure CeO₂-R exhibits almost negligible activity, achieving only 7.5 % at 850 °C. Similarly, pure BaCoO₃ displays poor activity, with the highest NTN2 at 750 °C being only 12.46 %. However, the activity sharply increased after loading BaCoO₃ on CeO₂ with different morphologies, particularly on the spherical and rod-like CeO₂ samples. The NTN2 at 800 °C follows a distinct order of BCR (78.8 %) > BCS (75.9 %) > BCP (56.9 %) > BaCoO₃ (8.6 %). The activity of BCS and BCR is about 19–22 % higher than that of the BCP sample, indicating the beneficial interaction between BaCoO₃ and CeO₂ with regular morphologies. The activity of BCR outperforms similar catalysts reported in the literature, as indicated in Table S1. The O₂ yield over BCR is also obtained, and it attains a value of 72.9 % at 800 °C. The amount of O₂ formation is always slightly less than that of N₂ at the same temperature. But if considering the formation of a small amount of NO₂, the quantities of N₂ and O₂ are nearly balanced for NO direct decomposition, which is also verified in the reference [25,26]. Hence, it seems likely that no accumulation of oxygen is anticipated on the catalyst surface. Subsequently, the NTN2 value is mainly discussed in the NO direct decomposition reaction. The intrinsic rates of the three catalysts at 800 °C are calculated and plotted in Fig. 1b. BCR exhibits the highest rate (2.81 × 10⁻⁷ mol g⁻¹ cat⁻¹ s⁻¹) among the three samples.

The O₂ resistance experiment results of BCR are presented in Fig. 1(c). The NTN2 declines from 78.77 % to 74.56 %, 60 % and 50 % at 800 °C under 1, 5 and 10 vol% O₂, respectively. Furthermore, the BCR sample shows a stable performance in the presence of 5 vol% O₂ over 12 h with only about 5 % decrease in conversion, see Fig. 1(d). The period drops and variation of the activity for the 8000 ppm NO/Ar + 5 vol% O₂ of Fig. 1d may be attributed to the slight change of vacuum degree in the mass spectrometer. Then under the coexistence of 5 vol% O₂ and 5 vol% H₂O, the NTN2 is stably sustained about 43 % over 12 h after slight decreases from 52 % to 44 % within the initial 5 h. Further, the BCR catalyst experiences an evident activity reduction from 53.1 % to 37.4 % after feeding 1 vol% NO/Ar + 5 vol% H₂O + 100 ppm SO₂ for 10.5 h. The NTN2 of BCR catalyst decreases from 50.9 % to 40 % for 11 h under the 1 vol% NO/Ar + 5 vol% O₂ atmosphere. The negative effect of CO₂ is due to the CO₂ can strongly adsorb on the Ba sites of BCR to form carbonate, in agreement with the results in the reference [27]. Therefore, the existence of O₂, CO₂, H₂O and SO₂ in actual flue gas all impede the catalytic performance of BCR catalyst for NO direct decomposition.
3.2. Structure and texture

The XRD patterns of the catalyst samples are presented in Fig. 2. For the BaCoO3/CeO2 samples, only the peaks of CeO2 phase appear and no diffraction peaks of BaCoO3 phase are observable no matter of the CeO2 morphology, indicating well dispersion of BaCoO3 on CeO2. Compared to the patterns of the pure CeO2 samples, i.e., the p, s and r, see Fig. S3, no shift of CeO2 peaks are observed. This result suggests that inter-diffusion may not occur between the BaCoO3 and CeO2 components. The BaCoO3 sample prepared under the same condition shows an XRD pattern typically for the perovskite phase (JCPDS 70-0363). The Raman spectrum is further performed in Fig. 2(c). For the catalysts containing cerium, the peak at 458–468 cm$^{-1}$ belongs to the F2g symmetric vibration of oxygen atoms with the fluorite-type structure cerium in CeO2 [28,29]. Compared with pure CeO2-r (468 cm$^{-1}$), the F2g peaks of BCP, BCS and BCR are slightly shifted to lower wave number (458 cm$^{-1}$) and peaks intensity becomes weaker, attributing to the addition of BaCoO3 causing distortion of CeO2 lattice structure to form more oxygen vacancies, particularly for BCR. There are three additional bands at 232, 598 and 1172 cm$^{-1}$, owning to modes of second-order transverse acoustic (2TA), defect-induced (D), and longitudinal optical (2LO), respectively [27]. The peak at 616 cm$^{-1}$ is assigned to the typical vibrational peak for perovskite-type structure of BaCoO3. The oxygen vacancy ($O_v$) concentration ($I_{598}/I_{458}$) was also obtained by semiquantitative analysis in Fig. 2(d). It is clearly seen that the $O_v$ concentration decreases with the sequence of BCR (0.13) > BCS (0.10) > BCR (0.06) > CeO2-r (0.02). The highest $O_v$ concentration in BCR is resulted from the stronger interface interaction between BaCoO3 and CeO2-r, which generates more lattice defects and oxygen vacancy.

The N2 adsorption-desorption isotherms and pore size distribution curves of the samples are plotted in Fig. 3. All the samples exhibit type IV isotherms with a H3-type hysteresis loop, illustrating that mesopores with good pore connectivity are dominant in Fig. 3a. The SSA of the samples follows an order of BCR (31.3 m$^2$/g) > BCS (28.6 m$^2$/g) > BCP (15.6 m$^2$/g) > BaCoO3 (1.2 m$^2$/g), which is in consistency with the activity order. It indicates that BCR displays the highest SSA, attributed to the rod morphology of CeO2 impedes effectively the catalyst sintering and is beneficial to promote the dispersion of active sites [28,29]. The average pore size of BCR is the smallest (15.98 nm) and the pore size distributes in the range of 2–20 nm in Fig. 3b.

Fig. 4 and S4 give the micrographs of the three catalyst samples. The particles in BCP are small agglomerates, while BCS and BCR display well-defined CeO2 spheres and rods with large particle sizes, Fig. 4a1-c1. The average particle sizes for the BCP, BCS and BCR samples are ~27 nm, ~184 nm and ~5 um, respectively. The HRTEM images, Fig. 4a2-c2, show that the (110), (102), (101), (211) planes of BaCoO3 in the samples are observed, corresponding to the lattice fringes of 0.28, 0.21, 0.34 and 0.17 nm, respectively. The CeO2 supports show d-spacing of 0.32 nm with (111) facets for BCP, 0.19 nm with (220) facets for BCS, and 0.19 and 0.32 nm with (220) and (111) facets for BCR. Herein, the major exposed facets are (111) for BCP, (220) for BCS and BCR, in good agreement with the literature [28,29]. Compared with (111) plane of CeO2 for BCP, the (220) plane of CeO2 is unstable and has lower formation energy of oxygen vacancy for BCS and BCR. Ba, Co, and Ce elements distribute evenly on all the catalyst samples surface, evidenced by the STEM-EDX mapping graphs in Fig. 4a3, b3 and c3.

3.3. Surface states and chemisorption

Fig. 5 plots the XPS Co 2p, Ce 3d and O1s spectra of the three catalyst samples in comparison with those of BaCoO3 and CeO2. The results of elemental analysis calculated with peak area integration are listed in Table 1. The Co 2p spectrum of pristine BaCoO3 is deconvoluted to distinguish the Co$^{3+}$ and Co$^{2+}$ contributions. The peaks at 779.6 and 794.8 eV are attributed to Co$^{3+}$ while 781.2 and 796 eV are assigned to Co$^{2+}$ in Fig. 5a [30]. The peak position of Co$^{3+}$ shifts to higher binding energy.
energy (~780.5 eV) in the three catalyst samples. Meanwhile, the part of Co\(^{3+}\) transfers to Co\(^{2+}\) at 796.1 eV owning to the electron transfers of Co species on the surface of CeO\(_2\) support during high temperature calcination step [31]. The semi-quantified Co\(^{3+}/Co^{2+}\) ratio of BCP, BCS, and BCR catalysts are 2.66, 3.32, and 3.58, respectively. For the Ce 3d spectra in Fig. 5b, the spectra of the three catalyst samples can be fitted to eight peaks (u, u’ , u”, ν, v, ν’ and ν”), which are assigned to u and v for Ce\(^{3+}\) that favors oxygen vacancies formation, as well as u’, u” , ν, ν’ and ν” for Ce\(^{4+}\) [32]. The amount of Ce\(^{3+}\) is related to the amount of oxygen vacancies (O\(_v\)) in CeO\(_2\). The Ce\(^{3+}/Ce^{4+}\) ratio is calculated based on the peak areas, which decreases in the sequence of BCR (0.19) > BCS (0.18) > BCP (0.16), suggesting that relatively higher amount of Ce\(^{3+}\) and O\(_v\) exist in BCR sample. Among the three catalyst samples, the higher ratios of Co\(^{3+}/Co^{2+}\) and Ce\(^{3+}/Ce^{4+}\) in BCR induce electron transfer at the interface between BaCoO\(_3\) and CeO\(_2\). As shown in Fig. 5, the O 1s spectrum has three peaks at ~529.2, ~531.2 and ~533.7 eV, corresponding to lattice oxygen (O\(_{\text{lat}}\)), adsorbed oxygen (O\(_{\text{ads}}\)) and hydroxyl oxygen (O\(_{\text{OH}}\)) species, respectively [33]. The O\(_{\text{ads}}\)/O\(_{\text{lat}}\)/(O\(_{\text{ads}}\)+O\(_{\text{lat}}\)+O\(_{\text{OH}}\)) ratio has an order of BCR (0.32) > BCS (0.30) > BCP (0.29), in agreement with the ranking of Ce\(^{3+}/Ce^{4+}\) ratio in the catalysts and Raman results (Fig. 2d). The ratio of adsorbed oxygen is correlated to the oxygen vacancy concentration [34]. Thus, the BCR shows a superior activation, transformation and desorption of oxygen via the function of Ce\(^{3+}/O_v\). Furthermore, the characteristic peaks of O\(_{\text{OH}}\) for BCP, BCS and BCR are located at lower binding energy at 530.6–530.8 eV than that for pure CeO\(_2\)-r (531.2 eV) and BaCoO\(_3\) (531.15). This observation indicates the higher mobility of lattice oxygen and formation of more oxide ion vacancies by introducing CeO\(_2\) supports in BCP, BCS and BCR catalysts, particularly in BCR, which promotes the catalytic activity.

The pure BaCoO\(_3\) shows three peaks in its H\(_2\)-TPR profile Fig. 6(a), labeled as α\(_1\) (362 °C), α\(_2\) (499 °C) and α\(_3\) (610 °C), corresponding to the reduction steps of Co\(^{4+}\) to Co\(^{3+}\), Co\(^{3+}\) to Co\(^{2+}\) and maybe Co\(^{2+}\) to Co\(^{0}\), respectively [35,36]. For pure CeO\(_2\)-r, two peaks at β\(_1\) (595 °C) and β\(_2\) (821 °C) are assigned to the reduction of surface and bulk Ce\(^{4+}\) to Ce\(^{3+}\), respectively [37]. It is explicit that the α peak (including α\(_1\) at 294 °C, α\(_2\) at 340 °C and α\(_3\) at 360 °C) for BCP, BCS and BCR catalysts shift to lower temperatures and still mainly exist as Co\(^{3+}\) compared to those of the pure BaCoO\(_3\). Meanwhile the β\(_1\) reduction peak almost disappears in all three catalysts. Obviously, the BaCoO\(_3\) component is highly dispersed on the CeO\(_2\) surface and strong synergy between two phases, benefiting to promote the reduction activity. In addition, the actual H\(_2\) consumption of α peak (Fig. 6d) follows order BCR (254.7 umol/g) > BCS (217.3 umol/g) > BCP (186.5 umol/g). The H\(_2\) consumption of Cerium species (β peak in Fig. 6d) is similar in these three catalysts. These mean that the more Co\(^{3+}\) species reduction and stronger interaction occurs in BCR, contributing to enhanced catalytic activity.

The mobility of lattice oxygen is examined by O\(_2\)-TPD in Fig. 6(b). Two peaks contain adsorbed oxygen on the oxide ion vacancies (~730 °C) and lattice oxygen (~860 °C) in the catalysts [38]. Compared with the desorption peak temperature of lattice oxygen in pure BaCoO\(_3\), the peaks of BCP, BCS and BCR shift to lower temperatures orderly, suggesting that different morphology of CeO\(_2\) effectively weakens the Co-O bond strength and promotes the desorption of oxygen from BaCoO\(_3\). Particularly, the BCR owns the highest lattice oxygen mobility, which is conducive to enhance activity and resistance of O\(_2\).

Fig. 6(c) displays the NO-TPD profiles to analysis NO sorption of catalysts. There are two NO desorption peaks at 314 and 507 °C for pure BaCoO\(_3\), meanwhile NO desorbs at 256 and 525 °C for pure CeO\(_2\)-r [39,40], which are attributed to weak and strong NO adsorption species on different components, respectively. The BCP, BCS and BCR show similar NO desorption profiles but all bigger than that of pure samples due to the interaction effect may produce more electron coordination environment on the active sites to adsorb NO. Combined to the calculated NO desorption amount of catalysts (Fig. 6d), decreasing in the order of BCR (318.3 umol/g) > BCS (260.3 umol/g) > BCP (228.1 umol/g) > BaCoO\(_3\) (122.3 umol/g) > CeO\(_2\)-r (75.8 umol/g). BCR exhibits the highest NO adsorption capacity and hence increase activity. Accordingly, the reducibility, mobility of lattice oxygen and NO sorption of catalysts decrease in the sequence of BCR > BCS > BCP > BaCoO\(_3\), in agreement with the variation of activity results (Fig. 1a).
3.4. In situ DRIFT and isotopic \(^{18}\text{O}_2\) tracing experiments

To further investigate the adsorbed intermediates and reaction behavior of the samples, in-situ DRIFT is applied with BCR sample (Fig. 7), and corresponding detailed assignment of NO adsorption bands are listed in Table 2. As shown in Fig. 7(a), the in-situ DRIFT profiles are recorded under 1 vol% NO/He from 30 to 600 \(^\circ\text{C}\). The intensity bands for nitrous oxide (N\(_2\)O) at 2240 and 2210 cm\(^{-1}\) are weak at 30 \(^\circ\text{C}\) and then gradually disappear with the temperature increase from 200 to 600 \(^\circ\text{C}\) (Fig. 7b) [41]. In general, the N\(_2\)O acts as main intermediate and decomposes quickly at higher temperature and few parts of N\(_2\)O still adsorb on the catalyst surface [41]. The adsorbed mononitrosyl (NO\(^-\)) at 1930 and 1845 cm\(^{-1}\) are assigned to Co\(^{3+}\) sites, revealing that the cobalt species acts as the main active center for NO decomposition reaction [20,42,43]. The intensity of mononitrosyl peak becomes weaker with the temperature increase, whilst the monodentate nitrite (NO\(^{-2}\) adsorbed on Co\(^{3+}\) (1520 cm\(^{-1}\)) becomes stronger, suggesting that the subsequent transformation from NO\(^-\) to NO\(^{-2}\) on the Co species of catalyst [44]. The band at 1774 cm\(^{-1}\) for asymmetric nitric oxide dimer ((NO)\(_2\)/-NOON) is formed on Co\(^{3+}\) during the NO adsorption process [20,21,45], in agreement with the IM2 structure in the following DFT results of Fig. 9(c) and (d). The intense bands of NO on Co\(^{3+}\) and weak -(NO)\(_2\) species formed on Co\(^{3+}\)/Co\(^{2+}\) cycles supply effective electrons for the NO adsorption and transformation, in consistency with the ratio of Co\(^{3+}\)/Co\(^{2+}\) in XPS results (Fig. 5a). In addition, the chelating bidentate nitrite (NO\(^{-2}\)) adsorbed on CeO\(_2\) (1238 and 813 cm\(^{-1}\)) turns weaker, probably due to NO\(^{-2}\) desorbs in the form of NO\(_2\) molecule at the certain temperature [46]. The strong bands at 1608–1013 cm\(^{-1}\) are attributed to the nitrate species with different adsorption states [26,47]. Those adsorption species become more stable and stronger with the temperature increase to 200 \(^\circ\text{C}\), particularly nitrate free ion to 400 \(^\circ\text{C}\), and then decompose at the higher temperature with the decreased band intensity until almost disappear at 600 \(^\circ\text{C}\). The contribution of Ba is more likely to promote the reduction of Co ions and formation of oxygen vacancy for stable ABO\(_3\) perovskite type catalyst [24].

The effect of O\(_2\) on the adsorbed intermediate species over the BCR was also explored under 1 vol% NO + 2 vol% O\(_2\)/He in Fig. 7(c) and (d). The intensity of bridging bidentate nitrate (1608 cm\(^{-1}\)) and monodentate nitrate (1408 cm\(^{-1}\)) become stronger than that of 1 vol% NO/...
state of each reaction stage. The effect of O2 on the NO direct decom-
position was illustrated in Fig. 8(b). Before the reaction, the fresh BCR
was pretreated and kept in Ar (30 mL/min) at 800 °C over the BCR catalyst after switching 1 vol% NO/Ar, while no by-product forms (Fig. S5). Then the mixed gas of 1 vol% NO + 1 vol% 18O2/Ar was fed to the reactor after purging with pure Ar. It is noted that the three different O2 isotopomer products (16O2, 16O18O and 18O2) were detected in turn. Firstly, large amounts of 16O2 produc-
ed quickly, which decreased rapidly and then increased gradually to steady value with increasing reaction time, attributed to NO decom-
position product and releasing from the BCR itself. The heteroexchange of 16O2 with original 16O in catalyst was noticed at high temperature with high oxygen mobility [49]. After 16O2 desorption from catalyst, the oxygen vacancy sites generated accordingly are simultaneously replenished by the surface adsorption and dissociation of gaseous 18O2 molecules followed by the lattice oxygen diffusion, resulting in the 16O2 signal produced quickly and then decreased [50]. The N18O should be considered to produce because of the same m/s (32) to 16O2. The signal of N18O-16O2 increased gradually to steady with increasing time reflects that the oxygen of NO can be exchanged with the part of adsorbed 18O from the catalyst surface to form N18O, which is also supported by Damma et al. [51]. Subsequently, the cross-labeled 16O18O generated quickly, corresponding to the rapid decrease of 16O2, suggesting that the suitable lattice 18O species adsorbed on the surface oxygen vacancies start to participate NO decomposition and promote 16O18O and N2 forma-
tion, in associate with the stable N2 signal at that time. Afterwards, the 18O2 increased gradually and tend to reach steady state with the increase of reaction time, attributed to excess gaseous 18O2 fills more oxygen vacancies firstly but can not exchange completely, resulting in a competitive adsorption with NO on the oxygen vacancies and the cata-
lytic activity decreases slightly, corresponding to a gradual decrease of N2 formation after about 60 min. It suggests that the updated and appropriate lattice oxygen (18O) species accelerate the reaction at the beginning of O2 addition. However, excess 18O2 produces an inhibition effect of NO adsorption and transformation. In addition, very small amounts of intermediate species (N2 18O and 16O18O) were monitored, indicating that no other impurity gas existed in the mixed reactant gases. The pure Ar (30 mL/min) was purged to remove reactants existed in the reactor after realizing steady state of each reaction stage. The effect of O2 on the NO direct decom-
position was illustrated in Fig. 8(b). Before the reaction, the fresh BCR was pretreated and kept in Ar (30 mL/min) at 800 °C. The evident and stable signals of N2 and 16O2 can be observed owning to the NO efficient decomposition at 800 °C over the BCR catalyst after switching 1 vol% NO/Ar, while no by-product forms (Fig. S5). Then the mixed gas of 1 vol% NO + 1 vol% 18O2/Ar was fed to the reactor after purging with pure Ar. It is noted that the three different O2 isotopomer products (16O2, 16O18O and 18O2) were detected in turn. Firstly, large amounts of 16O2 produc-
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tion, in associate with the stable N2 signal at that time. Afterwards, the 18O2 increased gradually and tend to reach steady state with the increase of reaction time, attributed to excess gaseous 18O2 fills more oxygen vacancies firstly but can not exchange completely, resulting in a competitive adsorption with NO on the oxygen vacancies and the cata-
lytic activity decreases slightly, corresponding to a gradual decrease of N2 formation after about 60 min. It suggests that the updated and appropriate lattice oxygen (18O) species accelerate the reaction at the beginning of O2 addition. However, excess 18O2 produces an inhibition effect of NO adsorption and transformation. In addition, very small amounts of intermediate species (N2 18O, N16O18O and N18O2) can be obtained, which may not decompose (Fig. S4). On the basis of isotopic (18O2) tracing studies, it can be found that moderate labile oxygen species on the surface oxygen vacancies of catalyst play a key role in NO direct decomposition reaction. 3.5. DFT calculation results

To deeply understand the synergy effect between BaCoO3 and CeO2 support of catalyst and direct decomposition mechanism of NO, the DFT-U method is performed to study this reaction. The reaction pathway is connected by initial state (IS), intermediate (IM), transition state (TS) and final state (FS). Based on catalyst morphology of TEM and XRD, pure BaCoO3(101), BaCoO3/CeO2 (110) for BCS and BCR, and BaO2/CeO2 (111) for BCP models are constructed in Fig. 9(a). The reaction mechanisms of NO on BaCoO3/CeO2 (110) and BaCoO3/CeO2

Table 1
Surface atomic ratios of catalysts obtained by XPS semi-quantitative analysis.

<table>
<thead>
<tr>
<th>Catalysts</th>
<th>Surface area ratio</th>
<th>Co3+/Co2+</th>
<th>Co4+/Co2+</th>
<th>Oads/(Oads+Oint+Ovol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CeO2-r</td>
<td>-</td>
<td>0.23</td>
<td>0.21</td>
<td></td>
</tr>
<tr>
<td>BaCoO3</td>
<td>2.69</td>
<td>0.16</td>
<td>0.29</td>
<td></td>
</tr>
<tr>
<td>BCP</td>
<td>3.32</td>
<td>0.18</td>
<td>0.30</td>
<td></td>
</tr>
<tr>
<td>BCS</td>
<td>3.58</td>
<td>0.19</td>
<td>0.32</td>
<td></td>
</tr>
<tr>
<td>BCR</td>
<td>3.58</td>
<td>0.19</td>
<td>0.32</td>
<td></td>
</tr>
</tbody>
</table>
(111) surfaces have been explored, as in Fig. 9. The adsorption of NO on the two different surfaces has been investigated. NO can be adsorbed on the surfaces through either the N-end or the O-end. Adsorption structures through the N-end are more stable on both BaCoO$_3$/CeO$_2$ (110) and BaCoO$_3$/CeO$_2$ (111) surfaces, with adsorption energies of 0.82 eV and 0.57 eV, respectively. Namely, NO adsorption on BaCoO$_3$/CeO$_2$ (110) releases ~0.25 eV more energy compared to on BaCoO$_3$/CeO$_2$ (111). On the other hand, adsorption through the O-end on the surfaces has lower energies, with values of 0.38 eV and 0.20 eV, respectively. The direct decomposition mechanism of NO on the BaCoO$_3$ (101) surface is also proposed in Fig. S5. Similarly, NO is also adsorbed through the N-end on the surface with the bigger adsorption energy (0.46 eV) than the O-end on the surface (0.33 eV), suggesting that the structure with NO adsorbed through the N on the surface is more advantageous on BaCoO$_3$. As the second NO molecule is adsorbed on the surface, the two NO molecules can directly react to form a bridging *NOON species (IM2). This reaction process releases energies of 1.72 eV for BaCoO$_3$/CeO$_2$ (110) and 1.99 eV for BaCoO$_3$/CeO$_2$ (111). Unlike on BaCoO$_3$ (Fig. S6), this transition state can occur spontaneously without the energy barrier on BaCoO$_3$/CeO$_2$. As the reaction progresses, the *NOON species undergoes a turnover to adsorb on the surface through the O-end (IM3). Different from that on BaCoO$_3$ with the negligible energy (0.08 eV), this process requires energies of 0.70 eV (BaCoO$_3$/CeO$_2$ (110)) and 1.01 eV on (BaCoO$_3$/CeO$_2$ (111)). This can be attributed to the stronger interaction between the N-end and the Co adsorption sites. Alternatively, NO can also directly form the IM3 structure through the O-end. After the second NO molecule is adsorbed through the O-end on the surface, it can directly react with the already adsorbed NO to form a *NOON species connected on the surface through the O-end. This process is also a barrierless reaction. As one of the N-O bonds in *NOON is broken (TS1), the dissociated *O atom fills an oxygen vacancy on the surface, while *N$_2$O molecule adsorbs on the surface through the O-end (IM4), in agreement with in situ DRIFT results. The energy barrier for this process is ~0.6 eV. Finally, after overcoming an energy barrier of about 1 eV, the N-O bond in *N$_2$O is broken (TS2). The *O atom fills the vacancy, while N$_2$ desorbs from the surface. The reaction energies for this process are 1.11 eV for BaCoO$_3$/CeO$_2$ (110) and 1.26 eV for BaCoO$_3$/CeO$_2$ (111), respectively. In addition, the energy barrier of O$_2$ adsorbed on the Co site is about 0.36 eV for BaCoO$_3$/CeO$_2$ (110), which is lower than that of NO adsorbed on the Co site, suggesting the NO is easier to adsorb on active sites.

4. Discussion

4.1. Intrinsic structure-activity relationship

The intrinsic catalytic activity of the BaCoO$_3$/CeO$_2$ catalyst shows a significant morphology-dependent behavior, which is displayed in Scheme 1. The addition of CeO$_2$ with different morphology induces a sensitive perovskite-support interaction (PSI). BCR owns the largest average particle size (~5 μm) and SSA with regular rod appearance, which provides enough reactive interface area to adsorb and transform reactants, which not only alleviates the sintering of BaCoO$_3$ effectively but also strengthens the synergy between BaCoO$_3$ and CeO$_2$ (Figs. 3 and 4). Hence, the well dispersed BaCoO$_3$ on the surface of CeO$_2$ support, and both components contact with each other closely to form effective reaction interface is beneficial for strong interaction to promote activity in NO direct decomposition. Compared with (111) plane of CeO$_2$ for BCP, the (220) plane is unstable and has lower formation energy of oxygen vacancy for BCS and BCR. Meanwhile, the higher ratios of Ce$^{3+}$/Ce$^{4+}$, Co$^{3+}$/Co$^{4+}$, O$_{ads}$/(O$_{ads}$+O$_{lat}$+O$_{st}$) and O$_{v}$ concentration of the BCR sample were induced by PSI, exhibiting more active electron
transfer and oxygen vacancy at the BCR interface (Fig. 5). The BCR surface contains Co$^{3+}$ and Co$^{2+}$ ions with an electron configuration of $t_{2g}^2e_g^2$ and $t_{2g}^5e_g^2$, respectively [52]. The Co$^{3+}$ possesses more unpaired electron than Co$^{2+}$, resulting in a promoting affinity for nitrogen donors when the ratio of Co$^{3+}$/Co$^{2+}$ increase [52]. By comparing the PSI on BaCoO$_3$ and BaCoO$_3$/CeO$_2$, we can observe that BaCoO$_3$/CeO$_2$ exhibits a stronger adsorption ability to NO. Furthermore, the NO adsorption sites on BaCoO$_3$/CeO$_2$ allow for the direct formation of *NOON species without the energy barrier. This characteristic enhances the catalytic performance of BaCoO$_3$/CeO$_2$, attributed to the synergistic effects between BaCoO$_3$ and CeO$_2$. The CeO$_2$ promotes the stabilization of the adsorbed species, leading to stronger NO adsorption. The proximity of the adsorption sites facilitates the direct formation of *NOON species, which further enhances the catalytic activity of BaCoO$_3$/CeO$_2$. Another factor contribute to the stronger catalytic performance of BaCoO$_3$/CeO$_2$ (110) is the influence of oxygen vacancy (O-vacancy). The electrons left behind with more oxygen vacancy formation which reduces Ce$^{4+}$ to Ce$^{3+}$ ions [53]. It is well known in DFT results that the O vacancy tends to be on the CeO$_2$ (110) surface (1.69 eV). On CeO$_2$ (111) surfaces, it is usually located at subsurface sites (1.83 eV). It is also consistent with the experimentally observed higher concentration of oxygen vacancies in BCR samples. Consequently, oxygen vacancies located on the surface are more prone to migrate to the BaCoO$_3$, enhancing the coordination effects and promoting catalyst activity. Therefore, compared to BaCoO$_3$/CeO$_2$ (111) for BCP, the combination of stronger NO adsorption and the presence of surface oxygen vacancies in BaCoO$_3$/CeO$_2$ (110) for BCS and BCR contribute to its superior catalytic performance in agreement with the activity results. That highlight the importance of surface orientation and oxygen vacancy distribution in determining the catalytic properties of composite materials.

Electronic interactions always occur at solid-solid interfaces and are simply driven by the principles of the system energy minimization and the continuity of the electric potential in the solid phases [54–56]. The redox interaction effect (Co$^{3+}$ + Ce$^{3+}$ ↔ Co$^{2+}$ + Ce$^{4+}$) produces long-range charge transfer and thereby causes more oxygen vacancies under the stable electric field on the concomitant BCR interface, suggesting the high availability of reaction environment on BCR with rod-like CeO$_2$. The BCR active interface is likely formed via a bonding between Co species in BaCoO$_3$ and the surface defect sites on both perovskite and ceria (Ce$^{3+}$–O$_2$) species. The similar results of interface effect between metal site and the morphology of CeO$_2$ supports for other catalytic reactions were also verified in the literatures [18,57–60]. To sum up, the construction of active interface on BCR improves PSI and activates electronic interactions, promoting reducibility, lattice oxygen mobility with more Oxygen vacancy and NO adsorption and boosting the catalytic performance in NO direct decomposition reaction.

4.2. Proposed reaction pathways

The active sites of Co species and oxygen vacancies on the interfaces in BCR were observed. Combined with in situ DRIFT, isotopic ($^{18}$O$_2$) exchange and DFT results, the NO molecules are effectively adsorbed on the Co species, but a competitive adsorption between NO and O$_2$ occurs on the oxygen vacancies after introducing O$_2$, resulting in a decrease of catalytic activity with the increasing operation time, in agreement with the results presented in Fig. 1d. Ba is more likely to promote the reaction of Co ions and formation of oxygen vacancy for stable ABO$_3$ perovskite type catalyst. Therefore, the reaction mechanism of NO direct decomposition over BCR catalyst is proposed in Scheme 2. NO molecule initially adsorbs on the active sites of Co species to form the mononitrosyl (NO$^*$), and then interacts with an adjacent NO$^*$ on [1] of active
sites over the active interface to produce NOON*, as discussed in literature [61]. The easy formation of NOON* decomposes into N2O* and the dissociated *O atom fills an oxygen vacancy on the surface to form active oxygen (O2*) species, verified in the in-situ DRIFT and DFT results. The high mobility of lattice oxygen promotes this process, which is evidenced by the O2-TPD and XPS results. The suitable amounts of lattice oxygen and oxygen vacancies on the surface of catalyst is beneficial to both the formation and consumption of NO*. Then the intermediates of N2O is broken with reaction energies at 1.11 eV to produce N2 and the *O atom fills the vacancy (Fig. 9b). The adjacent active *O atom to further form O2. Therefore, the decomposition of N2O* is likely the rate-determining step (RDS) in NO direct decomposition over BCR. In addition, the original lattice O16O is consumed and replaced with the formation of lattice 18O via occupying more [ ] of the catalyst after adding excess gaseous 18O2 in the isotopic (18O2) exchange experiment. Herein the competitive adsorption of NO and O2 on the [ ] mainly occurs in this step. The formed N2O* species quickly decomposes into N2 and another adsorbed oxygen (O*). Finally, the adjacent adsorbed oxygen (O*) species combine to produce O2 and release active sites, which is ready for next catalytic cycle. The production of 16O 18O is well verified with the formation and reaction of new lattice 18O in the above-mentioned isotopic (18O2) tracing results. In this reaction mechanism, not only Co species acted as main active sites, but also oxygen vacancies play an important role on the BCR active interface for NO decomposition. The intrinsic effect with O2 for NO direct decomposition shows that the suitable amount of lattice oxygen promotes the nitrite species formation at the beginning of the reaction. However, excessive amount of O2 occupies more oxygen vacancies and inhibits the NO* formation and transformation of NO* into N2O* due to the competitive adsorption of NO and O2 on the oxygen vacancies, impeding the NO direct decomposition. The mobility of lattice oxygen and timely desorption of O2 over the catalyst are crucial to further improve the catalytic activity and tolerance of O2. These results contribute to our understanding of the perovskite-support interaction and reaction mechanisms and energetics involved in the NO adsorption and reaction processes on BaCoO3/CeO2 systems, shedding light on the potential catalytic activity of these materials in NO-related reactions.

5. Conclusion

The distinct structure-activity relationships of the BaCoO3/CeO2 catalysts were investigated using CeO2 with conventional small particle (p), sphere (s) and rod (r) morphologies. The NTN2 at 800 °C decreases in the order of BCR (78.9 %) > BCS (75.9 %) > BCP (56.9 %) > BaCoO3 (8.6 %). BCR also exhibits a high durability (maintain NO conversion ~60 %) in the presence of 5 vol% O2 for 12 h. Such superior catalytic activity and O2 tolerance of BCR are attributed to the strong PSI on the oxide phases interface. The CeO2 rod enhances the sintering resistance with high surface area, redox activity, mobility of active lattice oxygen and NO adsorption capability. We proposed a reaction mechanism that

Table 2

<table>
<thead>
<tr>
<th>Wavenumber (cm−1)</th>
<th>Bond</th>
<th>Vibration</th>
<th>Adsorbed species</th>
<th>Structure</th>
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</thead>
<tbody>
<tr>
<td>2240, 2210</td>
<td>N-N</td>
<td>ν</td>
<td>N2O</td>
<td></td>
</tr>
<tr>
<td>1930, 1902, 1845</td>
<td>N-O</td>
<td>ν</td>
<td>NO on Co3+</td>
<td></td>
</tr>
<tr>
<td>1774</td>
<td>O-N</td>
<td>ν</td>
<td>Asymmetric nitric oxide dimer, (NO2) on Co3+ sites</td>
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</tr>
<tr>
<td>1608, 1044</td>
<td>N=O</td>
<td>ν</td>
<td>Bridging bidentate nitrate, (NO2)</td>
<td></td>
</tr>
<tr>
<td>1575, 1546</td>
<td>N=O</td>
<td>ν</td>
<td>Chelating bidentate nitrate, (NO2)</td>
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<tr>
<td>1520</td>
<td>O-N</td>
<td>ν</td>
<td>Monodentate nitrite (NO2) on Co3+</td>
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<tr>
<td>1408</td>
<td>N=O</td>
<td>ν</td>
<td>Monodentate nitrate, (M-NO2)</td>
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<tr>
<td>1347, 1281, 1013</td>
<td>N-O</td>
<td>ν</td>
<td>Nitrate free ion</td>
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</tr>
<tr>
<td>1238, 813</td>
<td>NO2</td>
<td>ν</td>
<td>Chelating bidentate nitrite (NO2) on CeO2</td>
<td></td>
</tr>
</tbody>
</table>

ν represents the stretching vibration; νas represents the asymmetric stretching vibration; νs represents the symmetric stretching vibration; M represents metal sites.

Fig. 8. Isotopic (18O2) exchange experiments (a) blank calibration without catalyst at room temperature, and (b) NO direct decomposition processes at 800 °C over BCR at 1 vol% NO/Ar and 1 vol% NO + 1 vol% 18O2/Ar conditions.
both Co species and adjacent oxygen vacancies act as active sites, and the decomposition of N₂O* intermediate is the RDS. Further based on the isotopic (¹⁸O₂) exchange results, the intrinsic effect of adding O₂ highlights that an appropriate amount of lattice oxygen promotes the formation of nitrite species at the initial stage of the reaction. Nevertheless, an excessive amount of O₂ occupies more oxygen vacancies and thus inhibits the NO* formation and transformation of NO* into N₂O*, due to the competitive adsorption of NO and O₂ on the oxygen vacancies of active sites, inhibiting the NO direct decomposition reaction.

**Scheme 1.** The morphology-activity relationship of catalysts.

**Scheme 2.** Reaction pathways for NO direct decomposition over BCR catalyst.

**CRediT authorship contribution statement**

**Lu Wang:** Writing – original draft, Methodology, Investigation. **Gang Wang:** Supervision, Funding acquisition. **Hong Chen:** Validation, Methodology, Investigation. **Cuijuan Zhang:** Validation, Resources, Investigation. **Running Kang:** Writing – original draft, Methodology, Investigation, Formal analysis, Data curation, Conceptualization. **Xuehai Wang:** Writing – review & editing, Resources, Project administration. **Junqin Huang:** Investigation, Data curation. **Sufeng An:** Resources, Investigation, Conceptualization. **Feng Bin:** Validation, Resources, Investigation, Data curation. **Yongdan Li:** Writing – review &
Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

No data was used for the research described in the article.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.apcatb.2024.123952.

References


