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Ni-Ru supported on CeO₂ obtained by mechanochemical milling for catalytic hydrogen production from ammonia

SYNTHESIS NH₃ N_2 NH₃ conversion **METHOD** at 450°C 72.1% Ce⁴⁺ Ce³⁺ nickel nitrate + CeO₂ - 1st step ruthenium acetate + CeO₂ - 2nd step 56.7% Ce⁴⁺ Ce³⁺ nickel nitrate + ruthenium acetate + CeO₂ - 1st step Mechanochemical milling Incipient wetness impregnation 40.0% Ce⁴⁺ Ce³⁺ nickel nitrate + ruthenium acetate + CeO₂ - 450°C Ni 68.2% Ce⁴⁺ Ce³⁺ nickel acetate + ruthenium chloride + CeO₂ - 450°C

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Highlights

Mechanochemical synthesis is an efficient method for catalyst preparation

NiRu/CeO₂ has proven to be a highly active catalyst for NH_3 decomposition

The high activity is attributed to a high number of vacancies in CeO₂ structure

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Ni-Ru supported on CeO₂ obtained by mechanochemical milling for catalytic hydrogen production from ammonia

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SUMMARY

Developing active and stable catalysts for carbon-free hydrogen production is crucial to mitigate the effects of climate change. Ammonia is a promising carbon-free hydrogen source, as it has a high hydrogen content and is liquid at low pressure, which allows its easy storage and transportation. We have recently developed a nickel-based catalyst with a small content of ruthenium supported on cerium oxide, which exhibits high activity and stability in ammonia decomposition. Here, we investigate mechanochemical milling for its synthesis, a faster and less energy-consuming technique than conventional ones. Results indicate that mechanochemical synthesis increases catalytic activity compared to the conventional incipient wetness impregnation method. The interaction between the metal precursors and the support is key in fine-tuning catalytic activity, which increases linearly with oxygen vacancies in the support. Moreover, the mechanochemical method modifies the oxidation state of Ni and Ru species, with a variation depending on the precursors.

INTRODUCTION

To limit climate change and environmental degradation, the European Commission approved the European Green Deal¹ in 2020. Among the suggested actions, the decarbonization of power generation is of prime importance, as it accounts for around three-quarters of global greenhouse gas emissions. In this context, hydrogen plays a key role. Its role has been described in detail in the Hydrogen Strategy,² which aims to develop and scale up renewable and low-carbon hydrogen supply and demand, turning hydrogen into a viable solution toward a climate-neutral economy. Among the proposed actions, the on-site catalytic production of hydrogen from ammonia is considered an interesting approach to feed proton exchange membrane fuel cells, which require a high hydrogen purity.³ Among numerous advantages, ammonia, a carbon-free molecule, has a high hydrogen content (17.8 wt.% and 108 kg H₂ m⁻³ at 20°C and 8.6 bar⁴), and it can be stored and transported easily as it can liquefy at low pressure. It is decomposed catalytically into hydrogen and nitrogen at moderate temperature, around 400°C–600°C at 1 bar (2NH_{3(g)} \leftrightarrows N_{2(g)} + 3H_{2(g)}; Δ H^o = 92 kJ mol⁻¹, i.e., the ammonia decomposition).

Currently, research on the catalysts for ammonia decomposition, aimed at hydrogen production, still focuses on ruthenium, with nickel and cobalt emerging as primary alternatives due to their cost-effectiveness. Alternative preparation techniques, including coprecipitation⁵ or sol-gel^{6,7} methods, are being investigated, together with the effect of different promoters,^{8–10} supports,^{11,12} or active phase precursors¹³ on the catalyst activity. Moreover, the utilization of oxide combinations, such as Al-La-Ce¹⁴ or CeO₂-promoted MgAl₂O₄,¹⁵ as catalyst supports, has allowed the enhancement of the stability of ruthenium, and this effect has been ascribed to the high presence of oxygen vacancies in the support structure. Innovative combinations based on aluminate,¹⁶ zeolite,¹⁷ or perovskite¹⁸ structures and bimetallic catalysts such as Ni-Mo¹⁹ and Fe-Cr²⁰ have also presented good results. We have recently developed a catalyst based on nickel (5 wt.%) with a small content of ruthenium (1 wt.%) supported on ceria nanoparticles for the ammonia decomposition prepared by classical incipient wetness impregnation (IWI), which showed an excellent catalystic performance in long-term stability tests with respect to monometallic Ru/CeO₂ and Ni/CeO₂ and in multicycle tests.²¹ The bimetallic catalyst prepared did not allow an increase in the conversion obtained with Ru/CeO₂, but the latter deactivates rapidly (as demonstrated in a previous work²²). The characterization has revealed the existence of an intimate contact between Ni, Ru, and CeO₂, which is considered the reason for the excellent catalytic activity and stability observed.²³

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Figure 1. HRTEM images of bimetallic samples Ni(N)-Ru(A)-IWI and Ni(N)-Ru(A)-MM (A and B) Ni(N)-Ru(A)-IWI and (C and D) Ni(N)-Ru(A)-MM, with the histograms of particle size.

The method employed to synthesize the catalysts, IWI, is widely used, although nowadays alternative technologies, such as mechanochemical milling (MM), are being investigated to precisely control the active sites formation according to specific properties of amorphous/crystalline interfaces. MM synthesis methods are particularly attractive due to their low energy and time requirements, versatility, and scalability. The possibility of preparing materials with an amorphous layer on the support using MM methods has been already reported in the literature.²⁴ It has been observed that the formation of the amorphous phase increases as the milling time increases. This phenomenon has been attributed to the fact that the energy stored as structural defects^{25,26} on the surface of the particles becomes greater than in the bulk; therefore, to minimize the energy, an amorphous phase or thin layer can form on the surface or at the edges of the particles, while maintaining the elementary crystalline matrix.²⁵ In previous works,^{27–30} the described amorphous layer was obtained using the MM method with different catalysts supported on ceria, and it has been demonstrated that the formation of this amorphous layer is closely linked to an increase in the activity of the catalyst. However, it is not always possible to obtain this amorphous/crystalline interface, as it strongly depends on the relative hardness of the precursors. Nonetheless, the amorphization of a part of the supported metal, using Ni or Ru, has been reported in the literature.^{31,32}

In this work, we study the MM method for the synthesis of the bimetallic catalyst composed of 5 wt.% of Ni and 1 wt.% of Ru supported on ceria. The aim is to reduce the time and energy required for the catalyst preparation while increasing the repeatability of the synthesis procedure. In particular, we have varied the precursors of the two metals among different forms (nitrate: N, acetate: A, chloride: C, and nanoparticles: np). Additionally, we have studied the effect on the catalyst activity of the introduction of the precursors into the MM vial in one or two steps (denoted by the absence or the presence of a line between the two precursors, starting with the precursor used in the first phase, respectively), to identify the synthesis procedure that maximizes the activity in ammonia decomposition for hydrogen production. Furthermore, through the characterization we establish correlations between catalyst characteristics and the synthesis method, comparing IWI and MM methods, while identifying the relationship between the precursor used and the resulting catalyst's properties. Specifically, we focus on the presence of defects within the ceria structure, on the oxidation states of Ni and Ru, and on the dispersion of these on the surface of the catalyst.

RESULTS

Morphological properties of the catalysts

HRTEM was employed to study the morphology, particle size, and crystal structure of catalysts prepared using the same precursors with the two different methods (Ni(N)Ru(A)-IWI and Ni(N)-Ru(A)-MM; Figure 1). The ceria particles that compose the analyzed catalysts exhibited an





Figure 2. Raman spectra of the samples as prepared

(A) Those prepared with Ni(A) and Ni(np) as precursors, and (B) those prepared with Ni(N) as precursor and monometallic samples. The inserts show the F_{2g} peak center for all the catalysts and the bare support.

average size of around 17 \pm 4 nm for the catalyst prepared by IWI and 13 \pm 3 nm for the one prepared by MM method, as determined from the histograms (Figure 1) obtained by measuring the particles in Figures 1A and 1C. In the high-resolution images (Figures 1B and 1D), differences in the d-spacing identified in the FFT images (insets) can be observed. In the case of the IWI catalyst, besides the identified planes attributed to cerium oxide [mainly (111), (002), (022), and (113) identified through d-spacing of 3.1, 2.7, 1.9, and 1.6 Å, respectively], d-spacings corresponding to the oxidized forms of Ni and Ru were identified as well, in particular, the planes (110) with a d-spacing of 3.2 Å; (210), 2.0 Å; and (211), 1.7 Å of RuO₂³³ and the planes (111) (d-spacing of 2.4 Å) and (200), 2.1 Å, of NiO.³⁴ It was not possible to estimate the average dimensions of the NiO and RuO₂ particles because their crystallites overlap with those of the ceria. This indicates that the particles of the two oxides and the support are intimately interconnected within the catalyst structure. For the MM-prepared catalyst, crystalline (210) planes of Ni(NO₃)₂ with a d-spacing of 3.3 Å³⁵ were identified. Similarly, these planes overlapped with ceria crystallites, hindering the unequivocal identification of the particles. No interplanar distances related to ruthenium acetate have been identified; however, d-spacings attributable to RuO₂ were identified, specifically at 3.2, 2.6, 2.0, and 1.7 Å corresponding to the (110), (101), (210), and (211) planes.

Interaction between the active phase and the support

The Raman spectra of the prepared catalysts are presented in Figure 2. The bands at 265, 463, 591, and 1177 cm⁻¹ observed in the bare ceria spectrum are assigned to the second-order transverse acoustic (2TA) mode, the vibrational mode of the fluorite phase (F_{2a}), the defectinduced mode (D), and the second-order longitudinal optical (2LO) mode, respectively.³⁶ All spectra were normalized to the height of the $F_{2\alpha}$ peak. These bands were observed in all spectra, with certain variations: a broadening coupled with a red shift of the $F_{2\alpha}$ peak, and an increase in the area of the bands related to defects in the ceria structure were observed for all the supported catalysts. These variations are linked to decreasing mean crystallite size, increasing lattice parameters, and higher disorder in the ceria structure.³⁷ In particular, a more pronounced red shift and broadening of the F_{2g} peak, and a larger area of the D and 2TA mode bands, indicate a higher level of defects in the ceria structure. The D band associated with defects in the ceria structure can be divided into three contributions: D1, due to intrinsic Frenkel defects of ceria; D2, due to extrinsic defects originating from metal-support interaction; and D3, assigned to oxygen vacancies coupled to the presence of Ce^{3+} or other aliovalent cations.³⁸ The ratio between the F_{2q} band area and the sum of the areas of three contributions D1, D2, and D3 (D) is indicative of the interaction between the support and the used precursors. The values of the F2a peak center, its FWHM, and the ratio between the intensity of the D band and that of F_{2q} (I_D/I_F) for all the prepared catalysts and the support are reported in Table 1. In addition to the ceria oxide bands, several additional bands were observed in the spectra of the prepared catalysts. Catalysts synthesized using nickel acetate displayed peaks related to the presence of carbon and/or oxygen bonds, in particular, a peak at 952 cm⁻¹, attributed to v_4 (C-C) and one at 680 cm⁻¹, assigned to v_5 (COO). In addition to these peaks, the spectra of the catalysts prepared using Ni acetate presented peaks at 364 and 270 cm⁻¹, corresponding to the stretching vibration of v(Ni-O) of the aquo complex of Ni(II) and of the acetatenickel complex,³⁹ respectively. In the case of the Ni(A)Ru(A) catalyst, the peak at 952 cm⁻¹ overlapped with a second peak (936 cm⁻¹), attributed to the presence of Ru acetate, as confirmed by analyzing the Ru(A)-MM catalyst spectrum (Figure 2B) where this peak appears at 930 cm⁻¹ (attributed to v(C-C) of Ru acetate). In the Ru(A)-MM spectrum, a band around 334–338 cm⁻¹ is visible, ascribed to hydrated RuO₂,⁴⁰ and this band is stronger in the spectra of the catalysts prepared using Ru(A) and Ni(N) as precursors. Furthermore, in the case of the monometallic catalyst Ru(A)-MM, a peak at 828 cm⁻¹ was identified, which is not visible in the spectrum of Ni(A)Ru(A)-MM. Bimetallic spectra containing Ru(A) also displayed a band around 900 cm⁻¹. These bands cannot be univocally attributed to a specific vibrational mode but likely originate from ruthenium acetate's presence or its interaction with the Ni precursor or ceria support. The analysis of the spectra of the catalysts prepared using Ru chloride as a precursor, apart from the peaks already described, revealed the presence of bands with peaks around 323 cm⁻¹ (attributed to a quasi-elastic scattering component in A_{1g} of $RuCl_3^{41,42}$) and 835–843 cm⁻¹ (not unequivocally

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Sample	Raman			XANES ^a	XPS	
	$F_{2g} \text{ cm}^{-1}$	FWHM F _{2g}	I _D /I _F	% Ce ³⁺ /Ce	% Ce ³⁺ /Ce	% O _S /O
CeO2	463.4	14.9	0.04	16.4(3)	-	-
Ni(A)Ru(A)-MM	462.3	16.9	0.04	16.9(5)	-	-
Ni(A)-Ru(C)-MM	460.0	19.2	0.07	17.3(2)	-	_
Ni(N)-MM	460.7	16.0	0.04	20.6(6)	14.0	37.7
Ni(N)-IWI	-	-	-	21.6(6)	33.4	68.0
Ni(np)-Ru(np)-MM	453.5	35.6	0.09	23.0(2)	-	-
RuA-IWI	-	-	-	23.7(1)	-	-
RuA-MM	461.3	21.4	0.06	24.1(1)	-	-
Ni(A)Ru(C)-MM	458.7	22.6	0.10	24.3(6)	-	-
Ni(N)Ru(A)-IWI	457.1	33.6	0.69	24.9(1)	19.8	76.2
Ni(np)Ru(np)-MM	457.1	38.0	0.46	28.3(8)	-	-
Ni(N)Ru(C)-MM	460.0	21.8	0.16	29.8(1)	-	-
Ni(N)Ru(A)-MM	460.0	26.2	0.44	29.8(2)	17.9	40.5
Ni(N)-Ru(A)-MM	462.3	21.3	0.45	31.3(6)	16.2	42.7
Ni(A)Ru(C)-IWI	452.2	51.2	1.04	32.1(1)	18.1	25.5

attributable to a specific vibrational mode, but likely resulting from the presence of Ru chloride). In the case of the catalysts prepared with Ni(NO₃)₂.6H₂O, a strong band between 1,010 and 1,080 cm⁻¹, corresponding to the nitrate group v₁(NO³⁻), was observed. This band is composed of two peaks: one relative to the coordinated NO³⁻ at around 1,020 cm⁻¹ and a second relative to the free NO³⁻ at around 1,050 cm⁻¹.⁴³ Monometallic Ni(N)-MM presented only the free NO³⁻ related band, whereas the other Ni-nitrate-containing catalysts exhibited both contributions in a coordinated/free proportion of approximately 1/3 for Ni(N)Ru(C)-MM and Ni(N)Ru(A)-MM and around 1/5 for Ni(N)-Ru(A)-MM. Three weaker bands between 1,260–1,350; 700–760; and 400–420 cm⁻¹, corresponding to the stretching modes of v_{as}(NO), v₄(NO³⁻) and v(Ni-N), respectively,^{44,45} were detected. The spectra of the catalysts prepared by MM using Ni(np) and Ru(np) as precursors presented a pattern very similar to those of the two catalysts prepared by IWI: the D band was much more intense than for the other catalysts [especially intense for IWI catalysts and then for Ni(np)Ru(np)-MM], and the only peak observable, apart from the bands related to cerium oxide, around 965–975 cm⁻¹, coupled with a peak around 680–700 cm⁻¹, was attributed to the formation of the Ru-O-Ce bond.^{46–48} The bands related to NiO and Ni-Ce-O are located at 520 and 635 cm⁻¹, respectively.⁴⁹ Therefore, in the samples examined they can be overlapped by the F_{2g} and D bands of ceria and consequently not distinguishable.

Composition of the catalysts and crystallite size

The X-ray diffraction (XRD) patterns of the prepared catalysts were collected at a wavelength of 0.05904 nm (21 keV) and are presented in Figure 3. All patterns present peaks attributed to cubic fluorite-type cerium oxide⁵⁰ with the four primary peaks identifiable at 20 of 10.8° (111), 12.5° (200), 17.8° (220), and 20.9° (311). The dimensions of ceria crystallites calculated using the Debye-Scherrer equation are 11 nm in bare ceria and 12 nm for all prepared catalysts. No noticeable shift of the ceria-related peaks of the prepared catalysts compared to the CeO₂ pattern was identified. In the case of the catalysts prepared by IWI (Ni(N)Ru(A)-IWI and Ni(A)Ru(C)-IWI), peaks at 14.0° (111), 16.2° (200), and 23.0° (220) attributed to an FCC NiO phase⁵¹ were observed, due to the calcination process at 450°C in the presence of air. Using the Debye-Scherrer equation the crystallite size of nickel oxide is estimated to be around 9 nm for both IWI catalysts. No ruthenium-related peaks were identified. The patterns of the catalysts prepared by MM method using Ni acetate as precursor presented several peaks related to Ni(A)⁵² (dotted lines in Figure 3). This was confirmed through the analysis under the same conditions of the unsupported precursors (Figure 4). Ni(A) first peaks were observed at 20 values of 4.9°, 5.7°, 7.1°, 7.7°, 8.0°, 8.4°, and 8.9°, corresponding to the peaks at 8.0°, 9.3°, 11.5°, 12.4°, 13.0°, 13.6°, and 13.8° of the unsupported Ni acetate, indicating that the peaks in the catalyst pattern have undergone a red shift. Using the Debye-Scherrer equation, the crystallite sizes of the highly crystalline unsupported material were estimated to be 59 nm, whereas in the case of the three prepared catalysts, these dimensions were estimated to be 50, 63, and 72 nm, in the order Ni(A)Ru(C)-MM < Ni(A)-Ru(C)-MM < Ni(A) Ru(A)-MM. In the XRD patterns of the catalysts prepared using the Ru chloride, Ru acetate, and Ni nitrate precursors, no peaks relative to Ru or Ni were identified. In the case of ruthenium, no considerations can be made as a content of 1 wt.% may be lower than the detection limit of the technique. In contrast, in the case of Ni, this may suggest that the particle size of Ni species is too small to be detected by XRD or that they are highly dispersed. Analyzing the unsupported precursors (Figure 4), Ru(C) main peaks were determined at 9.2°, 10.6°, 15.7°, and 21.4° 20 angles, 53 and the dimensions of the crystallites were estimated to be around 10 nm, whereas in the case of Ru(A) the XRD pattern indicates a low

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Figure 3. XRD patterns measured at 21 keV of the catalyst samples, as prepared

crystallinity level of the material. For catalysts prepared with Ni and Ru nanoparticles as precursors, the metallic FCC Ni (16.7°, 19.3°, 32.3°, and 33.8° 20 angles) and HCP Ru (15.9° 20 angle) phases were identified, along with the presence of Ni-oxidized phase in both analyzed catalysts. The size of the crystallites for the two highly crystalline precursors before being supported was estimated as 41 nm for metallic Ni [peaks 20 located at 27.2° (111), 31.5° (200), 45.1° (220), 53.4° (311), and 56.0° (222)⁵⁴] and 7 nm in the case of metallic Ru [main peak at 26.8° (101)⁵⁵]. Also, in this case, a strong red shift of the peaks can be observed. When supported metallic Ru particles and NiO did not allow the estimation of the dimensions of their crystallites.

Oxygen vacancies in the ceria structure

The amount of Ce^{3+} ions compared to the total amount of ceria has been indicated as directly proportional to the number of oxygen defects present in its structure.⁵⁶ Therefore, to assess and quantify the valence states present in the support of the different prepared catalysts, X-ray absorption near edge structure (XANES) spectra were collected and analyzed. The raw data were reduced using *Athena*. A curve-fitting analysis was performed within the range of -25 and 25 eV with respect to the edge jump using an *arctan* function to simulate the edge jump (with a height of 0.95 and a width of 1.5, around 5726 eV) centered at the maximum of the first derivative.⁵⁷ All spectra exhibited the typical doublet due to the interaction between the 4f orbitals of the Ce atoms and the 2p orbitals of oxygen, ^{58,59} composed of four peaks (Figure 5A). Gaussian functions were employed to fit the peaks, centered around the energy values: 5721 (D), 5726 (C), 5731 (B), and 5738 eV (A). Component A is assigned to the transition from the Ce 2p shell to the 5d shell with no electron in the Ce 4f shell, whereaS component B is assigned to the excitation from the 2p shell to the 5d shell, along with an electron being excited from the O 2p shell to the Ce 4f shell, thus leaving a hole in the valence band. Components A and B are associated with the Ce⁴⁺ ions.⁵⁹ Component C, assigned to Ce³⁺, results from the 5d to 4f emission.⁶⁰ Component D, in the pre-edge region, was assigned to the 2p_{3/2} to 4f quadrupole transition, which is a consequence of 5d ad-mixtures to the 4f states.⁶¹ The quantification of [Ce³⁺] was done by calculating the area ratio of Ce³⁺ (C) with respect to all Ce (A + B + C).⁵⁷ The results, reported in Table 1, indicate a variable presence of Ce³⁺. The CeO₂ support presented an initial concentration of 16%, and the amount of reduced ceria increased with the presence of Ni and Ru, up to a maximum in the case of the Ni(N)-Ru(A)-MM and Ni(A)Ru(C)-IWI catalysts around 31% and 32%, respective



Figure 4. XRD patterns measured at 21 keV of the unsupported precursors







Figure 5. Normalized XANES spectra at the Ce L₃-edge for selected catalysts

(A) illustrates an example of fitting (CeO₂ bare support), whereas (B) presents a comparison of spectral intensity between CeO₂, Ni(A)Ru(A)-MM, Ni(N)-Ru(A)-MM, and Ni(A)Ru(C)-IWI.

observable: in Ni(A)Ru(A)-MM catalyst the estimated amount of reduced ceria remained unchanged compared to the support (16.9%), possibly indicating a low or absent interaction between the precursors, whereas in the case of the Ni(N)-Ru(A)-MM and Ni(A)Ru(C)-IWI catalysts a decrease in the Ce⁴⁺ contribution can be observed, indicated by the lower intensity of the doublets. These results support the findings from Raman spectroscopy: the MM method guarantees an excellent contact between Ni, Ru, and the support, since their presence leads to an increase in the presence of Ce³⁺, associated with the presence of oxygen vacancies and defects in the ceria structure.

Oxidation state of the active phase

The XANES spectra collected at the K-edges of Ni and Ru were analyzed to identify their respective oxidation state. The obtained E₀ values for Ni (Figure 6A) and Ru (Figure 6B) are shown in Figure 6. These spectra revealed the presence of the oxidized forms in all catalysts prepared via MM. The percentage of NiO and RuO₂ obtained through linear combining fitting of the XANES spectra of the catalysts using those of the precursors and oxides is reported in Table 2. The catalysts prepared using nanoparticles of Ni and metallic Ru are mainly composed of the metallic form, with minor proportions of NiO [4.7 and 12.4% for Ni(np)-Ru(np)-MM and Ni(np)Ru(np)-MM, respectively] and RuO₂ [2.6 and 4.3% for Ni(np)-Ru(np)-MM and Ni(np)Ru(np)-MM, respectively]. The presence of RuO₂ is lower with respect to NiO, and in both EXAFS spectra the main contribution coincides with the spectrum of metallic form (Ni-Ni bond at 2.2 Å and Ru-Ru bond at 2.5 Å; Figures 7A and 8C). In contrast to catalysts employing metal nanoparticles, the analysis suggests a higher oxidation state of ruthenium compared to nickel in the other catalysts (Table 2). Generally, Ni acetate-based catalysts prepared via MM exhibited a lower fraction of the oxidized forms for both Ni and Ru. The IWI-prepared catalysts presented nearly complete oxidation of the precursors, particularly in the case of ruthenium, showing a 96%–98% oxide state according to the linear combination fitting of the XANES spectrum. Analyzing the magnitude of the FT, however, the IWI catalysts FT spectra (Figure 8D) lack the Ru oxide second shell peak (Ru–O–Ru) at 3.1 Å. This absence implies that the Ru species are likely anchored on the surface through Ru-O-Ce bonds, suggesting their presence as surface species rather than existing as RuO nanoclusters.⁶² On the contrary, in the case of nickel, the spectrum exhibits both shells characteristic of Ni oxide (Figure 7D). This is supported by the Raman spectrum, wherein the presence of Ru-O-Ce bonds was identified in the catalysts prepar



Figure 6. E₀ values obtained by analyzing XANES spectra of the catalysts prepared (A) Ni K-edge and (B) Ru K-edge of the prepared catalysts.

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Table 2. Properties associated with the active phase (Ni-Ru) of the prepared catalysts						
Sample	XRD		XANES ^a		XPS	
	Form	Crystallite nm	% NiO	% RuO ₂	% NiO/Ni	% RuO ₂ /Ru
Ni(np)-Ru(np)-MM	Ni(np)	51	4.7(1)	2.6(1)	-	-
Ni(np)Ru(np)-MM	Ni(np)	48	12.4(1)	4.3(1)	-	-
Ni(A)Ru(A)-MM	Ni(A)	72	6.7(2)	32.2(4)	-	-
Ni(A)-Ru(C)-MM	Ni(A)	63	10.3(2)	24.3(2)	-	-
Ni(A)Ru(C)-MM	Ni(A)	50	13.9(2)	26.6(2)	-	-
Ni(N)-Ru(A)-MM	-	-	26.5(3)	46.7(7)	59.9	31.1
Ni(N)Ru(C)-MM	_	-	34.7(4)	45.1(3)	-	-
Ni(N)Ru(A)-MM	-	-	32.5(4)	45.1(4)	49.7	56.1
Ni(A)Ru(C)-IWI	NiO	9	81.2(2)	93.6(2)	96.5	74.8
Ni(N)Ru(A)-IWI	NiO	9	96.2(3)	97.5(3)	92.4	63.7
^a Uncertainty (Least Squa	ares Method) is giv	ven in parentheses.				

Surface atomic composition

XPS was employed to determine the surface composition of the catalysts. For the cerium 3d spectra, six peaks for Ce⁴⁺ (V, V", V", U, U", and U") were employed, representing three pairs of spin-orbit doublets, along with four peaks (two pairs of spin-orbit doublets) for Ce³⁺



Figure 7. Magnitude of the FT of the EXAFS spectra at the Ni K-edge of the catalysts prepared (A) Ni nanoparticles, (B) Ni nitrate, and (C) Ni acetate as Ni precursor and (D) prepared by IWI.







Figure 8. Magnitude of the FT of the EXAFS spectra at the Ru K-edge of the catalysts prepared (A) Ru chloride, (B) Ru acetate, and (C) Ru nanoparticles as Ru precursor and (D) prepared by IWI.

 $(V_0, V', U_0, and U')$, using peak positions and FWHM as reported in the literature.⁶³ "U" and "V" denote the $3d_{3/2}$ and $3d_{5/2}$ spin-orbit components, respectively. Deconvolution of the nickel 2p spectra was conducted by cross-referencing peak positions and FWHM reported in the literature.⁶⁴⁻⁶⁷ Nickel 2p spectra were deconvoluted using two peaks and two satellites for Ni₂O₃ (two doublets), four peaks and four satellites for NiO (four doublets), two peaks and four satellites for nickel acetate (three doublets), two peaks and two satellites for nickel nitrate (two doublets), and two peaks and two satellites for Ni⁰ (two doublets). The deconvolution of the ruthenium 3d spectra followed methodologies described in the literature.⁶⁸⁻⁷¹ For the deconvolution of Ru 3d spectra four peaks were used for RuO₂ (two doublets), two peaks for RuCl₃ (one doublet), two peaks for Ru(OH)₃ (one doublet), two peaks for ruthenium acetate (one doublet), and two peaks for Ru⁰ (one doublet). The obtained results are shown in Figure 9, whereas the surface atomic ratios are shown in Table 3. The spectra are normalized with respect to the U"" peak of ceria and to the adventitious carbon peak in the case of Ru 3d spectra. From the Ce 3d spectra deconvolution, the presence on the surface of all the catalysts of both CeO₂ (Ce⁴⁺) and, to a lesser extent, Ce₂O₃ (Ce³⁺) was identified, consistent with XANES results. The relative proportion of the two oxidation states is shown in Table 1. The surface Ce³⁺ percentage is smaller compared to the bulk, displaying diverse trends among the catalysts prepared using IWI and MM methods. Generally, IWI-prepared catalysts exhibited a higher surface percentage of Ce³⁺ compared to the catalyst prepared using the same precursors with MM, probably due to the formation of Ni-Ce-O and/or Ru-Ce-O bond, as identified from Raman spectra, resulting in higher oxygen defect concentrations.⁷² Similar trends were observed in the deconvoluted O 1s spectra (Figure 10), where the peak around 531 eV is usually attributed to oxygen species weakly absorbed on the ceria surface associated with Ce³⁺ ions (O_s), whereas the peaks around 529 eV and 527 eV are attributed to ceria lattice oxygen (O_L).^{73–77} The spectra are normalized to the O_L peak. The absorption of oxygen on the IWI-prepared catalyst surfaces was higher compared to the MM counterparts (Table 1), and the two catalysts prepared with Ni nitrate and Ru acetate with MM (Ni(N)Ru(A)-MM and Ni(N)-Ru(A)-MM) presented very similar values. IWI-prepared catalysts showed oxidized forms of Ni₂O₃, NiO, and RuO₂ on their surfaces, and in the case of Ru 3d spectra, metallic Ru presence is an indication of precursor decomposition during calcination at 450°C,²¹ demonstrating the complete oxidation (or partial decomposition) of the precursors used. Conversely, MM-prepared catalysts exhibited precursor forms together with oxidized forms of nickel and ruthenium, confirming XAS results of partial precursor oxidation through the milling process. Furthermore, an IWI-prepared catalyst analyzed before calcination (Ni(A)Ru(C)-preIWI) revealed the presence of precursors (Ni acetate and RuCl₃), along with NiO, and Ru(OH)₃ on its surface, indicating the initial precursor oxidation. Table 2





Figure 9. XP spectra of Ce 3d, Ni 2p, and Ru 3d

XP spectra of Ce 3d and Ni 2p (A) and Ru 3d (B) of (1) Ni(N)-IWI, (2) Ni(N)-MM, (3) Ni(N)Ru(A)-MM, (4) Ni(N)-Ru(A)-MM, (5) Ni(N)Ru(A)-IWI, (6) Ni(A)Ru(C)-preIWI, and (7) Ni(A)Ru(C)-IWI. The C 1s peak at 284.8 eV corresponds to adventitious carbon, whereas the peak at around 289.0 eV is related to C in the O-C=O form (acetate).

displays the percentage of NiO and RuO_2 identified on the surface in relation to total Ni and Ru amounts, respectively. For IWI-prepared catalysts, oxidation appeared gradient-like within the particle: the surface oxidation was more advanced than in the bulk. Conversely, in MM-prepared catalysts, the surface oxidation varied based on the metal type; for Ni, surface forms were more oxidized compared to the bulk, whereas for Ru, precursor amounts on the surface were higher than in the bulk. The Ni/Ru surface ratio was higher in IWI-calcined catalysts compared to MM counterparts, whereas the IWI catalyst analyzed before its calcination presents a value similar to MM catalysts.

Table 3. Surface atomic ratios and binding energies of Ni (2p3/2) and Ru (Ru 3d5/2) main peak of each form detected in the catalysts					
Sample	Ni/Ru	(Ni+Ru)/Ce or Ni/Ce	Ni forms and main peaks BE (eV)	Ru forms and main peaks BE (eV)	
Ni(N)-IWI	-	0.48	Ni ₂ O ₃ (855.0) NiO (853.5, 855.9)	-	
Ni(N)-MM	-	2.65	Ni nitrate (857.5) NiO (854.7, 855.9)	-	
Ni(N)Ru(A)-MM	1.1	1.11	Ni nitrate (857.2) NiO (853.1, 855.3)	Ru acetate (282.1) RuO ₂ (279.7)	
Ni(N)-Ru(A)-MM	0.7	2.00	Ni nitrate (857.3) NiO (854.0, 855.6)	Ru acetate (282.0) RuO ₂ (280.0)	
Ni(N)Ru(A)-IWI	4.2	0.75	Ni ₂ O ₃ (856.5) NiO (854.2, 856.4)	RuO ₂ (281.5) Ru (280.3)	
Ni(A)Ru(C)-preIWI	1.6	0.84	Ni acetate (856.5) NiO (854.7, 856.2)	RuCl ₃ (282.7) Ru(OH) ₃ (281.0)	
Ni(A)Ru(C)-IWI	8.6	0.72	Ni ₂ O ₃ (855.9) NiO (854.3, 856.2)	RuO ₂ (281.2) Ru (279.5)	







Figure 10. XP spectra of O 1s of (1) Ni(N)-IWI, (2) Ni(N)-MM, (3) Ni(N)Ru(A)-MM, (4) Ni(N)-Ru(A)-MM, (5) Ni(N)Ru(A)-IWI, (6) Ni(A)Ru(C)-preIWI, and (7) Ni(A)Ru(C)-IWI. O_s indicates the peak related to oxygen species weakly absorbed on the ceria surface, whereas O_L indicates the peak related to ceria lattice oxygen

The atomic ratio between the two metals Ni and Ru with respect to Ce can be used as an indicator of metal dispersion on the support surface, which was higher in MM-prepared catalysts.

Reducibility of the catalysts

The reducibility of selected catalysts was analyzed through H₂-TPR (Ni(N)-Ru(A)-IMI, Ni(N)Ru(A)-IWI, and Ni(A)Ru(C)-IWI). The results have been reported in Figure 11. All profiles presented two peaks at high temperatures related to the ceria support: the one at around 732°C–778°C corresponds to the reduction of bulk Ce(IV) to Ce(III) and that close to 400°C is ascribed to the surface reduction of ceria involving bridging OH group formation.⁷⁸ In the case of the Ni(A)Ru(C)-IWI catalyst, a band with a maximum at 608°C is also visible, attributed to the reduction of the Ni-O-Ce entities.⁷⁹ The Ni(N)Ru(A)-IWI sample exhibited a prominent peak at 300°C, assigned to the reduction of NiO species present on the surface.⁸⁰ Additionally, this profile displayed two peaks at lower temperatures, attributed to the reduction of Ru, at 85°C and 130°C. The presence of multiple peaks suggests the coexistence in this sample of RuO_x in a well-dispersed state and well-crystal-lized RuO₂ species, with the latter being more difficult to reduce.⁸¹ The other catalysts presented a single peak at around 100°C, assigned to the simultaneous reduction of Ni and Ru species. The presence of only one low-temperature peak associated with the reduction of both











Figure 12. Variable-temperature ammonia conversion

(A) catalysts prepared using the MM method and different precursors and (B) catalysts prepared with nickel nitrate and ruthenium acetate using different methods. The blank test was performed using the same volume of SiC without catalyst. Reaction test conditions: GHSV of 3660 h^{-1} , 0.1 g of catalyst, F/W of 15.0 L h^{-1} g⁻¹, Ar:NH₃ = 1.2:1 M, 1 bar.

metals indicates a strong interaction between them in the prepared bimetallic samples,⁸² which can explain the superior catalytic performance of these two samples compared to the other catalysts tested.

Ammonia decomposition performance

The catalysts were tested for the decomposition of ammonia reaction between 350°C and 600°C. The ammonia conversion results are shown in Figure 12. The catalyst prepared using nickel and ruthenium acetates demonstrated the lowest activity (6.5% and 19.3% at 400°C and 450°C, respectively, Figure 12A), whereas the catalyst prepared under the exact same conditions of milling (15 Hz, 10 min, B/P = 8.5) using nickel nitrate and ruthenium acetate presented an ammonia conversion of 26.2% at 400°C and 56.7% at 450°C. These findings underscore the key role of precursor types, particularly highlighting the influence of the interactions among the three precursors (including cerium oxide), since the same precursor showed results that vary with the second metal precursor, as in the example above. Notably, the two-step catalyst preparation (10 min with the Ni and ceria precursor followed by 10 min after adding the Ru precursor) revealed enhanced catalyst activity in specific precursor combinations, as in the case of the catalyst prepared using Ni nitrate and Ru acetate, which is the one that showed the highest activity. If prepared in two steps, this catalyst allows to increase the conversion of ammonia by 11.1% and 15.4% respectively, at 400°C and 450°C. However, altering the precursor order-incorporating the support and Ru acetate in the first step (10 min) and adding Ni nitrate in a subsequent 10-min step—resulted in a significant decrease in conversion (Figure 12B). This may be related to the residence time of the precursors in the synthesis system. Furthermore, by comparing the results obtained with those of the catalyst prepared by IWI method using the same precursors, a higher activity was determined in the case of the catalysts prepared with MM (Figure 12B). The comparison of the ammonia conversion levels obtained with the catalyst prepared by IWI using Ni acetate and Ru chloride reported in our previous works²¹ (Figure 12A) revealed higher conversion rate in the case of the catalyst prepared via MM prepared in two steps using Ni nitrate and Ru acetate.

The catalyst prepared with MM that presented the highest catalytic activity was tested in a long-term reaction (100 h) at 450°C. The result obtained for the conversion of ammonia is shown in Figure 13. The catalyst presented an initial deactivation, where the conversion decreased



Figure 13. Stability test in the decomposition of ammonia at 450°C Reaction test conditions: GHSV of 3660 h^{-1} , 0.1 g of catalyst, F/W of 15.0 L h^{-1} g⁻¹, Ar:NH₃ = 1.2:1 M, 1 bar.







Figure 14. EDX maps of Ni(N)-Ru(A) catalysts

(A) Ni(N)-Ru(A)-IWI and (B) Ni(N)-Ru(A)-MM (using Ni, Ru, and Ce K-peaks).

from 67% to 61%, and after around 20 h of reaction, the catalyst was stable over the subsequent 80 h of reaction at an average conversion of 56%.

Post-reaction characterization

Ni(N)-Ru(A)-MM and Ni(N)Ru(A)-IWI samples have been analyzed using HRTEM/STEM and XPS after being tested in ammonia decomposition reaction up to 600°C. From HRTEM images, the average size of the ceria particles has been estimated: 19 nm for MM-prepared catalysts and 22 nm for IWI-prepared one. Compared to pre-reaction sizes (Figure 1), there was an increase of around 6 nm for the MM-prepared catalyst ceria particles, and around 5 nm for the IWI-prepared catalyst, attributed to their partial sintering at elevated temperatures. The distribution of Ni and Ru particles remained uniform on the surface, as evidenced by EDX mapping (Figure 14).

In Figure 15, the XPS spectra of the same samples post-reaction are reported. The spectra presented oxidized forms of both metals, NiO and RuO₂, and, to a lesser extent, metallic Ni [2.7 and 0.7% of the total for Ni(N)-Ru(A)-MM and Ni(N)Ru(A)-IWI, respectively] and Ru (34.2 and 33.8%) were also identified. The analysis of the amount of Ce³⁺ compared to total Ce showed an increase in reduced ceria for both catalysts after the reaction, particularly notable for the MM-catalyst, with values of 27.0% and 23.1% for the MM- and IWI-prepared catalysts, respectively, compared to their initial states (Table 1). The surface dispersion of Ru and Ni slightly decreased for the MM-prepared catalyst to a value of 1.28, while remaining around 0.69 for the IWI catalyst, suggesting partial sintering of metal particles on the MM-prepared catalyst's surface, probably due to the absence of the calcination step during its synthesis, unlike the IWI-prepared catalyst. The Ni to Ru ratio increased to 2.6 for the MM-prepared catalyst, whereas it stayed at 4.2 for the IWI-prepared catalyst (compared to the values in Table 3 for the catalysts as prepared).

DISCUSSION

Understanding the relationship among the catalyst properties and activity

HRTEM analysis revealed initial morphology and size differences between the catalysts prepared by the two methods, mainly due to the calcination step of the catalyst prepared by IWI, which led to the complete oxidation of the precursors, as confirmed by XPS, XAS, and XRD results. Both IWI and MM catalysts showed crystallites associated with ceria and Ni and Ru forms; however, distinguishing their individual particles was difficult, indicating an intimate contact between the two active metals and with the support. The size of the metal particles was estimated through the crystallite size obtained with XRD. NiO particles in IWI-prepared catalysts measured approximately 9 nm, whereas the catalysts prepared via MM using nickel nitrate, those that showed the highest catalytic activity, did not exhibit peaks related to Ni species, suggesting their extremely small size and high dispersion. Catalysts prepared with Ni acetate displayed larger particles, ranging between 50 and 72 nm (Table 2) and those prepared using Ni nanoparticles presented supported Ni metallic particles of about 50 nm. It was not possible to determine the average crystallite dimensions of ruthenium species in the catalysts, due to its low content (1 wt.%). Notably, ammonia conversion with MM catalysts increased proportionally with decreasing the nickel particle size. The catalytic activity of the IWI-prepared catalysts also exhibited significant dependency on the precursors used, since the catalyst prepared using Ni nitrate and Ru acetate demonstrated a conversion of 40% at 450°C, whereas Ni(A)Ru(C)-IWI achieved 68.2% at the same temperature. Although no specific differences related to Ni or Ru particle size were identified in these two catalysts, distinct interactions of these metals with the ceria support were observed. In particular, the catalyst that exhibited the highest activity showed the largest amount of reduced ceria (Table 1). This trend was consistent across all catalysts,





Figure 15. XP spectra of Ce 3d and Ni 2p (A) and Ru 3d (B) of the catalysts (1) Ni(N)-Ru(A)-MM and (2) Ni(N)Ru(A)-IWI post-reaction. The C 1s peak at 284.8 eV corresponds to adventitious carbon

showing a progressive increase in conversion with the increase of reduced ceria (Figure 12). This can be attributed to a relationship between oxygen vacancies and Ce^{3+} presence, as well as enhanced electron transfer capacity.⁸³ The increase in the reduced form of ceria also correlates well with the increase in the intensity of the band related to defects in the structure of ceria of the Raman spectra. It has been reported in the literature that the interaction between oxygen vacancies in the support and the active phase facilitates the recombinative desorption of N atoms, achieving higher catalytic performance in the decomposition of NH₃, due to the increase in electron density that could be transferred from the basic sites of the support at the surface of the active phase.^{84,85} The XPS analysis of the surface composition of some of the prepared catalysts revealed that the presence of reduced surface ceria shows minimal differences between the IWI- and MM-prepared bimetallic catalysts, although the former exhibited a slightly higher Ce³⁺ content. Moreover, there were no substantial distinctions found among catalysts prepared using identical precursors (Ni nitrate and Ru acetate) but differing in one or two milling steps. Figure 16 shows the relationship between the conversion, the amount of bulk Ce³⁺, and the relationship between the D and F_{2g} areas. The strong interaction between the metal precursors and the support is further supported by the results obtained with XANES and EXAFS, where the partial



Figure 16. Relationship between the ammonia conversion, the amount of Ce^{3+} (XANES), and the relationship between the D and F_{2g} areas (Raman spectroscopy)



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oxidation of the metals due to the milling process was determined (Table 2), confirmed by XPS analysis. By guantifying the oxidized form of the two metals, considerations can be made regarding the intensity of interaction among the three precursors in the catalysts prepared via MM, given that the milling conditions employed are identical. Catalysts demonstrating a higher presence of Ni- and Ru-oxidized forms in the bulk exhibited increased ammonia conversion, particularly those prepared using nickel nitrate as a precursor. Notably, the catalyst showing the highest activity (Ni(N)-Ru(A)-MM) presented the highest presence of bulk RuO₂, followed by the same catalyst prepared with the same precursors but in a single milling step. This suggests that although the size of the nickel particles is influential in tuning the catalyst activity, the interaction between ruthenium and the other precursors is equally crucial. Regarding the surface composition, analyzed with XPS, these two catalysts exhibited distinct behaviors: the catalyst prepared in two milling steps shows a higher quantity of Ni-oxidized forms on the surface compared to the bulk, whereas for ruthenium, the opposite trend is observed. This indicates the presence of oxidation gradients within the catalyst particles. Upon analyzing the surface atomic ratios among the identified elements of the catalysts, it was possible to conclude that MM-prepared catalysts exhibit a notably higher metal dispersion on the surface compared to those prepared by IWI (1-2 vs. 0.8 in the case of bimetallic catalysts, respectively), thereby increasing the number of available active sites. Additionally, MM-prepared catalysts show a higher presence of Ru, as evidenced by the diminishing Ni/Ru ratio, dropping from 4.2-8.6 (IWI catalysts) to 0.7 in the catalyst displaying the highest catalytic activity. Comparing the catalysts prepared using Ni nitrate and Ru acetate in one or two milling steps, it can be observed that the catalyst prepared in two steps displayed the greatest presence of Ru on the surface, even surpassing that of Ni, and a higher dispersion. As regards the preparation of the catalyst by milling in two steps, depending on the type of precursors it has been demonstrated that different results are achieved: in the case of Ni(A)-Ru(C)-MM and Ni(np)-Ru(np)-MM the catalyst activity decreased, whereas in the case of Ni(N)-Ru(A)-MM it increased, indicating that the increase in milling time of the Ni precursor with ceria, from 10 to 20 min, has different effects depending on the precursor used. In general, catalysts prepared in one step presented different characteristics compared to those prepared in two steps, underlining the possibility of tuning the catalytic activity by varying the milling conditions.

The analysis of the catalyst reducibility revealed interesting differences, particularly between the two catalysts synthesized using IWI. In the catalyst prepared using Ni nitrate and Ru acetate as precursors, which exhibited the lowest activity among the two, the reduction of Ru and Ni species occurs at different temperatures, whereas in the catalyst prepared with Ni acetate and Ru chloride, which showed higher activity, the reduction of Ni and Ru species occurred simultaneously and at lower temperatures. The MM-catalyst with the highest activity among those prepared showed an H₂-TPR profile closely similar to that of Ni(A)Ru(C)-IWI. This similarity further underscores the importance of the interaction between Ni and Ru in influencing catalytic activity: higher interaction corresponds to increased activity. Prior to the reaction, the catalysts are reduced at 300°C with H₂, which means that in the case of the most active catalysts, at this temperature both the Ni and Ru species are already completely reduced. The characterization of the catalysts employed in the reaction revealed slight sintering of ceria particles, similar for both synthesis methods. Additionally, the surface dispersion of Ru and Ni slightly decreased in the MM-prepared catalyst. The estimation of the surface atomic ratio between Ni and Ru showed an increase in the surface Ni amount in the MM-prepared catalyst compared to Ru, suggesting the sintering of Ru particles. The IWI-prepared catalyst exhibited higher stability than its MM-prepared counterpart, also during the stability test, where the catalyst exhibited an initial deactivation, followed by a phase of constant activity. This enhanced stability is likely attributed to the calcination step used in the preparation of the IWI catalyst. Comparing the hydrogen production achieved at 450°C in the stable phase of the catalyst prepared by MM (305.8 mmol_{H2} g_{cat}^{-1} h^{-1}) with results obtained in the literature under similar or more favorable reaction conditions (WHSV of 136000 mL_{NH3} g_{met}^{-1} h^{-1} or lower), the prepared catalyst demonstrates superior activity compared to nickel-based catalysts, such as Ni/MgO (19.1 mmol_{H2} $g^{-1} h^{-1}$),⁵ or promoted Ni catalyst, such as 2wt.%K-15wt.%Ni/Al₂O₃ (20.5 mmol_{H2} $g^{-1} h^{-1}$),⁸⁶ and bimetallic configurations like Ni-Co supported on Y_2O_3 (168.6 mmol_{H2} g⁻¹ h⁻¹ at 500°C).⁸⁷ However, the MM-catalyst exhibits lower activity compared to catalysts containing ruthenium.⁸¹

Conclusions

Mechanochemical milling synthesis of NiRu/CeO₂ catalysts demonstrated the ability to achieve better ammonia conversion for hydrogen production compared to catalysts prepared by the conventional incipient wetness impregnation method. The use of different types of metal precursors enabled significant modulation of catalyst activity. Moreover, catalysts prepared by ball milling in two steps, milling first the Ni precursor with the ceria support and then the Ru precursor in a second step, performed better than catalysts prepared in a single step by milling together the Ni and Ru precursors with the ceria support. Specifically, catalysts with smaller Ni particle size and with a strong interaction between the metals and cerium oxide proved to be the main characteristics enhancing the catalytic activity in the decomposition of ammonia. In particular, the mechanochemical method employed demonstrated the capacity to change the oxidation state of Ni and Ru, and this capacity strongly depends on the type of precursors used. A specific interaction between the precursors and the support achieved by mechanochemistry allows an increase in oxygen vacancies in the ceria structure, usually related to an increased electron transfer capacity and better catalytic performance.

Limitations of the study

The measurement of the relative hardness of the precursors could provide valuable information regarding the characteristics of the resulting catalysts. Further transmission electron microscopy studies are needed to precisely determine the structural characteristics of the active phase.





STAR***METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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AUTHOR CONTRIBUTIONS

I.L. was responsible for the preparation and characterization of the catalysts, for the analysis of results and for writing the manuscript. I.S. was responsible for the preparation and reaction test of the catalysts. X.G. was responsible for the characterization. A.G., J.A., X.H., L.P., J.P., E.E.V., and C.M. contributed to the characterization. C.E. and J.L. provided the resources and supervised the project. All the authors reviewed the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, peptides, and recombinant proteins		
NH ₃ (28 vol. %)	Scharlab	AM02562500
Ce(NO ₃) ₃ ·6H ₂ O	Alfa Aesar	11361089
Ni(CH ₃ COO) ₂ ·4H ₂ O	Probus	Not available
$Ni(NO_3)_2 \cdot 6H_2O$	Thermo Fisher Scientific	A15540.0B
RuCl ₃	Tokyo Chemical Industries	R0074
Ru(CH ₃ COO) ₃	Biosynth	3D-XCA19632
Ru nanoparticles	Sigma Aldrich	326712-1G
Ni nanoparticles	Sigma Aldrich	577995-5G

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Jordi Llorca (jordi. llorca@upc.edu).

Materials availability

No new reagents were created in the presented research.

Data and code availability

- All data reported in this paper will be shared by the lead contact upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

METHOD DETAILS

Preparation of catalysts

CeO₂ was prepared by adding NH₃ (28 vol. %, Scharlab) to a magnetically stirred solution of Ce(NO₃)₃·6H₂O (Alfa Aesar) in water, until pH~9. The precipitate was filtered (with a porous plate n° 4) and calcined at 500°C (4 h, 5°C min⁻¹). All prepared catalysts have a content of 5 wt.% Ni and 1 wt.% Ru. The catalysts synthesized using the incipient wetness impregnation (IWI) method were prepared by co-impregnation with an aqueous solution of Ni and Ru precursors. Between each impregnation phase the catalyst is dried at 100°C for approximately 10 min and ground. After the impregnation, the catalysts were dried at 100°C for 24 h, ground, and then calcined at 450°C for 4 h using a ramp of 5°C min⁻¹. The monometallic counterparts were prepared following the same method, but dissolving only the respective precursor in distilled water. The precursors that were used for IWI catalysts are Ni(CH₃COO)₂·4H₂O (Probus) and Ni(NO₃)₂·6H₂O (Thermo Fisher Scientific) for Ni, while those for Ru were RuCl₃ (Tokyo Chemical Industries) and Ru(CH₃COO)₃ (Biosynth). The catalysts synthesized using the mechanochemical (MM) method were prepared employing a moderate-energy vibrating ball mill (Pulverisette 23, FRITSCH). Ceria and the chosen Ni and Ru precursors were loaded into a 15 mL zirconia vial equipped with a 15 mm diameter high wear-resistant zirconia ball. The vial was agitated at an oscillation frequency of 15 Hz for 10 min, with a ball/powder ratio of 8.5/1. The same precursors used for the IWI method were also employed for the catalysts prepared with the MM method, however, in this case, Ru nanoparticles (Sigma Aldrich) and Ni nanoparticles (Sigma Aldrich) were also used. The monometallic counterparts were prepared following the same method, introducing in the vial only the corresponding precursor and ceria. Furthermore, a comparison was made by introducing the precursors into the vial in two steps: in a first step the first precursor and CeO₂ were milled together, followed by the addition of the second precursor in a subsequent step. For these catalysts the same milling conditions were used, varying the total time and both steps lasted 10 min. The nomenclature of the catalysts was defined by indicating the type of precursors (nitrate: N, acetate: A, chloride: C and nanoparticles: np). In the case of the two-step preparation, this was indicated with a line between the two precursors, starting with the precursor used in the first phase. Finally, the method used was specified by adding IWI or MM at the end.

Catalyst characterization techniques

High-resolution transmission electron microscopy (HRTEM) images were acquired in a field emission gun FEI *Tecnai* F20 microscope operated at 200 kV. The crystal structure was analyzed using the frequency spots in the corresponding fast Fourier transformations (FFTs) using





Gatan Digital Micrograph software. The HAADF images and EDX maps were acquired in a double corrected and monochromated Thermo Fisher Spectra 300 microscope operated at 300 kV using scanning transmission electron microscopy (STEM) mode. The HAADF images and EDX maps have been analyzed by Velox software. Raman spectroscopy was performed with a Renishaw inVia Qontor confocal Raman microscope (532.1 nm laser) and a Leica DM2700 M microscope (×50). An exposure time of 0.5 s and 12 repetitions were used to acquire the spectra. Ex situ X-ray absorption spectroscopy (XAS) and X-ray diffraction (XRD) analysis were conducted at the NOTOS beamline of the ALBA Synchrotron Light Source facility (Cerdanyola del Vallès, Barcelona, Spain). The synchrotron light coming from a bending magnet has been first vertically collimated, then monochromatized using two pairs of water-cooled Si(111) crystals and finally focused on the sample position down to \sim 500 x 500 μ m². Si and Rh stripe coatings of the two mirrors were opportunely chosen to guarantee the higher harmonics rejection. The beamline was initially set to an energy configuration capable of exciting Ce atoms around the Ce L3-edge (5723 eV) and the samples were measured in transmission mode using three DC current-type ionization chambers (IO, I1 and I2, where a reference is placed between the chambers I1 and I2, Oken) filled with the appropriate mixture of inert gases. Subsequently, Ni was analyzed by changing the configuration to excite the Ni atoms around its K-edge (8333 eV) and the samples were measured in fluorescence (13-element Si detector, Canberra). For Ru fluorescence measurement, the setup was modified using the Rh stripe to reach the high energies required by its K-edge (22117 eV). Just before the Ru K absorption edge, XRD diffractograms were recorded at 21000 eV using a Mythen-I detector (Dectris) in a 20 range of 42°, measuring with three different detector positions for 5 s each. All catalysts were analyzed in pellet form, and prepared by diluting the sample with cellulose. A minimum of three extended X-ray absorption fine structure spectra (EXAFS) were acquired to ensure spectral reproducibility and a good signal-to-noise ratio. Data analysis and treatment were conducted using the Demeter software package.⁸⁹ The Fourier Transform (FT) was obtained on k^2 -weighted EXAFS oscillations in the range of 2.9–10.8 Å⁻¹ and R space range of 0–6 Å. The average crystallite size, denoted as L [nm], was calculated from the highest intensity peak from XRD patterns using the Debye-Scherrer equation (Equation 1):

$$L = \frac{K\lambda}{\beta \cos \gamma}$$
 (Equation 1)

Where 0.89 was chosen for the shape factor K, λ [nm] is the wavelength, β [rad] is the full width at half maximum (FWHM), and γ [rad] is the Bragg angle.

X-ray photoelectron spectroscopy (XPS) analysis utilized a *SPECS* system equipped with a *PHOIBOS* 150 EP Hemispherical Energy Analyzer with an MCD-9 detector, using a Mg K α X-ray source of 1253.6 eV energy and 150 W power, and a pass energy of 20 eV. The X-ray source is placed at 54° with respect to the analyzer axis and is calibrated by using the Ag 3d_{5/2} line with a full width at half maximum of 1.211 eV. A flood gun operating at 15 μ A and 1.5 eV was used to compensate the charge. The measurements were calibrated against the C 1s peak at a binding energy (BE) of 284.7 eV and the U''' Ce 3days peak at 916.9 eV. CasaXPS software (*Casa Software Ltd.*, Teignmouth, UK) was employed for result analysis. Atomic fractions were estimated by normalizing peak areas based on acquisition parameters following background subtraction, experimental sensitivity factors, and the analyzer transmission efficiency provided by the manufacturer. H₂-temperature programmed reduction (H₂-TPR) experiments were conducted using a *Chemstar-TPX* instrument equipped with a Thermal Conductivity Detector (TCD). Initially, 50 mg of the sample were heated from room temperature (RT) to 450°C in an argon flow rate of 50 mL min⁻¹, with a heating rate of 10 °C min⁻¹. Then the samples were maintained at 450°C for 10 min, before being cooled down to 50°C under an argon flow. Subsequently, H₂-TPR was carried out from 50°C to 850°C at a rate of 10 °C min⁻¹ in a mixture of 10% H₂ in Ar, with a total flow rate of 50 mL min⁻¹. The samples were then held at 500°C for 30 min.

Catalytic tests

Ammonia decomposition experiments were conducted in a stainless-steel reactor under atmospheric pressure conditions. 0.1 g of the catalyst were combined with SiC to achieve a bed volume of approximately 0.41 cm³. Temperature-dependent tests for the reaction were performed within the temperature range of 350°C–600°C, at steps of 50°C, each lasting 30 min. The blank test was performed without any catalyst using the same volume of SiC. Stability tests were conducted for 100 h at 450°C. The gas mixture used had an Ar:NH₃ molar ratio of 1.2:1, and the total gas flow rate was set at 25 mL min⁻¹. The control of the gas flow was achieved using *Bronkhorst* mass flow controllers. Before starting the catalytic tests, the samples were activated at 300°C for 1 h in a 10 vol. % H₂ atmosphere (80 mL min⁻¹, N₂ balance). The reaction progress was continuously monitored using a mass spectrometer (*OmniStar*) with a measurement error of \pm 5%. Ammonia conversion (denoted as x [%]) was determined using Equation 2 using the inlet and outlet molar flow rates of ammonia, *F_{inNH3}* and *F_{outNH3}*, respectively (measured in mol NH₃ s⁻¹).

$$x = \frac{F_{inNH_3} - F_{outNH_3}}{F_{inNH_3}} \times 100$$
 (Equation 2)

QUANTIFICATION AND STATISTICAL ANALYSIS

The standard deviation of the particle size measured from TEM images (Figure 1) is estimated by determining the width of the Gaussian curve when the frequency is half of the maximum value, and then dividing it width by 2.

In Demeter software package, Athena uses Least Squares Method for XANES spectra fitting (Table 1) and linear combination analysis (Table 2).