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# High pressure ammonia decomposition on Ru-K/ CaO catalysts†

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Potassium-promoted ruthenium supported on CaO is a very efficient catalyst for ammonia decomposition, surpassing the performance of other Ru-supported solids. At an optimum Ru loading of 3% wt, catalysts with a K/Ru atomic ratio of 0.9 showed the best catalytic performance under a wide range of operating conditions, P = 1-40 bar, T = 250-550 °C and WHSV = 9000-30000 mL g<sup>-1</sup> h<sup>-1</sup>. Although NH<sub>3</sub> conversion levels decrease considerably upon increasing the reaction pressure ( $X_{550}$   $_{\circ C,~40}$   $_{\text{bar}}$  = 0.8), high pressure ammonia decomposition offers the possibility of CO<sub>v</sub>-free compressed hydrogen and hydrogen productivities and TOFs 40 times bigger than when applying atmospheric pressure. Extensive characterization by CO chemisorption and HR-TEM demonstrates that potassium promotion increases metal dispersion by decreasing the Ru particle size. Electronic effects derived from the close proximity between K and Ru result in a decrease in the reaction apparent activation energy, as shown by a detailed kinetic analysis.

# Introduction

The use of hydrocarbons as energy carriers has facilitated human development over the last century. However, the derived environmental consequences and scarcity issues are a strong driving force to look for cleaner and more sustainable alternatives. 1-3 Among the different possibilities, the use of hydrogen offers great advantages<sup>4-7</sup> especially if its production8-12 is accompanied by CO2 capture or if it is directly produced from water.

Hydrogen storage, however, is still an issue for which several alternatives are being proposed: metal hydrides, such as NaAlH<sub>4</sub> and LiAlH<sub>4</sub>, <sup>13,14</sup> or methanol <sup>15,16</sup> are among the most studied, but there is also growing interest in the use of ammonia. NH3 offers several advantages: high volumetric (108 kg H<sub>2</sub> per m<sup>3</sup> NH<sub>3</sub> at 20 °C and 8.6 bar) and gravimetric energy density (17.8 wt%), ease of storage and transport, and the fact that NH3 production is a very well established technology (global production of 176 million metric tons in 2014 (ref. 17-21)); these all point to ammonia as a very promising energy carrier readily compatible with the current distribution network. Moreover, ammonia decomposition is

economically more viable than reforming processes, including methanol reforming,  $^{22,23}$  and hydrogen and  $N_{\rm 2}$  are the only decomposition products, in contrast to the coproduction of CO, CO2, and CH4 from other liquid H2 carriers.24,25 Last but not least, this reaction can be carried out at lower temperatures (450 °C or below) than reforming processes (above 800 °C). 17,26,27

Ammonia decomposition proceeds through stepwise dehydrogenation followed by recombination of two N and two H adatoms to form N2 and H2 followed by desorption, and it is thought that NH cleavage is the rate limiting step. Ru is at the top of the volcano-type relationship observed for the classical ammonia decomposition rate vs. nitrogen binding enthalpy plot.<sup>28-30</sup> Consequently, the best ammonia decomposition catalysts known to date are based on Ru. K-promoted catalysts supported on either MgO or carbon nanotubes (CNTs) are among the most active catalysts for this reaction. 31-37 The nature of the support has been shown to strongly influence the catalytic performance of Ru-based catalysts. Widely reported is the use of basic supports like MgO and La<sub>2</sub>O<sub>3</sub>, 33,38,39 neutral supports, such as CNTs, 40,41 and acidic supports, like Al<sub>2</sub>O<sub>3</sub>. 31,32,35,42 In general, the use of basic and neutral supports results in better catalytic properties.43 In addition, the use of alkali promoters has been shown to further decrease the nanoparticle size and the associative desorption of nitrogen, decreasing the activation energy of the reaction.44,45

As can be derived from the previous paragraphs, ammonia decomposition is a relatively well-known process. However,

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for most potential applications of ammonia derived H<sub>2</sub> (i.e. fuel cells), hydrogen should be supplied at elevated pressures, or separated from nitrogen or stored at high pressure. Therefore, in order to avoid additional compression costs<sup>46-48</sup> and derived emissions (estimates are about 6.0 kW h kg<sup>-1</sup> for compression of H<sub>2</sub> to 70 MPa, which leads to approximately 1.3 kg of CO<sub>2</sub> per kg of hydrogen), <sup>49</sup> high pressure ammonia decomposition would be preferred, in spite of the obvious thermodynamic limitations.<sup>50,51</sup> Surprisingly, to the best of our knowledge, this is among the first publications on this topic.

Here we demonstrate the potential and limitations of Rubased catalysts for the high-pressure decomposition of ammonia. In a first step, by performing experiments at atmospheric pressure, CaO is found to be the most adequate support for Ru. Successive promotion with K results in further improvements in the catalytic performance. Finally, the optimized catalyst is studied at different pressures. Extensive characterization by CO chemisorption and HR-TEM demonstrates that potassium promotion increases metal dispersion by decreasing the Ru particle size. Electronic effects derived from the close proximity between K and Ru result in a decrease in the reaction apparent activation energy, as shown by a detailed kinetic analysis.

# **Experimental**

### Catalyst synthesis

(MWCNTs, Commercial multiwall carbon nanotubes Aldrich), magnesium oxide (Sigma-Aldrich), natural sepiolite (NS) (Aldrich) and calcium oxide (Sigma-Aldrich) were used as supports.

Rehydrated ruthenium(III) chloride (Aldrich) was selected as the Ru precursor, and it was incorporated onto the different supports by incipient wetness impregnation using acetone, in order to achieve different Ru loadings (from 1 to 7 wt%). After drying at 60 °C for 3 hours and a subsequent thermal treatment under argon at 500 °C for 3 hours, a potassium promoter was introduced by incipient wetness impregnation using KOH in ethanol, to reach the desired K loading (0 to 15 wt%). Finally, a second thermal treatment under Ar at 500 °C for 3 h was performed.

#### **Catalyst characterization**

Nitrogen adsorption and desorption isotherms were recorded on a Micromeritics ASAP 2040 system at 77 K. The samples were previously evacuated at 373 K for 16 h. The Brunauer-Emmett-Teller (BET) method was used to calculate the specific surface area. The  $P/P_0$  range for the BET analysis was  $0.067 < P/P_0 < 0.249$ .

X-ray diffraction (XRD) patterns were obtained using a Bruker D8 instrument in the Bragg-Brentano configuration using Cu Ka radiation. The diffractograms were scanned with a step size of 0.02° in the  $2\theta$  range of 10–90°. The crystalline phase was identified by comparison with the Joint Committee on Powder Diffraction Standards (JCPDS).

CO pulse chemisorption studies were carried out on a Micromeritics ASAP 2920 analyzer. The samples were previously reduced with a 50%H<sub>2</sub>/50%Ar gas mixture from room temperature to 500 °C, and this temperature was kept for two hours. After the reduction, the samples were cooled down to 35 °C, and treated with several CO pulses (10%CO/ 90%He). For further calculations, a stoichiometric factor Ru/ CO = 1 and a spherical geometry for the ruthenium metal particles were assumed.

Inductively coupled plasma-optical emission spectrometry (ICP-OES) was used to analyze the composition and metal loadings of the catalysts on a Thermo-Electron 3580 instrument. Complete digestion of the powders was achieved at 240 °C and 35 bar using an UltraWAVE apparatus (Milestone) and aqua regia in a ratio of 1 mg of catalyst: 1 mL of aqua regia.

Transmission electron microscopy (TEM) of the samples was performed with a Titan Themis-Z microscope from Thermo-Fisher Scientific operated at an accelerating voltage of 300 kV and a beam current of 0.5 mA. Dark-field imaging was performed on a scanning TEM (STEM) coupled to a highangle annular dark-field (HAADF) detector. Furthermore, a high throughput X-ray energy dispersive spectrometer (EDS) was also utilized in conjunction with DF-STEM imaging to acquire STEM-EDS spectrum-imaging datasets. During the acquisition of these datasets, at every image pixel, a corresponding EDS spectrum was also acquired to simultaneously generate the elemental maps of Ru, O, K, and Ca atoms. It is also pertinent to note herein that the spectrum-imaging datasets were acquired in the so-called frame mode, in which the electron beam was allowed to dwell at each pixel for only a few microseconds in order to keep the total frame time to 6 s or less. Both imaging and spectroscopy datasets for each sample were acquired as well as analyzed with a newly developed software package called Velox from Thermo-Fisher Scientific. The elemental maps for Ru, O, K and Ca atoms were computed using the extracted intensities of their respective Ka lines after background subtraction. The generated maps were slightly post-filtered by applying a Gaussian filter ( $\sigma = 0.5$ ).

#### Catalytic testing

Ammonia decomposition catalytic tests were carried out in a PID Microactivity Reference system, using a continuous fixed bed stainless steel reactor coated with alumina to avoid any activity of the reactor. Prior to the activity measurement, the catalysts (200 mg pelletized between 300 µm and 500 µm and diluted with 1 g of SiC) were reduced/activated in situ with hydrogen (25 mL min<sup>-1</sup>) at 500 °C for 3 h. The catalytic performance was evaluated at different temperatures in the range of 150-550 °C, achieving 7 h of total reaction time. For the experiments at atmospheric pressure, ammonia in the gas phase (30-100 mL min<sup>-1</sup>, WHSV = 9000-30000 mL  $g_{cat}^{-1}$ h<sup>-1</sup>) was flowed, using a mass flow controller, over the catalytic bed. For the experiments at high pressure (10-40

bar), a liquefied feeding system was used pumping liquid ammonia with an HPLC Gilson pump (30 N mL min<sup>-1</sup>, WHSV = 9000 mL g<sub>cat</sub><sup>-1</sup> h<sup>-1</sup>) through a preheater at 100 °C, in order to expand the liquid ammonia into gas before reaching the catalytic bed, where the operating pressure was automatically controlled using a back pressure regulator.

Reaction products (nitrogen, hydrogen and ammonia) were analyzed under isothermal conditions with an online gas chromatograph (3000A Micro-GC gas analyzer, Agilent) using helium (1 ml min<sup>-1</sup>) as the internal standard for quantitative analysis. The Micro-GC is equipped with two columns: a PLOTU precolumn/Molsieve column using argon as the carrier gas for nitrogen and hydrogen and a PLOTU column with helium carrier gas for ammonia. Both columns are equipped with thermal conductivity detectors.

# Results and discussion

#### Catalyst characterization

The BET area and ICP results are shown in Table 1. Very small specific surface areas were observed for all the catalysts based on CaO (in the range of 4-10 m<sup>2</sup> g<sup>-1</sup>), independent of Ru and K loadings while moderate to very high surface areas were obtained using other support materials (see MWCNTs or natural sepiolite). The ICP results are in good agreement with the nominal composition for all the samples, corroborating the successful incorporation of the active species (Ru) and promoter (K).

The Ru-K/CaO catalysts with different Ru and K loadings were characterized by XRD at different stages: after thermal decomposition at 500 °C under an Ar atmosphere, after activation under H2 at 500 °C for 3 h, and after being exposed to ammonia under reaction conditions for 7 h. The XRD patterns are shown in Fig. 1. It is important to note that the CaO support tends to carbonate upon exposure to the atmosphere. Nevertheless, this phenomenon does not alter the properties of the catalysts since it does not occur during the catalytic testing under the activation or reaction conditions.

The XRD patterns recorded after thermal treatment under Ar, Fig. 1A, show CaO, K2O and RuO2 phases. As expected,

Table 1 BET surface area and ICP analysis for Ru-based catalysts

Sample	$\frac{S_{\text{BET}}}{(\text{m}^2 \text{ g}^{-1})}$	$\frac{\mathrm{Ru}^a}{\mathrm{(wt\%)}}$	$\frac{K^a}{(wt\%)}$
5%Ru10%K/MWCNT	758	4.7	9.7
5%Ru10%K/MgO	32	4.8	9.7
5%Ru10%K/NS	138	4.9	9.4
1%Ru10%K/CaO	6	0.9	9.6
2%Ru10%K/CaO	7	1.8	9.7
3%Ru10%K/CaO	5	2.8	9.6
5%Ru10%K/CaO	4	4.7	9.5
7%Ru10%K/CaO	8	6.6	9.4
3%Ru/CaO	10	2.9	_
3%Ru5%K/CaO	5	2.8	4.9
3%Ru15%K/CaO	5	2.7	13.8

<sup>&</sup>lt;sup>a</sup> Calculated by ICP-OES measurements.

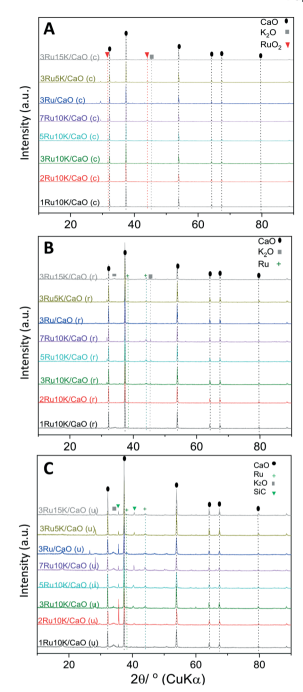


Fig. 1 XRD patterns of the Ru-K/CaO samples. A) Calcined samples, B) reduced samples and C) after catalytic testing

the most intense reflections are assigned to CaO (JCPDS 00-37-1497) with  $2\theta$  angles of 32.2°, 37.4°, 53.9°, 64.2°, and 67.4°. K<sub>2</sub>O with a small diffraction peak (JCPDS 00-47-1701) at a  $2\theta$  angle of 45.3° is formed under Ar thermal treatment after the incorporation of the promoter by impregnation. Additionally, diffraction peaks corresponding to RuO2 are also detected, due to the incorporation of Ru by impregnation (JCPDS 00-43-1027) with  $2\theta$  angles of 31.7 and 44.1°; the intensity of these reflections increases when increasing the Ru loading. After H2 treatment, Fig. 1B, apart

from the reflections assigned to CaO and K2O, full reduction of Ru is observed: see the disappearance of RuO2 associated with diffraction and the appearance of metallic Ru (JCPDS 00-06-0663) at  $2\theta$  angles of 38.4° and 44°. In the samples with different loadings of K, a well-defined Ru pattern was also observed (Fig. 1B). After the catalytic test, CaO, K2O and Ru are still visible in the XRD patterns (Fig. 1C).

The ruthenium dispersion and particle size were evaluated by CO pulse chemisorption in the samples after the activation under H2 treatment. Table 2 summarizes the obtained results for the metal dispersion, active metal surface area (AMSA), Ru particle size and number of surface active sites. It is interesting to observe that, in spite of the low surface area of the support, the metal dispersion and particle size remain constant for Ru loadings up to 3 wt%. Addition of Ru further results in bigger nanoparticles and therefore lower dispersions. Addition of K further contributes to enhancing dispersion, most probably by inhibiting Ru sintering as previously observed. 52-54

The comparison of the obtained results reveals 3%Ru10%K/CaO as the composition with the highest active surface area.

The local structures of the Ru-K/CaO catalysts were evaluated using high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) and energydispersive X-ray spectroscopy (EDX). Representative TEM images are shown in Fig. 2 for 3%Ru10%K/CaO and 3%Ru/ CaO catalysts after activation treatment under H<sub>2</sub> at 500 °C for 3 h. In the TEM images of the sample without K (3%Ru/CaO), Ru nanoparticles are dispersed on the CaO support with a quite broad particle size distribution in the range of 2.5 to 22.5 nm. The mean particle size, 9 nm, is in agreement with that calculated by CO chemisorption (Tables 2 and S4†). Elemental mapping shows that the Ru dispersion is not completely regular in these samples, with Ru localized preferably in certain regions. In the TEM images for the sample containing K (3%Ru10%K/CaO), significant differences are observed compared to the sample without potassium. Ru nanoparticles are better dispersed, with a narrower particle size distribution,

Table 2 Ruthenium dispersions, active metal surface areas and particle sizes for Ru-based catalysts

Sample	$\frac{\text{Dispersion}^a}{(\%)}$	$\frac{\text{AMSA}^b}{\left(\text{m}^2\ \text{g}_{\text{metal}}^{-1}\right)}$	$\frac{{d_{\rm p}}^c}{({\rm nm})}$	$\frac{N_{\rm AS} \times 10^{-3}}{\left(\text{mol g}^{-1}\right)}$
1%Ru10%K/CaO	16.2	87	6	1.6
2%Ru10%K/CaO	15.2	81	6	3.0
3%Ru10%K/CaO	15.5	83	6	4.6
5%Ru10%K/CaO	8.5	45	11	4.2
7%Ru10%K/CaO	5.2	28	17	3.6
3%Ru/CaO	9.8	52	9	2.9
3%Ru5%K/CaO	11.2	60	8	3.3
3%Ru15%K/CaO	14.8	79	6	4.4

<sup>&</sup>lt;sup>a</sup> Metal dispersion (%): weight of exposed metal atoms/weight of total metal atoms. <sup>b</sup> Active metal surface area (AMSA). <sup>c</sup> Active particle size; (spherical geometry).  $d_p$  (nm) = 6/(AMSA) $\cdot \rho_{Ru}$  (where  $\rho_{Ru}$ = 12.2 g cm<sup>-3</sup>). Number of surface active sites:  $N_{AS}$  (mol g<sup>-1</sup>) = Ru loading (wt)  $\times$  dispersion/Ru mass (g mol<sup>-1</sup>).

from 2 to 15 nm, and a smaller mean particle size, 7 nm, that matches with the particle size calculated by CO chemisorption (6 nm). Mapping shows that K, although is dispersed throughout the catalyst surface, it is preferably located on top of the Ru nanoparticles.

#### Catalytic activity

In a preliminary stage, the catalytic performance of potassium-promoted Ru catalysts on different supports was evaluated. Table 3 shows the ammonia conversion levels and hydrogen production rates at 400 °C for the Ru-K catalysts on different supports. The Ru-K catalysts supported on CaO and NS were found to be more efficient catalysts compared to those on other supports reported for this process, such as MWCNTs or MgO, with Ru-K/CaO showing the highest ammonia conversion (54%) and hydrogen productivity (574.3 mol H<sub>2</sub> per mol Ru h<sup>-1</sup>) at 400 °C. It is worth highlighting that these results are not correlated with the measured surface areas of the different catalysts. Considering that both Ru loadings and Ru: K ratios are similar for all the catalysts, this suggests that the chemical nature of the support has an important influence on the catalytic performance. For instance, the use of basic supports resulted in better catalytic performance, in line with this fact, CaO is a more basic support with a larger number and strength of basic sites. 55,56

In view of these results, the optimal metal and promoter loadings were subsequently investigated using CaO as the support. Fig. 3 shows the ammonia conversion versus reaction temperature, while the insets compare the activity at 400 °C for different Ru and K loadings. The conversions are not affected by heat or mass transport limitation (Fig. S1-S3†).

First, the samples with Ru loadings in the range of 1-7 wt% and constant K loading (10 wt%) were evaluated. The positive effect of Ru is clearly observed up to 3 wt% loading. After that, the activity decreases significantly. Secondly, the samples with the optimum Ru loading (3 wt%) and K loadings in the range of 0-15 wt% were also analyzed. For K promotion, also an optimum loading is observed, in this case 10 wt%. For CaOsupported samples, the catalytic performance is very well in line with the obtained characterization results, especially in terms of Ru dispersion, active metal surface area and TEM characterization, according to the beneficial effect of K preventing Ru agglomeration under activation and reaction conditions. 52-54 The best catalytic results are obtained for Ru nanoparticles of 6-7 nm. Ammonia decomposition on Ru has been shown to be a highly structure-sensitive reaction, with terrace sites being the main active sites<sup>57</sup> and with particles below 6 nm being hardly active. Our experimental results further confirm this earlier observation.

Once the catalyst composition was optimized, we studied the high-pressure (10-40 bar) decomposition performance of the 3%Ru10%K/CaO sample. Fig. 4A shows the impact of operating pressure on the catalytic performance. The ammonia conversion is greatly decreased when the pressure increases from 1 to 40 bar. Meanwhile a high conversion, around 80%, is

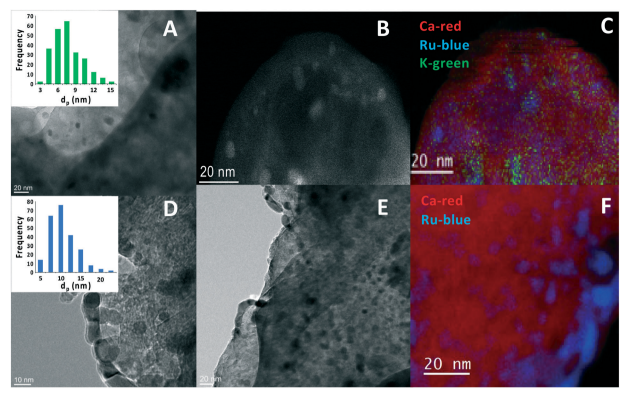


Fig. 2 Representative images for 3%Ru10%K/CaO (A–C) and 3%Ru/CaO (D–F) after activation under hydrogen at 525 °C for 3 h, high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) and energy-dispersive X-ray spectroscopy (EDX) images. Elemental mapping images for Ru (blue), calcium (red), and potassium (green).

still observed at 550 °C and 40 bar. This is further analysed in Fig. 4B. Interestingly, although a drastic decrease is observed in the conversion level, the hydrogen productivity increases with pressure (Fig. 4C), already anticipating a positive reaction order for NH<sub>3</sub> (*vide infra*) (Fig. 4C and D).

Finally, turnover frequencies (TOFs) were calculated (Fig. 4D and 5) considering the number of active surface sites evaluated by CO chemisorption studies (see above in section 3.1, Table 2). Rates far from equilibrium were considered to compare the TOF calculated for both samples. The promotion effect of potassium on the catalytic activity of the Ru catalysts is clearly observed in Fig. 5.

#### Kinetic analysis

Kinetic studies at atmospheric pressure were performed for the optimized 3%Ru10%K/CaO catalyst, as well as for the

Table 3 Ammonia conversion and hydrogen productivity of Ru-based catalysts on different supports at 400 °C (P = 1 atm; NH<sub>3</sub> flow rate 30 ml min<sup>-1</sup>; WHSV = 9000 mL h<sup>-1</sup> g<sup>-1</sup>)

Catalyst	$X_{\mathrm{NH_{3}}}\left(\% ight)$	$H_2$ production (mol $H_2$ per mol Ru $h^{-1}$ )
5%Ru10%K/MWCNT	30.8	322.7
5%Ru10%K/MgO	39.4	407.8
5%Ru10%K/NS	46.6	474.4
5%Ru10%K/CaO	53.7	574.3

unpromoted catalyst in order to elucidate the effect of potassium from a kinetic point of view. Reaction orders for NH<sub>3</sub> and H<sub>2</sub> were calculated from differential experiments varying the NH<sub>3</sub> and H<sub>2</sub> partial pressures and the temperature (details of the kinetic study in the ESI† and Fig. S4–S6). The results are summarized in Table 4. The apparent activation energy was calculated from the Arrhenius plots for both catalysts. Incorporation of K results in a decrease in the activation energy from 96 to 75 kJ mol<sup>-1</sup>. Promotion with alkali metals is well known to decrease the hydrogenation ability of Ru *via* electronic promotion.<sup>27</sup>

Inhibition of the ammonia decomposition rate by the hydrogen produced is observed. The negative orders observed for hydrogen for both catalysts can be explained, considering that chemisorbed hydrogen blocks surface sites that are necessary for ammonia decomposition, or reacts chemically and hydrogenates  $NH_x$  intermediates generated during the ammonia decomposition. As shown in Table 4, potassium promotion results in a decrease of both the reaction order on  $NH_3$  (from 0.8 to 0.5) and the negative reaction order on  $H_2$  (from -1.9 to -1.2), highlighting the positive electronic promotion of the alkali metal.

For Ru-based catalysts, it is generally agreed that the limiting reaction step is the dissociation of ammonia  $^{59,60}$  and the observed negative order on hydrogen is related to the rehydrogenation of partially dehydrogenated species (NH $_x$  intermediates, x=1, 2). The kinetic data clearly show the

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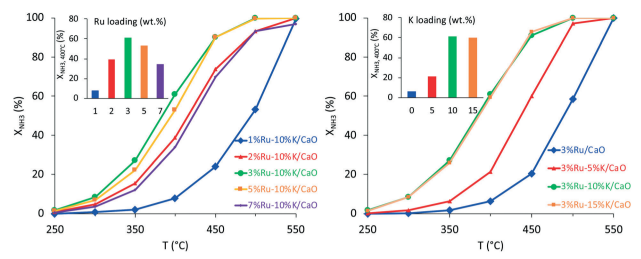


Fig. 3 Ammonia conversion versus temperature of Ru-K/CaO catalysts; effect of Ru and K loadings. P = 1 atm; T = 250-550 °C; NH<sub>3</sub> flow rate 30 N ml min<sup>-1</sup>;  $W_{cat} = 200$  mg; WHSV: 9000 mL g<sup>-1</sup> h<sup>-1</sup>.

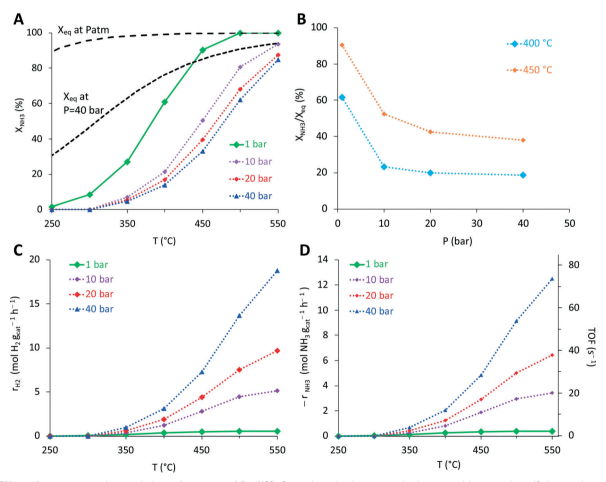


Fig. 4 Effect of pressure on the catalytic performance of Ru-K/CaO catalysts in the ammonia decomposition reaction. A) Ammonia conversion (%); B) conversion related to the equilibrium; C) hydrogen productivity (mol  $H_2$  per  $g_{cat} h^{-1}$ ); D) ammonia decomposition rate (mol  $NH_3$  per  $g_{cat} h^{-1}$ ) and TOF (s<sup>-1</sup>) versus temperature. The reaction rates of ammonia conversion and hydrogen production were calculated from the ammonia conversion values assuming a differential reactor and taking into account the operating pressure. P = 1-40 atm; T = 250-550 °C; NH<sub>3</sub> flow rate 30 N ml min<sup>-1</sup>; WHSV = 9000 mL h<sup>-1</sup> g<sup>-1</sup>.

ability of K to decrease the hydrogenation capability of Ru. In this way, inhibition of ammonia decomposition by hydrogen becomes less important and this is translated into an enhancement of the catalytic activity.

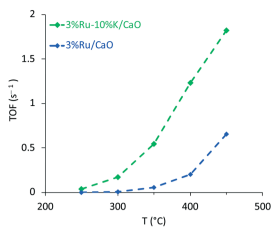


Fig. 5 Calculated turnover frequencies (TOFs) for K-promoted and unpromoted Ru/CaO catalysts. P =1 atm; T = 150-500 °C; NH<sub>3</sub> flow rate 30 ml min<sup>-1</sup>; WHSV = 9000 mL  $h^{-1}$   $g^{-1}$ 

Table 4 Kinetic parameters of the Ru-K/CaO catalysts for the ammonia decomposition reaction at 350  $^{\circ}$ C and  $P_{\text{atm}}$ 

Sample	$E_{\rm a}$ (kJ mol <sup>-1</sup> )	а	b
3%Ru/CaO 3%Ru10%K/CaO	96 75	0.8 0.5	-1.9 -1.2
3%Ku10%K/CaO	73	0.3	-1.2

The observed kinetics for the Ru-K/CaO catalysts at atmospheric pressure are well described using the power law given by eqn (1).

$$(-r_{NH_3}) = k \times P_{NH_3}{}^a P_{H_2}{}^b$$
 (1)

In order to develop a reliable kinetic model able to predict catalytic performance under relevant operating conditions for the industrial application of this process, where almost complete NH3 conversion is achieved, such as the usually high temperatures needed or the high pressures used in this work, the overall approach to thermodynamic equilibrium needs to be considered in order to contemplate the effect of the reverse reaction that suppresses the decomposition rate. Under these operating conditions, to attain the large conversions, the reaction rate is better described by eqn (2), including a term that accounts for the influence of the reverse reaction with the approach to the equilibrium.<sup>35,61,62</sup> This term becomes increasingly significant at temperatures above 400 °C or when ammonia conversion is close to the equilibrium conversion, and will definitively be important at the high pressures used here, above the atmospheric pressure.

$$(-r_{\text{NH}_3}) = k \times P_{\text{NH}_3}{}^a P_{\text{H}_2}{}^b \left( 1 - \frac{1}{K_{\text{eq}}} \left( \frac{P_{\text{N}_2} P_{\text{H}_2}{}^3}{P_{\text{NH}_3}{}^2} \right) \right)$$
(2)

The kinetic model (eqn (2)) considers that the rate is inhibited by the hydrogen product and also by the reverse reaction, non-negligible at temperatures below 500 °C.61

Considering the inhibition of the decomposition rate by the hydrogen produced, and taking into account that the gas composition is very different along the reactor with the increasing conversion level, an integral reactor must be considered for the kinetic analysis in this case, to account for the increasing hydrogen concentrations at the different conversion levels along the catalytic bed. 61 The integration of egn (3) allows the calculation of the ammonia conversion in a plug-flow reactor, where the ammonia conversion rate  $(-r_{\rm NH})$  is given by the expression in eqn (2), and can be compared with the experimental values. See details of the kinetic study in an integral reactor in the ESI.†

$$W/F = \int_0^X \frac{dX_{\rm NH_3}}{(-r_{\rm NH_3})} \tag{3}$$

In order to check the validity of the kinetic model based on the power-law to predict the catalytic performance at elevated pressures, the ammonia conversions were calculated under these operating conditions, by assuming an integral reactor and solving eqn (3) with the reaction rate given by eqn (2). In Fig. 6 the experimental conversions are compared with those predicted by the kinetic model given by eqn (2), showing an excellent matching and corroborating the goodness of the kinetic model (further details in the ESI,† with the fittings in Fig. S7 and S8). This excellent correspondence between the experimental and calculated values indicates that the kinetic model (eqn (2)) is not only very robust to predict low and high conversion levels, under a wide range of reaction conditions of temperature and space velocity, but also very reliable when predicting the catalytic performance at high pressure. This study demonstrates the validity of the selected kinetic model to predict the performance outside the data used for the fitting and suggests that experiments at higher pressures are probably not per se needed to assess the highpressure performance of ammonia decomposition catalysts.

Furthermore, other kinetic models taking into account different mechanistic considerations and assuming different rate determining steps (given by eqn (S5)-(S8) in the ESI†)

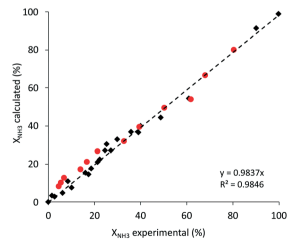


Fig. 6 Experimental vs. modeled ammonia conversion using the proposed power-law model (eqn (2)) for ammonia decomposition over the 3%Ru10%K/CaO catalyst. Red: ammonia conversion at high pressure. Black: conversion at atmospheric pressure.

were explored here to fit the experimental data, as well as to predict the catalytic performance at high pressure. Fittings obtained with all the kinetic models selected are shown in Fig. S7 and S8 in the ESI.† Considering the parity plots that compare the experimental and calculated values of ammonia conversion, it is difficult to draw conclusions related to the limiting step, since almost identical fittings are obtained with the mechanistic models that consider different rate determining steps, such as those assuming the ammonia dissociation (eqn (S7)†) or nitrogen desorption as the rate limiting step (eqn (S5)†). This is the main reason why, in spite of the research over the past decades in this reaction, there is no agreement about the limiting step or the kinetically relevant surface species. In fact, the results in Fig. S7 and S8† clearly corroborate that all the species present at the catalyst surface must be considered as kinetically relevant. The fitting of the experimental data with the models that consider a few adsorbed species as the most abundant ones (such as kinetic eqn (S6) and (S8)†) provides worse matching. This aspect becomes more significant when the process is conducted at high pressure (Fig. S8†).

Interestingly, from Fig. 6 and S7 and S8 in the ESI,† with the parity plots for different kinetic models, it is concluded that the power law given by eqn (2) is as good as the mechanistic models that take into account different rate limiting steps, while offering, by far, the simplest equation for a kinetic model. Furthermore, this model is the most robust to predict the catalytic performance for ammonia decomposition at high pressure (Fig. S8†).

# Conclusions

In summary, our work further demonstrates that Ru-based catalysts are very efficient for the decomposition of ammonia. Considering the relatively low Ru loadings desired and the strong structure sensitivity of this reaction, with an optimal Ru particle size in the order of 7 nm, the chemical nature of the support, especially its basicity, plays a much bigger role than other physical properties such as surface area. Considering these facts, among the studied supports, CaO has been identified as a very promising one.

The reaction proceeds with inhibition of the decomposition reaction rate by the hydrogen produced, suggesting that competitive adsorption of H2 and/or re-hydrogenation of NHx intermediates occur, and pointing to the ammonia dissociation as the limiting step of the reaction rate. By promoting Ru with K, the competitive adsorption of H2 is reduced and the hydrogenation capability of Ru is decreased, as shown by the strong decrease in the H<sub>2</sub> negative reaction order. This effect translates into higher intrinsic catalytic activity.

Last but not least, our modelling results demonstrate that kinetic analysis in the low-pressure regime using the power law-derived equation can be used to accurately predict catalytic performance at higher decomposition pressures, which, from an application perspective, are much more attractive but may be difficult to measure in many laboratories.

## Conflicts of interest

There are no conflicts to declare.

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## References

- 1 A. J. Chapman, B. C. McLellan and T. Tezuka, Appl. Energy, 2018, 219, 187-198.
- 2 V. Ş. Ediger, Energy Procedia, 2019, 156, 2-6.
- 3 R. York and S. E. Bell, Energy Res. Soc. Sci., 2019, 51, 40-43.
- 4 A. Singh, P. Baredar and B. Gupta, Energy Convers. Manage., 2017, 145, 398-414.
- 5 E. Abokyi, P. Appiah-Konadu, F. Abokyi and E. F. Oteng-Abayie, Energy Rep., 2019, 5, 1339-1353.
- 6 H. A. D. Hdom, Renewable Energy, 2019, 139, 186-197.
- N. M. Jalal, A. R. Jabur, M. S. Hamza and S. Allami, Energy Procedia, 2019, 157, 1494-1505.
- 8 A. Chica and S. Sayas, Catal. Today, 2009, 146(1), 37-43.
- 9 S. Sayas and A. Chica, Int. J. Hydrogen Energy, 2014, 39(10), 5234-5241.
- 10 M. Menor, S. Sayas and A. Chica, Fuel, 2017, 193, 351-358.
- 11 D. Guban, I. K. Muritala, M. Roeb and C. Sattler, Int. J. Hydrogen Energy, 2019, DOI: 10.1016/j.ijhydene.2019.08.145.
- 12 D. Xu, W. Li, X. Ren, W. Shen and L. Dong, Int. J. Hydrogen Energy, 2019, DOI: 10.1016/j.ijhydene.2019.09.030.
- 13 C. Weidenthaler, J. Energy Chem., 2020, 42, 133-143.
- 14 M. Lototskyy, I. Tolj, Y. Klochko, M. W. Davids, D. Swanepoel and V. Linkov, Int. J. Hydrogen Energy, 2019, 45(14), 7958-7967.
- 15 P. T. Aakko-Saksa, C. Cook, J. Kiviaho and T. Repo, J. Power Sources, 2018, 396, 803-823.
- 16 H. Jorschick, M. Vogl, P. Preuster, A. Bösmann and P. Wasserscheid, Int. J. Hydrogen Energy, 2019, 44(59), 31172-31182.
- 17 S. Mukherjee, S. V. Devaguptapu, A. Sviripa, C. R. F. Lund and G. Wu, Appl. Catal., B, 2018, 226, 162-181.
- 18 N. Itoh, A. Oshima, E. Suga and T. Sato, Catal. Today, 2014, 236, 70-76.
- 19 A. Kantürk Figen, M. B. Pişkin, B. Coşkuner and V. İmamoğlu, Int. J. Hydrogen Energy, 2013, 38(36), 16215-16228.
- 20 Y. Kojima, Int. J. Hydrogen Energy, 2019, 44(33), 18179–18192.
- 21 J. W. Makepeace, T. He, C. Weidenthaler, T. R. Jensen, F. Chang, T. Vegge, P. Ngene, Y. Kojima, P. E. de Jongh, P. Chen and W. I. F. David, Int. J. Hydrogen Energy, 2019, 44(15), 7746-7767.
- 22 R. Metkemeijer and P. Achard, Int. J. Hydrogen Energy, 1994, **19**(6), 535–542.
- C. H. Christensen, T. Johannessen, R. Z. Sørensen and J. K. Nørskov, Catal. Today, 2006, 111(1), 140-144.
- Y. W. Cheng, M. R. Khan, K. H. Ng, S. Wongsakulphasatch and C. K. Cheng, Renewable Energy, 2019, 138, 1114-1126.
- L. P. Didenko, L. A. Sementsova, P. E. Chizhov and T. V. Dorofeeva, Int. J. Hydrogen Energy, 2019, 44(48), 26396-26404.

- 26 F. R. García-García, Y. H. Ma, I. Rodríguez-Ramos and A. Guerrero-Ruiz, Catal. Commun., 2008, 9(3), 482-486.
- A. K. Hill and L. Torrente-Murciano, Int. J. Hydrogen Energy, 2014, 39(15), 7646-7654.
- 28 J. C. Ganley, F. S. Thomas, E. G. Seebauer and R. I. Masel, Catal. Lett., 2004, 96(3-4), 117-122.
- 29 A. Boisen, S. Dahl, J. K. Nørskov and C. H. Christensen, J. Catal., 2005, 230(2), 309-312.
- 30 C. J. H. Jacobsen, S. Dahl, B. S. Clausen, S. Bahn, A. Logadottir and J. K. Nørskov, J. Am. Chem. Soc., 2001, 123(34), 8404-8405.
- 31 S. Bajus, F. Agel, M. Kusche, N. Ní Bhriain and P. Wasserscheid, Appl. Catal., A, 2016, 510, 189-195.
- 32 A. Di Carlo, L. Vecchione and Z. Del Prete, Int. J. Hydrogen Energy, 2014, 39(2), 808-814.
- 33 C. Huang, Y. Yu, J. Yang, Y. Yan, D. Wang, F. Hu, X. Wang, R. Zhang and G. Feng, Appl. Surf. Sci., 2019, 476, 928–936.
- 34 D.-C. Huang, C.-H. Jiang, F.-J. Liu, Y.-C. Cheng, Y.-C. Chen and K.-L. Hsueh, Int. J. Hydrogen Energy, 2013, 38(8), 3233-3240.
- 35 K. Lamb, S. S. Hla and M. Dolan, Int. J. Hydrogen Energy, 2019, 44(7), 3726-3736.
- 36 L. Li, Y. Wang, Z. P. Xu and Z. Zhu, Appl. Catal., A, 2013, 467, 246-252.
- 37 L. Li, Z. H. Zhu, Z. F. Yan, G. Q. Lu and L. Rintoul, Appl. Catal., A, 2007, 320, 166-172.
- 38 X.-C. Hu, W.-W. Wang, Z. Jin, X. Wang, R. Si and C.-J. Jia, I. Energy Chem., 2019, 38, 41-49.
- 39 X. Ju, L. Liu, P. Yu, J. Guo, X. Zhang, T. He, G. Wu and P. Chen, Appl. Catal., B, 2017, 211, 167-175.
- J. Chen, Z. H. Zhu, S. Wang, Q. Ma, V. Rudolph and G. Q. Lu, Chem. Eng. J., 2010, 156(2), 404-410.
- 41 S.-F. Yin, B.-Q. Xu, C.-F. Ng and C.-T. Au, Appl. Catal., B, 2004, 48(4), 237-241.
- 42 S. Chiuta, R. C. Everson, H. W. J. P. Neomagus and D. G. Int. J. Hydrogen Energy, Bessarabov, 3774-3785.
- 43 I. Lucentini, A. Casanovas and J. Llorca, Int. J. Hydrogen Energy, 2019, 44(25), 12693-12707.
- 44 S. J. Wang, S. F. Yin, L. Li, B. Q. Xu, C. F. Ng and C. T. Au, Appl. Catal., B, 2004, 52(4), 287-299.
- 45 P. Chen, X. Ju, L. Liu, X. Zhang, J. Feng and T. He, ChemCatChem, 2019, 11, 4161-4170.

- 46 J. Bellosta von Colbe, J.-R. Ares, J. Barale, M. Baricco, C. Buckley, G. Capurso, N. Gallandat, D. M. Grant, M. N. Guzik, I. Jacob, E. H. Jensen, T. Jensen, J. Jepsen, T. Klassen, M. V. Lototskyy, K. Manickam, A. Montone, J. Puszkiel, S. Sartori, D. A. Sheppard, A. Stuart, G. Walker, C. J. Webb, H. Yang, V. Yartys, A. Züttel and M. Dornheim, Int. J. Hydrogen Energy, 2019, 44(15), 7780-7808.
- 47 C. Deng, M. Zhu, Y. Zhou and X. Feng, Energy, 2018, 159, 203-215.
- 48 G. Sdanghi, G. Maranzana, A. Celzard and V. Fierro, Renewable Sustainable Energy Rev., 2019, 102, 150-170.
- P. Di Profio, S. Arca, F. Rossi and M. Filipponi, Int. J. Hydrogen Energy, 2009, 34, 9173-9180.
- 50 M. Al-Zareer, I. Dincer and M. A. Rosen, Int. J. Hydrogen Energy, 2019, 44(33), 18214-18224.
- 51 B. Wang, H. Kong, H. Wang, Y. Wang and X. Hu, Int. J. Hydrogen Energy, 2019, 44(49), 26874-26887.
- K. Aika, K. Shimazaki, Y. Hattori, A. Ohya, S. Ohshima, K. Shirota and A. Ozaki, *J. Catal.*, 1985, 92(2), 296-304.
- 53 G. Ertl, Catal. Rev.: Sci. Eng., 1980, 21(2), 201-223.
- 54 F. F. Gadallah, R. M. Elofson, P. Mohammed and T. Painter, Stud. Surf. Sci. Catal., ed. G. Poncelet, P. Grange and P. A. Jacobs, Elsevier, 1983, vol. 16, pp. 409-420.
- 55 H. Hattori and Y. Ono, Catalysts and catalysis for acid-base reactions, in Metal Oxides in Heterogeneous Catalysis, ed. J. C. Védrine, Elsevier, 2018, ch. 4, pp. 133-209.
- 56 Y. Ono and H. Hattori, Preparation and Catalytic Properties of Solid Base Catalysts - I. Metal Oxides, in Solid Base Catalysis, Springer, Berlin, Heidelberg, 2011, pp. 69-156.
- 57 A. M. Karim, V. Prasad, G. Mpourmpakis, W. W. Lonergan, A. I. Frenkel, J. G. Chen and D. G. Vlachos, J. Am. Chem. Soc., 2009, **131**(34), 12230–12239.
- 58 W. Tsai, J. J. Vajo and W. H. Weinberg, J. Am. Chem. Soc., 1985, 89, 4926.
- 59 T. E. Bell and L. Torrente-Murciano, Top. Catal., 2016, 59(15-16), 1438–1457.
- 60 F. Schüth, R. Palkovits, R. Schlögl and D. S. Su, Energy Environ. Sci., 2012, 5, 6278-6289.
- 61 S. Armenise, E. García-Bordejé, J. L. Valverde, E. Romeo and A. Monzón, Phys. Chem. Chem. Phys., 2013, 15(29), 12104-12117.
- 62 A. Di Carlo, A. Dell'Era and Z. Del Prete, Int. J. Hydrogen Energy, 2011, 36, 11815-11824.