Catalytic decomposition of $\text{N}_2\text{O}$ over $\text{Cu}–\text{Al}–\text{O}_x$ mixed metal oxides

Magdalena Jabłońska, a,b,⁎ Miren Agote Arán, c Andrew M. Beale, d,⁎⁎ Kinga Góra-Marek, e Gérard Delahay, f Carolina Petitto, f Kateřina Pacultová f and Regina Palkovits, f a,b,⁎

Cu–Al–O mixed metal oxides with intended molar ratios of Cu/Al = 85/15, 78/22, 75/25, 60/30, were prepared by thermal decomposition of precursors at 600 °C and tested for the decomposition of nitrous oxide (deN$_2$O). Techniques such as XRD, ICP-MS, $\text{N}_2$ physisorption, O$_2$-TPD, H$_2$-TPR, in situ FT-IR and XAFS were used to characterize the obtained materials. Physico-chemical characterization revealed the formation of mixed metal oxides characterized by different specific surface area and thus, different surface oxygen default sites. The O$_2$-TPD results gained for Cu–Al–O mixed metal oxides conform closely to the catalytic reaction data. In situ FT-IR studies allowed detecting the form of Cu$^{+}$$+$$\text{N}_2$ complexes due to the adsorption of nitrogen, i.e. the product in the reaction between N$_2$O and copper lattice oxygen. On the other hand, mostly nitrate species and NO were detected but those species were attributed to the residue from catalyst synthesis.

1. Introduction

Nitrous oxide ($\text{N}_2\text{O}$) significantly contributes to the greenhouse effect and ozone destruction in the stratosphere. The main anthropogenic source of $\text{N}_2\text{O}$ is nitric acid production (about 1% of all greenhouse gas emission).1 The catalytic decomposition of $\text{N}_2\text{O}$ up to 450–500 °C provides an attractive solution for reducing $\text{N}_2\text{O}$ emissions in tail gas from the point of both application and operation costs, respectively.2 To control the emission of $\text{N}_2\text{O}$, many catalysts have been reported for the catalytic decomposition of $\text{N}_2\text{O}$, including supported metals, pure and mixed oxides, and zeolites, etc.3 Among them, mixed oxides containing cobalt spinels revealed excellent activity in deN$_2$O, however, cobalt toxicity represents another serious issue. On the other side, copper-based materials represent one of the classes of catalysts dedicated to $\text{N}_2\text{O}$ decomposition with their low cost and high catalytic activity. E.g. commercial and mesoporous CuO,3 CuO supported on different carriers, such as Al$_2$O$_3$, ZnAl$_2$O$_4$, ZrO$_2$ [e.g. ref. 4 and 5], Cu-containing zeolites [e.g. ref. 6], CuO–CeO$_2$ [e.g. ref. 7] were investigated for $\text{N}_2\text{O}$ decomposition. Several works aimed to study $\text{N}_2\text{O}$ decomposition over hydrotalcite derived mixed metal oxides. In particular, Chmielarz et al.8 reported significantly higher activity of Cu–Mg-Al–O$_x$ hydrotalcite derived mixed metal oxides than analogues Co–Mg-Al–O$_x$. Cu–Mg–Al–O$_x$ with molar ratios of Cu/MgAl = 10/61/29 and calcined at 600 °C, reached full N$_2$O conversion at 600 °C in the presence of O$_2$ (0.1 g catalyst, 0.5 vol% N$_2$O/He, 4.5 vol% O$_2$/He, 50 cm$^3$ min$^{-1}$). Kannan9 found 48% N$_2$O conversion at 450 °C over Cu–Al–O$_x$ (Cu/Al = 3/1, mol. ratio, 0.1 g catalyst, 0.0985 vol% N$_2$O/He, 100 cm$^3$ min$^{-1}$). Co-containing catalysts were more active than corresponding Cu-systems (84 versus 48% at 450 °C). The best results were obtained for the Co–Al–O$_x$ catalyst with a Co/Al molar ratio of 3/1 among ratios of 3–1/1. Pan10 tested samples with different Cu/Al molar ratios (2–4/1) in the presence of O$_2$ (1.0 g catalyst, 2.0 vol% N$_2$O/Ar, 4.0 vol% O$_2$/Ar, 140 cm$^3$ min$^{-1}$) and also found out that Cu/Al = 3.1/1, mol. ratio, was the most active. However, no simple correlation between activity and Cu/Al ratio was obtained and deeper insight into the effect of material composition would be desirable. Thus, the above studies motivated us to prepare Cu–Al–O$_x$ mixed metal oxides and explore the effects of different molar ratios of used metals (Cu/Al = 85/15, 78/22, 75/25, 60/30). We investigated the relationship between the physicochemical properties and the catalytic activity in deN$_2$O over Cu–Al–O$_x$ mixed metal oxides using XRD, ICP-MS, N$_2$ physisorption, O$_2$-TPD, H$_2$-TPR, in situ...
FT-IR and EXAFS combined with microreactor catalytic tests. The in situ FT-IR experiments allowed monitoring the surface nitrogen groups, while application of in situ EXAFS revealed the oxidation/coordination state of copper oxides species of Cu–Al–Ox mixed metal oxides during deN₂O.

2. Experimental

2.1. Catalyst preparation

A series of Cu–Al precursors with intended molar ratios of Cu/Al = 85/15, 78/22, 75/25, 60/30 were prepared by coprecipitation. An aqueous solution containing appropriate amounts of Cu(NO₃)₂·3H₂O (Sigma), Al(NO₃)₃·9H₂O and 1 M NaOH (Chemsolute) was dropped simultaneously to a vigorously stirred aqueous solution containing a slight over-stoichiometric (Chemsolute) was dropped simultaneously to a vigorously stirred aqueous solution containing a slight over-stoichiometric excess of Na₂CO₃ (Sigma) at 60 °C. The pH of the reaction mixture was maintained constant at 10.0 ± 0.2 throughout the whole synthesis by NaOH addition. The obtained suspension was aged at 60 °C for another 0.5 h after complete coprecipitation. The solid was filtered, washed carefully with distilled water and dried at room temperature. Finally, the prepared precursors were crushed and calcined at 600 °C for 6 h with a heating ramp of 10 °C min⁻¹ and in static air. For catalytic experiments, a fraction of particle size in the range of 0.250–0.500 mm was used.

2.2. Catalyst characterization

The X-ray diffraction (XRD) analysis of the mixed metal oxides was performed applying a Siemens D5000 XRD diffractometer using Cu-Kz radiation (λ = 1.54056 Å, 45 kV, 40 mA).

The chemical composition of mixed metal oxides was determined by ICP-MS using an Agilent Technologies 8800 Triple Quad spectrometer. Prior to the measurement, the sample [50 mg] was dissolved in 6 cm³ mixture of concentrated acids (HCl : HNO₃ : H₂O = 1 : 1), and afterwards, the resulting mixture was diluted with 64 cm³ deionized water before warming up to 40 °C for 24 h.

The specific surface area (S_BET) of the mixed metal oxides was determined by low-temperature (−196 °C) N₂ sorption using a Quantachrome Quadrasorb SI. Prior to nitrogen adsorption, the samples were outgassed at 250 °C for 12 h using a Quantachrome Fluvac degasser. The specific surface area (S_BET) was calculated using the Brunauer–Emmett–Teller (BET) multiple point method in the p/p₀ range from 0.05 to 0.3.

The temperature-programmed desorption of O₂ (O₂-TPD) was performed to investigate oxygen desorption behaviour using a Micromeritics AUTOCHEM 2910. 49–58 mg of sample was loaded into a quartz tube and pretreated in reconstituted air (20.0 vol% O₂/80.0 vol% N₂) at 600 °C (30 cm³ min⁻¹). After cooling down to 80 °C, the pretreated sample was heated up to 600 °C with a linear heating rate of 10 °C min⁻¹ in a carrier gas of He (15 cm³ min⁻¹) and kept for 10 min at 600 °C and cooled down up to 80 °C in the same flow (He). Then this material was heated, under a flow of 1.0 vol% N₂O/He (30 cm³ min⁻¹), at 600 °C [plateau 10 min] and cooled to 80 °C. Afterward the second run of O₂-TPD was performed under He flow (15 cm³ min⁻¹). O₂-TPD was also carried out (15 cm³ min⁻¹ of He, 10 K min⁻¹, 80–600 °C, 10 min at 600 °C) over 52–56 mg of sample after heating (up to 600 °C, plateau 10 min) and cooling down (80 °C) in the presence of 1.0 vol% N₂O/He (30 cm³ min⁻¹).

The redox properties of the mixed metal oxides were studied by the temperature-programmed reduction (H₂-TPR) using Quantachrome ChemBET Pulsar TPR/TPD. H₂-TPR runs for the samples (50 mg) were carried out starting from room temperature to 1000 °C, with a linear heating rate of 5 °C min⁻¹ and in a flow of 5.0 vol% H₂/Ar (25 cm³ min⁻¹). Water vapour was removed from the effluent gas by the means of a cold trap placed in an ice-water bath. The H₂ consumption was detected and recorded by a TCD detector.

2.3. Catalytic tests

Steady-state catalytic measurements of N₂O decomposition were performed in an integral fixed bed stainless steel reactor of 5 mm internal diameter in the temperature range of 300–450 °C under atmospheric pressure. The space velocity (SV) of 30 or 60 l h⁻¹ (20 °C, 101325 Pa) was applied. The inlet gas contained 0.1 vol% of N₂O in N₂ as balance. O₂ (5.0 vol%) and H₂O (2.5 vol%) were added to feed in order to simulate real waste gas from nitric acid plants. Before the first catalytic run, the catalyst was pre-treated in N₂ flow at 450 °C for 1 h. Then the catalyst was cooled to the reaction temperature, the steady-state of N₂O concentration level was measured and used for calculation of N₂O conversion. An infrared analyser N₂O (GMS 810 Series Sick) was used to analyse N₂O. The conversion of N₂O (X(N₂O)) was determined according to X(N₂O) = ([c(N₂O)in − c(N₂O)out])/c(N₂O)in × 100%, where: c(N₂O)in and c(N₂O)out – concentration of N₂O in the inlet gas, and concentration of N₂O in the outlet gas.

2.4. In situ experiments

The X-ray absorption spectra (XAS) of selected mixed metal oxides were performed in situ using the quartz capillary flow reactor cells, and gas delivery systems available on the beamline, on station B18 at the Diamond Light Source synchrotron facility. The measurements were carried out using a Si (111) monochromator at the Cu K-edge, Cu foil (10 μm) was used as an energy calibrant. The catalyst diluted with SiO₂ (1 : 5) was sieved into 0.200–0.250 mm and placed into the capillary reactor (with an internal diameter of 3 mm). Prior to the reaction, the catalyst was outgassed at 600 °C for 1 h in a flow of pure He (10 cm³ min⁻¹), and subsequently cooled down to 100 °C. The reactant concentrations at the reactor inlet were composed of [N₂O] = 0.1 vol% and [He] = 99.9 vol% (10 cm³ min⁻¹). The temperature was raised in steps of 50–100 °C up to 450 °C and each temperature was held for 30 min. X-ray absorption spectra at Cu K-edge for all the samples were collected in transmission with the exception of the catalyst with Cu/Al = 60/30, mol. ratio, which was collected in fluorescence mode. At least 3 spectra for each sample were taken at room temperature, appropriate temperatures and after reaction at room temperature. CuO reference was measured at room temperature in pellet form.
oxides were confirmed by elemental analysis and varied to some extent with respect to the desired ratios (Table 1). In particular, Cu/Al molar ratios were significantly lower than expected from synthesis for materials with Cu/Al > 3.0, mol. ratio. The chemical analysis identified by ICP-MS evidenced sodium residual from the preparation procedure up to 1.1 wt% for Cu/Al = 78/22, mol. ratio.

Cu-Al-Ox mixed metal oxides show specific surface area (S_{BET}) in the range of 34–84 m² g⁻¹. S_{BET} varied between materials with different molar ratios. While no correlation existed between copper content (wt%) and specific surface area, the specific surface area increased with decreasing Cu/Al molar ratio. Fig. 2 presents O₂ desorption rates evaluated for the pretreated materials under reconstituted air and subsequently reoxidized by 1.0 vol% N₂O/He, or for pretreated materials under 1.0 vol% N₂O/He. The O₂-TPD profiles of Cu-Al-Ox mixed metal oxides showed peaks related to the desorption of surface oxygen species around 150–400 °C, whilst the peak above 400 °C is attributed to the desorption of lattice oxygen. The O₂-TPD profiles are dominated by high-temperature peaks (Table 1), indicating that the increasing Cu/Al molar ratios decreased the molar amount of desorbed lattice oxygen. Also, the presence of copper in Cu₇₅Al₂₅O and Cu₆₀Al₃₀O led to a lower amount of desorbed O₂ compared to Co₃O₄. Nevertheless, the quantity of oxygen desorbed seems to increase with the specific surface area. It should be emphasized that the amount of oxygen desorbed per gram of catalyst is negligible compared to the amount of oxygen consumed by hydrogen during the TPR (see below H₂-TPR results). This point seems to indicate that the desorbed oxygen comes rather from the surface of the material.

The H₂-TPR profiles of Cu-Al-Ox mixed metal oxides revealed one main broad peak between 200 and 400 °C corresponding to the reduction of bulk copper oxide species to metallic copper, as shown in Fig. 3. The shape of peak maxima and H₂ uptake (Table 1) obtained for mixed metal oxides matched to that of pure CuO (maximum at about 350 °C, H₂ uptake of 10.7 mmol g⁻¹). According to the XRD analysis, the peaks associated with the CuO were the main peaks observed in the mixed metal oxides. Otherwise, for Cu₅₇Al₃₅O and Cu₆₀Al₃₀O the reduction of copper in CuAl₂O₄ cannot be excluded. A quantitative analysis of H₂ consumption based on integrating

### Table 1: Theoretical and determined molar ratios, specific surface area (S_{BET}), molecular amount of O₂ desorbed from O₂-TPD measurements and H₂ uptake of the Cu-Al–Ox mixed metal oxides

<table>
<thead>
<tr>
<th>Cu-Al-Ox</th>
<th>CuO particle size (nm)</th>
<th>Cu/Al molar ratio</th>
<th>Cu/wt%</th>
<th>Na content (wt%)</th>
<th>S_{BET} (m² g⁻¹)</th>
<th>H₂ uptake (mmol g⁻¹)</th>
<th>O₂(des) (µmol g⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>60/30</td>
<td>15</td>
<td>Theoretical</td>
<td>Determined</td>
<td>48.4</td>
<td>84</td>
<td>7.8</td>
<td>46.9/27.3/36.3</td>
</tr>
<tr>
<td>75/25</td>
<td>14</td>
<td>Theoretical</td>
<td>Determined</td>
<td>55.0</td>
<td>68</td>
<td>8.4</td>
<td>32.1/24.9/25.1</td>
</tr>
<tr>
<td>78/22</td>
<td>17</td>
<td>Theoretical</td>
<td>Determined</td>
<td>41.8</td>
<td>70</td>
<td>9.4</td>
<td>30.1/17.5/17.5</td>
</tr>
<tr>
<td>85/15</td>
<td>21</td>
<td>Theoretical</td>
<td>Determined</td>
<td>51.3</td>
<td>34</td>
<td>9.4</td>
<td>26.2/17.5/14.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Determined</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Estimated by the Scherrer’s formula for (111) reflection. * Determined with ICP-MS analysis. * Calculated by the equation: Y = 9 \times 10^{-9}X + 2 \times 10^{-7}, R² = 0.9996, and X, Y referred to the area of each reduction peak and the H₂ consumption, respectively. * Estimated from direct O₂-TPD of known amount of as-stored Ag₂O (Stream Chemicals). O₂ desorbed during O₂-TPD for pretreated materials under reconstituted air (20.0 vol% O₂ / 80.0 vol% N₂), subsequently reoxidized by 1.0 vol% N₂O/He (f), or for pretreated materials under 1.0 vol% N₂O/He (g).
the H2-TPR curves confirmed that H2 uptake did not change significantly over mixed metal oxides (7.8–9.4 mmol g\(^{-1}\)).

Fig. 4 shows the results of N\(_2\)O decomposition over Cu-Al-O\(_x\) mixed metal oxides with varying Cu/Al molar ratios. The effect of sample composition on the activity was studied to find the optimum materials for maximum conversion. The highest activity among the tested catalysts reached material with a molar ratio Cu/Al = 60/30 (conversion of 25% at 450 °C). The other catalysts reached significantly lower activities (below 20% at 450 °C). Comparable results were obtained by Kannan,\(^9\) who found a conversion of 48% at 450 °C over Cu-Al-O\(_x\) (Cu/Al = 3/1 mol. ratio, 0.1 g catalyst, 0.0985 vol% N\(_2\)O/He, 100 cm\(^3\) min\(^{-1}\)). The most active material was also tested in simulated waste gas conditions – in the presence of 5.0 vol% O\(_2\) and 2.0 vol% H\(_2\)O. O\(_2\) present in the feed caused a steep drop in conversion; what is important, the inhibiting effect was fully reversible. The stepwise addition of H\(_2\)O (results not shown) had a detrimental effect on N\(_2\)O conversion probably due to the strong adsorption of the water molecule on the surface, since the initial activity in inert gas was not fully recovered after removal of O\(_2\) and H\(_2\)O. Table 2 lists examples of Cu-containing catalyst tested for N\(_2\)O decomposition. It can be found that either (supported) copper oxide or hydrotalcite derived mixed metal oxides are inherently not active in deN\(_2\)O. Further modification of Cu-containing material with noble/rare earth metals can significantly improve their catalytic activity [e.g. ref. 1 and 9].
The most active catalyst (Cu/Al = 60/30, mol. ratio) was studied by in situ EXAFS during N₂O decomposition. The Cu K-edge absorption spectra were collected at room temperature, during the temperature ramp (100 to 450 °C) under N₂O/He feed, and finally at room temperature after deN₂O. The Cu K-edge EXAFS spectra for the catalyst collected at different stages of the experiment (Fig. 5A) presents similar phases as the CuO reference; in line with the XRD results, suggesting CuO as the main species discernible in the samples. The reduced EXAFS oscillation amplitudes in Cu₆₀Al₃₀Oₓ can be attributed to differences in sample measuring conditions: CuO was measured in pellet form while the catalyst was measured as granulated powder in a microreactor, inhomogeneities and pinhole effects in the latter case resulted in EXAFS amplitude reduction. No significant changes in the Cu₆₀Al₃₀Oₓ spectra were observed during activation or N₂O exposure indicating the CuO local structure remains constant.

The Fourier transform of the X-ray absorption fine structure (FT-EXAFS) spectra of the catalyst in Fig. 5B gives insight into the distance of neighboring atoms around the absorber atom. The peaks with maxima around 0.19 and 0.29 nm correspond to neighbor O and Cu atoms, respectively. A decrease in peak intensity was observed for the data collected at increasing temperatures. This decrease is attributed to the increasing atom vibration due to thermal effects which smear out the EXAFS oscillations affecting the signal intensity in the Fourier transform. The position of the peaks did not vary significantly throughout the experiment up to 250 °C evidencing no changes in bond distance or local Cu geometry in the sample measured at 450 °C. When the catalyst is active, there seems to be a slight shift to shorter Cu–O distance which could suggest the formation of Cu²⁺. Nonetheless, a definite conclusion cannot be drawn here as no changes were discernible in the position of the rising absorption edge in the XANES to account for Cu²⁺ reduction to Cu⁺.

For FT-IR, the selected materials – Cu₆₀Al₃₀Oₓ and Cu₈₅−

Al₁₅Oₓ (the most active and less active sample in deN₂O, respectively) studies were contacted with the N₂O dose (60 Tr of N₂O per 10 mg of the sample) at room temperature, then the IR spectrum was collected (Fig. 6). Next, the catalysts were heated to 330 °C, kept for 10 min and then cooled down to RT to collect IR spectrum of all the reaction products. The procedure was repeated for 390 and 450 °C. An intense absorption band around 2220–2230 cm⁻¹ recorded at room temperature appeared due to adsorbed N₂O. In the spectra recorded at higher temperatures, bands appeared below 1700 cm⁻¹ due to the appearance of nitrites, nitrates and nitro compounds. Also, the gaseous NO can be easily identified in the IR spectra due to

![Fig. 5](image_url)  
**Fig. 5** Cu K-edge absorption spectra acquired during the *in situ* experiments: EXAFS (A) and FT-EXAFS (B) spectra (phase corrected) for material with Cu/Al = 60/30, mol. ratio, and CuO reference at room temperature.
the vibration-rotation bands at 1950–1850 cm \(^{-1}\). Nitric oxide was found to play a crucial role in the kinetic oscillations caused by a complex interaction of different reactions. Both NO and molecular oxygen can be formed by the decomposition of the residual nitrate species from the catalyst synthesis. \(^{17}\) It is generally accepted that nitrate species on Cu-ZSM-5 and other zeolite catalysts modified with copper are stable even at high temperatures. Furthermore, nitrate moieties have been proposed to be important intermediates in the decomposition of NO, \(^{18}\) the selective catalytic reduction of NO by hydrocarbons, \(^{19}\) and the SCR of NO by ammonia. \(^{20}\) The spectra of Cu–Al–O\(_x\) mixed metal oxides were consistent with the formation of bridged nitrates at 1674 cm \(^{-1}\), surface nitrates at 1614 and 1576 cm \(^{-1}\), monodentate nitrates at 1416 cm \(^{-1}\), nitro groups at 1353 cm \(^{-1}\), as well as bidentate nitrates at 1312 cm \(^{-1}\). \(^{21,22}\)

For the most active Cu\(_{60}\)Al\(_{30}\)O\(_x\), the existence of N-species implied a reaction between N\(_2\)O and copper lattice oxygen. What is more, the formation of end-product of the reaction, i.e. N\(_2\) molecule was strongly supported by the presence of 2301 cm \(^{-1}\) band attributed to Cu\(^+\)/N\(_2\) complexes. \(^{23}\) N\(_2\) molecule interacting with Cu\(^+\) ions inside zeolites leads to a 1(N≡N) band in the 2300–2290 cm \(^{-1}\) region, which is significantly downshifted with respect to gas-phase value of 2321 cm \(^{-1}\). This bathochromic shift could be explained in terms of chemical interactions involving molecular orbitals of the probe molecule and a suitable d-orbital of the metal cation. \(^{24}\) Basing on the wavenumber and half-bandwidth of 2301 cm \(^{-1}\) band identical to the band of Cu\(^+\)–N\(_2\) adducts in Cu-zeolites we advocated on the formation of this adducts also in Cu–Al–O\(_x\) mixed metal oxides. The presence of stable Cu\(^+\)–N\(_2\) complexes indicated the co-presence of both Cu\(^+\) and Cu\(^{2+}\) cations on the catalyst surface, as expected for a redox mechanism requiring a balance between these two sites. It is also in line with the CO sorption results on spent material Cu\(_{60}\)Al\(_{30}\)O\(_x\) (spectra not shown) which indicated that the surface Cu is poorer in the Cu\(^{2+}\) cationic species. Such observation allows for concluding on the efficient catalyst which should be characterized by the high redox ability to reduce Cu\(^{2+}\)/Cu\(^+\) redox pair. Furthermore, in this particular case, Cu\(^+\) cations are not able to efficiently transform back to a Cu\(^{2+}\) state. The poorest activity of Cu\(_{85}\)Al\(_{15}\)O\(_x\) was evidenced as no completed decomposition of N\(_2\)O (the 2224 cm \(^{-1}\) band) could be achieved even at a temperature as high as 450 °C (Fig. 6). The activity varied among the tested materials, however, no clear trend related either to the Cu or Na content became evident. While alkali metals can act as basic centres and significantly influence catalytic activity, \(^{24}\) for example, Obalová et al. \(^{25}\) pointed out that 1.15 wt% of Na introduced by impregnation already slightly enhanced activity of Co\(_4\)MnAlO\(_x\). In our case, Na residual remained a preparation procedure, and actually with lower values than 1.15 wt% (Table 1). At this stage, it is not possible to precisely justify the influence of residual Na on materials catalytic activity in deN\(_2\)O.

N\(_2\)O decomposition on Cu–Al–O\(_x\) yielded N\(_2\) and O\(_2\), and the reoxidation of copper side took place with gas phase N\(_2\)O (eqn (1)):
\[
\text{Cu}^* + \text{N}_2\text{O} \rightarrow \text{Cu}^*\text{O} + \text{N}_2. \quad (1)
\]

In the decomposition of \(\text{N}_2\text{O}\) by an oxidation–reduction mechanism, the desorption of adsorbed \(\text{O}_2\) is the rate-determining step of the \(\text{N}_2\text{O}\) decomposition.\(^{15,26}\) Consequently, materials, that possess better mobility of lattice oxygen, can promote desorption of \(\text{O}_2\) and regeneration of the active sites (eqn (2)):

\[
2\text{Cu}^*\text{O} \rightarrow 2\text{Cu}^* + \text{O}_2. \quad (2)
\]

Moreover, a larger specific surface area may promote this specificity since a higher specific surface area may generate more surface oxygen defaults sites. The \(\text{O}_2\)-TPD results correspond closely to the results of \(\text{N}_2\text{O}\) decomposition over \(\text{Cu–Al}\) oxides for \(\text{N}_2\text{O}\) decomposition.

### 4. Conclusions

In this study, a series of \(\text{Cu–Al–O}_x\) mixed metal oxides with different molar ratios (\(\text{Cu/Al = 85/15, 78/22, 75/25, 60/30, mol. ratio}\) was successfully obtained by coprecipitation, followed by their thermal decomposition. The catalysts were investigated in \(\text{N}_2\text{O}\) decomposition confirming that the activity is strongly dependent on the \(\text{Cu/Al}\) molar ratio. Decreasing the amount of \(\text{Cu/Al}\) molar ratio in \(\text{Cu–Al–O}_x\) leads to the most active catalysts in an order as follows (\(\text{Cu/Al, mol. ratio}: 60/30 > 75/25 > 78/22 > 85/15\)). The highest activity of \(\text{Cu/Al = 60/30, mol. ratio, systems appeared possibly due to its high oxygen mobility combined with its specific surface. Further research is carried out in order to clarify the promotion effects of alkali and rare earth metals on \(\text{Cu–Al–O}_x\) mixed metal oxides for \(\text{N}_2\text{O}\) decomposition.}

### Conflicts of interest

There is no conflicts to declare.

### Acknowledgements

The authors acknowledge the Federal Ministry of Education and Research (BMBF) for funding in the frame “Material Innovations” within the research project “Efficient DeNOx-strategies for lean-operated combustion engines” (BMBF-PTJ FKz 13XP5042A). This work was partially funded by the Excellence Initiative of the German Federal and State Governments in the frame of the Center for Automotive Catalytic Systems Aachen (ACA) at RWTH Aachen University by the Grant No. 2015/18/E/ST4/00191 from the National Science Centre, Poland and by EU structural funding in Operational Programme Research, Development, and Education, project No. CZ.02.1.01/0.0/0.0/16_019/0000853 “IET-ER”. The authors acknowledge the Diamond Light Source (project SP14834) for the provision of beamtime on the beamlines B18, and Diego Gianolio for assistance in performing the XAFS measurements.

### References

4. R. Zhang, C. Hua, B. Wang and Y. Jiang, \(\text{N}_2\text{O}\) decomposition over \(\text{Cu–Zn}/\gamma\text{-Al}_2\text{O}_3\) catalysts, Catalysts, 2016, 6, 200.
16. T. Franken and R. Palkovits, Investigation of potassium doped mixed spinels \(\text{Cu}_{x}\text{Co}_{3-x}\text{O}_4\) as catalysts for an


22 S. Parres-Escalazp, I. Such-Basañez, M. J. Illán-Gómez, C. S.-M. de Lececa and A. Bueno-López, Study by isotopic gases and \textit{in situ} spectroscopies (DRIFTS, XPS and Raman) of the N\textsubscript{2}O decomposition mechanism on Rh/CeO\textsubscript{2} and Rh/γ-Al\textsubscript{2}O\textsubscript{3} catalysts, \textit{J. Catal.}, 2010, \textbf{276}, 390–401.


