

In-Situ Generation of CO<sub>x</sub>-Free H<sub>2</sub> by Catalytic Ammonia Decomposition over Ru-Al-Monoliths.  
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## Abstract

Ru catalysts supported on alumina coated monoliths has been prepared employing three different precursor, which are ruthenium chloride, ruthenium nitrosyl nitrate and ruthenium acetyl acetonate, by an equilibrium adsorption method. The Ru particle sizes could be controlled varying the metal precursor salt. Among the prepared catalysts, Ru catalyst prepared from nitrosyl nitrate exhibited the highest activity which is concomitant to the largest mean Ru particle size of 3.5 nm. The values of the apparent activation energy calculated from the Arrhenius equation are according to the Temkin-Phyzev model, indicating that the recombinative desorption of N ad-atoms is the rate-determining step of the reaction. However, the ratio between the kinetic orders with respect to ammonia and hydrogen ( $-\alpha/\beta$ ), is not in agreement to the value predicted by Temkin formalism. This fact could be related to the different operational conditions used during the reaction, and/or catalyst nature, but not to any change on the controlling step of the reaction.

**Keywords:** Structured Reactors; Ru-Catalyst; Ammonia Decomposition; Kinetics; CO<sub>x</sub>-Free; Nanoparticle.

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## 29 **1. Introduction**

30 Hydrogen combined with PEM fuel cells have attracted great interest recently as  
31 substitute of fossil fuels in locomotion and other fields. However, hydrogen storage and  
32 transportation involves great technical challenges to overcome. Its low energy per  
33 volume unit, along with its flammability are factors which make difficult its storage as  
34 gas at normal pressure [1]. Because of that, several technologies for hydrogen storage  
35 have been proposed. These include storage at high pressures, either in liquid form or  
36 adsorbed in a solid [2,3]. However, liquid H<sub>2</sub> requires high energy consumption and  
37 expensive leak-free tanks. A more interesting option is the chemical storage in form of  
38 easily storable and with high hydrogen content compounds, such as light hydrocarbons  
39 or ammonia. These compounds must be decomposed in-situ to release a hydrogen  
40 stream to feed the fuel cell. Recently, the production of hydrogen from the  
41 decomposition of light hydrocarbons as methane, ethane or ethanol has attracted a great  
42 researching interest [4]. Nevertheless, all these processes have the inherent drawback of  
43 producing CO<sub>x</sub> along with the H<sub>2</sub> stream. CO<sub>x</sub> acts as poisons for PEM fuel cells, even  
44 at concentrations as low as a few ppm [5]. Therefore, systems based on hydrocarbons  
45 decomposition must include a series of purification steps, like desulfurization, water gas  
46 shift, methanation, and preferential oxidation to reduce CO<sub>x</sub> levels in the reformer outlet  
47 gas. Ammonia, on the other hand, produces a CO<sub>x</sub>-free stream of hydrogen, and  
48 unconverted ammonia can be easily reduced to safe levels in one-step adsorption. In  
49 addition, ammonia has a high H<sub>2</sub> content (17.7% wt) and can be stored in liquid form at  
50 mild conditions (room temperature and 6 atms). For the whole process of H<sub>2</sub> storage as  
51 NH<sub>3</sub> to be environmentally sustainable, the H<sub>2</sub> and the energy required for synthesis and  
52 decomposition process should be produced using renewable energies. On the other  
53 hand, the energy needed for decomposition of NH<sub>3</sub> could be generated using a small

54 percentage of the energy stored in  $\text{NH}_3$ . This percentage will be minimized if a catalyst  
55 active at the lowest temperature as possible is used and obtaining a high-purity  $\text{H}_2$   
56 stream. All this characteristic make more economically feasible the hydrogen  
57 production from ammonia than others career [6].

58 In the last years, ammonia decomposition reaction has been extensively studied for  $\text{H}_2$   
59 generation, including the catalytic active phase to the reactor structure and design [7].

60 Yin et al., reported that ruthenium exhibits the highest TOF in ammonia decomposition  
61 [8]. Therefore, research has been focused on ruthenium supported on different carriers,  
62 such as alumina [9–11] or others transition metal oxides like  $\text{MgO}$  [12], activated  
63 carbon [13,14], or carbon nanotubes/nanofibers [15–18]. Bimetallic combinations of  
64 transition metals (MoCo) have also proved to be active enough to replace ruthenium  
65 catalyst [19–23]. From the point of view of reactor design, microstructured reactors  
66 have improved the performance of the conventional packed bed reactors [24–27].

67 One century after the development of Habber-Bosch process, ammonia mechanism still  
68 being investigated by several authors to obtain more precise information about reaction  
69 mechanism and to explain the wide range differences observed on the kinetic  
70 parameters.

71 The mechanism of ammonia decomposition reaction has been grouped in two limiting  
72 cases: (i) effect of ammonia concentration, (ii) effect of hydrogen concentration.

73 At low ammonia concentrations and low temperatures [28,29], the reaction rate shows  
74 non-dependence with respect to ammonia. However, at high temperatures the reaction  
75 becomes first-order with respect to ammonia [30,31]. These transition temperature has  
76 been investigated by Tamaru [32] and Chellappa [33], and explained by a change of  
77 reaction mechanism. At temperature below 650 K the recombinative desorption of N-

78 adsorbed species is the rate-determining step (RDS), while at high temperature (< 750  
79 K), cleavage of N-H bond on NH<sub>3</sub> adsorbed species is the RDS [30,31].

80 At high hydrogen partial pressure and low temperature, the reaction is found to be  
81 inhibited by hydrogen [33]. In these operational conditions the reaction is explained by  
82 Temkin-Phyzev mechanism [32,34]. This reaction mechanism assumes that both  
83 recombinative desorption of N-species and N-H cleavage are the RDS. Inhibitory  
84 effects of hydrogen could be associated to the re-hydrogenation of N-species adsorbed  
85 or by blocking the active sites by competitive adsorption, where ammonia  
86 decomposition takes place. At high temperature and low hydrogen partial pressure (low  
87 conversion), Temkin-Phyzev model could be re-written as Tamaru's model [32,35].

88 This work is a new contribution of our group to the study of the ammonia  
89 decomposition reaction, using nickel catalysts supported on honeycomb monoliths  
90 prepared by electrostatic adsorption structured reactors [27]. Herein, our aim is to  
91 investigate the performance of Ru-based reactors, and in particular, to obtain uniformly  
92 dispersed Ru nanoparticles in alumina coated monoliths. To this end, different Ru  
93 precursors were tested. The catalysts after reduction were characterized by STEM, TPR  
94 and XPS and tested in the decomposition of pure ammonia. Also a kinetic study is  
95 carried out in order to determine the effect of precursor synthesis on the main apparent  
96 kinetic parameters: activation energy and reaction orders.

## 97 **2. Experimental**

### 98 2.1. Materials and Catalyst Preparation

99 Cordierite monoliths were carved out from blocks supplied by Corning (400 cpsi) to the  
100 desired dimensions (65 mm long, 10 mm diameter, ca. 2,4 g). These monoliths were  
101 impregnated with alumina by a dip coating method, as described elsewhere [27]. A sol-  
102 gel was prepared by mixing pseudoboehmite (AlOOH pural, from Sasol), 0.3 M nitric

103 acid and urea, in 2:5:1 proportion. After stirring for 24h, monoliths were immersed in  
104 the mixture until ensuring the absence of air bubbles in the channels. Following up, the  
105 excess of liquid was removed with pressurized air and the monoliths were dried for 24  
106 hours at room temperature while rotating around its axis. Later, the monoliths were  
107 calcined under 100 ml/min air flow for 2 hours at 873 K, at a heating rate of 1 K/min.

108 The deposition of ruthenium nanoparticles was carried out by equilibrium adsorption or  
109 electrostatic adsorption method [36]. In this method, the ions of precursor in solution  
110 are bound by electrostatic forces to the sites of opposite charge at support surface. Three  
111 different Ru precursors were used. Ruthenium Chloride ( $\text{RuCl}_3 \cdot x\text{H}_2\text{O}$ ) dissolved in 10  
112 ml of 0.1M HCl, ruthenium nitrosyl nitrate ( $\text{Ru}(\text{NO}_3)_3(\text{NO})$ ) dissolved in 10 ml of  
113 distilled water and ruthenium acetyl acetonate ( $\text{RuC}_{15}\text{H}_{21}\text{O}_6$ ) dissolved in 10 ml of  
114 toluene. The weight of ruthenium in solution was calculated on the basis of a nominal  
115 weight percentage of 5 wt% with respect to alumina coating. The alumina coated  
116 monoliths were fitted to a vial with the impregnating solution and rotated continuously  
117 perpendicular to its axis during 24 hours. In this process, the liquid with the precursor  
118 flows through the monoliths channels and it is concomitantly homogenized due to the  
119 rotating movement. After this process, the liquid inside the channels was removed with  
120 pressurized air and the monoliths were dried under rotation during 24 hours. Finally,  
121 they were calcined in 100 ml/min of  $\text{N}_2$  at 873 K during 2 h, using a heating rate of 1  
122 K/min. The monolith samples were denoted as Ru(Cl)/Al/M, Ru(NN)/Al/M and  
123 Ru(acac)/Al/M for catalysts prepared from ruthenium chloride, ruthenium nitrosyl  
124 nitrate and ruthenium acetyl acetonate, respectively. The actual Ru content was  
125 analyzed by inductive coupled plasma optical emission spectroscopy (ICP-OES) after  
126 digestion of the samples in lithium borohydride. In addition, Ni catalyst was also

127 prepared from a nickel nitrate solution as described previously [23]. The actual Ni  
128 loading determined by ICP-OES was 15 wt%.

129

130

## 131 2.2. Catalytic Testing

132 Catalyst reactivity system consists of a continuous-flow 15 mm i.d. quartz reactor inside  
133 horizontal furnace with a temperature controller (Eurotherm). The monolithic nickel  
134 catalyst was wrapped with quartz stripe and it was tightly fitted to the wall of the quartz  
135 reactor to avoid channeling. Subsequently, the reactor was placed in the constant  
136 temperature zone of the furnace. The reaction testing was carried out after catalyst  
137 reduction in hydrogen atmosphere (100 ml/min H<sub>2</sub>:N<sub>2</sub>, 50:50) at 773 K for 1 h.  
138 Catalysts were tested between 573 and 973 K using 100 ml/min of pure anhydrous  
139 ammonia as feed gas. The outlet reaction gases were analyzed with an Agilent Micro  
140 GC 3000A. H<sub>2</sub> and N<sub>2</sub> were analyzed in a molsieve column and ammonia in a Plot-Q  
141 column. To ensure repeatability, 2–3 separate GC samples were taken and averaged for  
142 each experimental data point, and analyses were typically within  $\pm 3\%$  of each other.  
143 The conversion was calculated taking into account the variations in the flow due to the  
144 mole increase in the reaction.

## 145 2.3. Catalyst Characterization

146 The catalysts after reaction were tested by different techniques. Temperature  
147 programme reduction (TPR) was carried out in a Micromeritics AutoChem II 2920. In  
148 TPR experiments, the catalyst was heated in 50 ml/min (STP) of 10% H<sub>2</sub> in Ar up to  
149 1273 K with a heating rate of 10 K/min.

150 XPS technique was used to measure the Ru oxidation state. The apparatus was an  
151 ESCAPlus Omnicrom system equipped with a Mg K $\alpha$  radiation source to excite the

152 sample. Calibration of the instrument was done with Ag 3d<sub>5/2</sub> line at 368.27eV. All  
 153 measurements were performed under UHV, better than 10<sup>-10</sup> Torr. Internal referencing  
 154 of spectrometer energies was made using the dominating Al 2p peak of the support, at  
 155 74.0 eV and corroborated with the C 1s signal at 284.4 eV. The curve fitting of the  
 156 spectra was performed using CASA XPS software after applying a Shirley baseline.  
 157 Catalyst particle size was studied by scanning transmission electron microscopy  
 158 (STEM) using a FEI TECNAI F30 electron microscope equipped with Gatan Energy  
 159 Filter and cold field emission gun (FEG) operated at 300 kV with 1.5 Å lattice  
 160 resolution. TEM specimens were prepared by ultrasonic dispersion in ethanol of powder  
 161 retrieved from the monoliths. A drop of the suspension was applied to a holey carbon  
 162 support grid.

### 163 3. Results and Discussion

#### 164 3.1 Textural and morphological characterization

165 Table 1 shows the textural properties of the prepared catalysts. All Ru catalysts exhibit  
 166 similar textural properties and the surface area is comparable to that of the support (246  
 167 m<sup>2</sup>/g). There are not significant differences between the Ru content deposited using the  
 168 different salt precursors.

169 Table 1. Textural properties measured by N<sub>2</sub> physisorption and surface composition  
 170 measured by XPS of Ru catalysts on alumina coated monoliths prepared from different  
 171 precursors.

Catalyst	Ru/Al <sup>1</sup> wt%	Surface Area (m <sup>2</sup> /g)	Pore diameter (Å)	Ru/Al <sup>2</sup> wt %	Ru <sup>0</sup> /Ru <sub>total</sub> <sup>2</sup> at. %
Ru(NN)/Al/M	3.0	238	60	6.7	69.1
Ru(Cl)/Al/M	3.6	215	60	6.4	5.7
Ru(acac)/Al/M	3.0	233	55	5.6	7.4

172 <sup>1</sup>Measured by ICP-OES

173 <sup>2</sup>Measured by XPS

174

175 The ruthenium particle size was characterized by STEM. Fig. 1 shows representative

176 STEM images and particle size distribution (inset) for the several catalysts prepared.

177 For Ru(NN)/Al/M (Fig. 1a, and 1b) the particles are spherical with heterogeneous

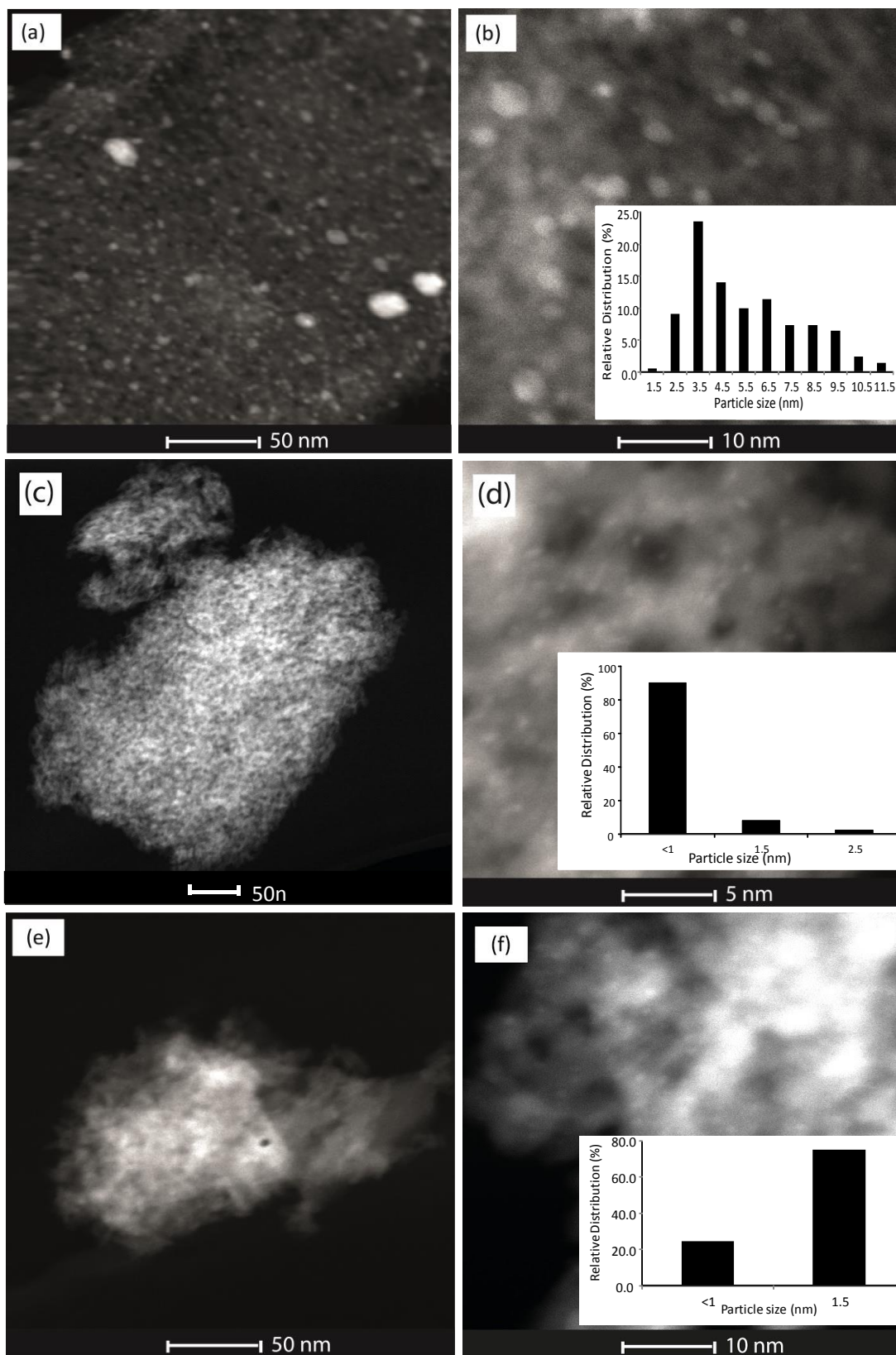
178 distribution particle sizes, ranging between 1 and 10 nm with a mean size of 3.5 nm. On

179 the other hand, for Ru(Cl)/Al/M and Ru(acac)/Al/M (Fig. 1c to 1f) the particle size are

180 predominantly smaller than 1.5 nm, with a homogeneous distribution. The mean particle

181 size is subnanometric for Ru(Cl)/Al/M and 1.5 nm for Ru(acac)/Al/M.





182

183 **Fig. 1.** Representative STEM images at two different magnifications and particle size  
 184 distribution of catalysts used in reaction: (a,b) Ru(NN)/Al/M; (c,d) Ru(Cl)/Al/M y (e,f)  
 185 Ru(acac)/Al/M.

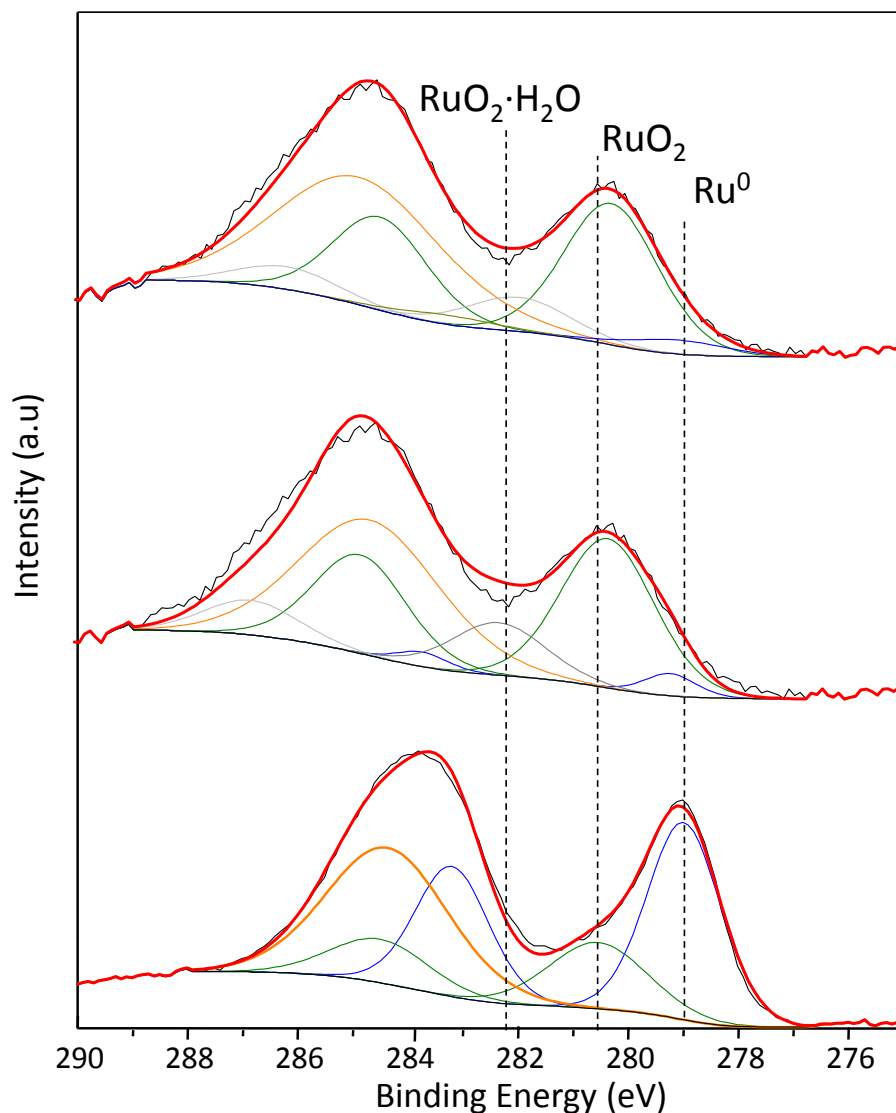
186

### 187 3.2. Ru oxidation state characterization

188 Fig. 2 shows the deconvolution of XPS Ru signal according to the literature  
189 assignments [25-28]. In addition, the quantification of Ru/Al and  $\text{Ru}^0/\text{Ru}_{\text{total}}$  atomic  
190 ratios determined by XPS are displayed in table 1. The Ru/Al ratio determined by XPS  
191 is similar for all the catalysts synthesized and these ratios are around twice that those  
192 determined by ICP. This result indicates that the surface of the alumina coating is  
193 enriched by Ru.

194 In the deconvolution of Ru 3d XPS signal of Ru(NN)/Al/M (spectrum a of Fig. 2) only  
195 the peaks at 279.1 eV and 280.5 eV are present, which are associated to  $\text{Ru}^0$  and  $\text{RuO}_2$   
196 species [37–40], respectively. However, for the catalyst Ru(Cl)/Al/M and  
197 Ru(acac)/Al/M, besides the peaks associated to  $\text{Ru}^0$  and  $\text{RuO}_2$ , is observed a new signal  
198 centered at 282.3 eV, which can be ascribed to hydrated  $\text{RuO}_2$  species [41].  
199 Additionally, a peak centered at 284.4 eV has been observed in all the catalysts which is  
200 attributed to carbon contamination.

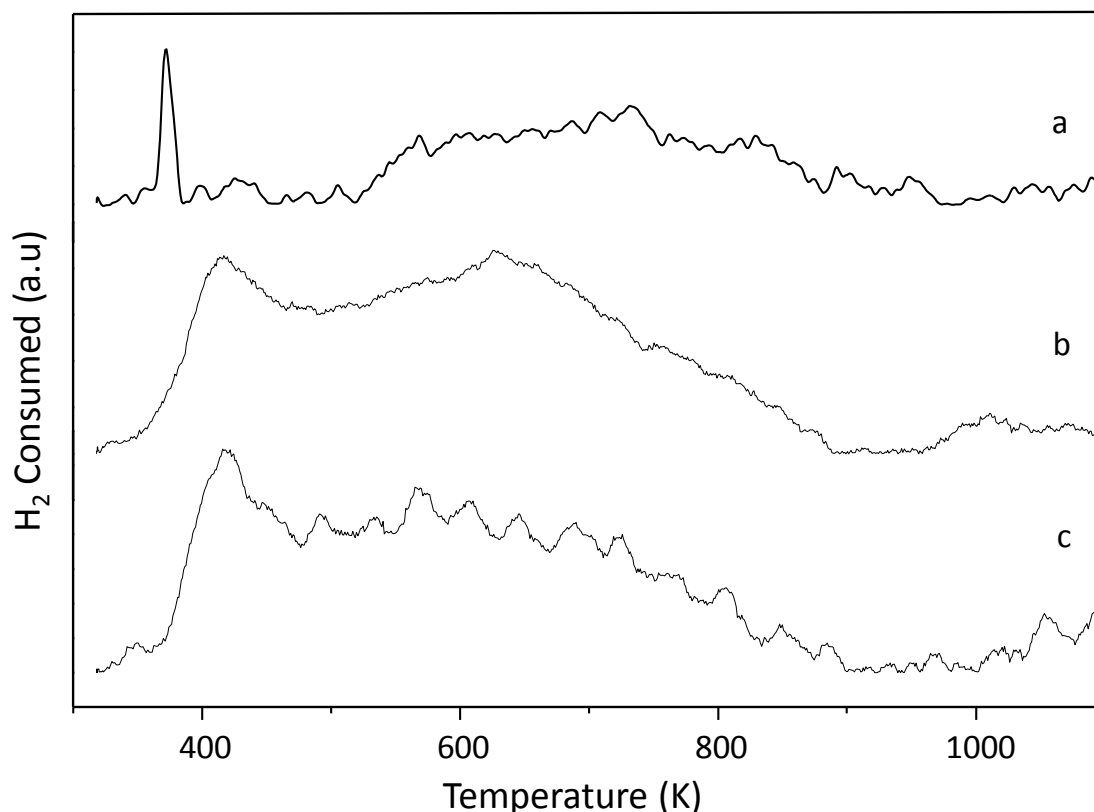
201 The catalysts Ru(Cl)/Al/M and Ru(acac)/Al/M show  $\text{Ru}^0/\text{Ru}_{\text{(total)}}$  ratios substantially  
202 smaller than the catalyst Ru(NN)Al/M. In the XPS spectra of Ru(Cl)/Al/M, a signal at  
203 198.5 eV was observed indicating the presence of chlorine ions. It is well known that  
204 electronegative species such as chlorine can increase the oxidation state of metals  
205 [42,43], which could explain the lower  $\text{Ru}^0/\text{Ru}_{\text{(total)}}$  ratios. However, the effect of  
206 electron-withdrawing species cannot be invoked to explain the lower  $\text{Ru}^0/\text{Ru}_{\text{(total)}}$  ratios  
207 of Ru(acac)/Al/M. Therefore, the most plausible explanation is that metal oxidation  
208 state is governed by interaction with the support, which is particle-size dependent. Thus,  
209 the smaller the Ru particles, the stronger is the metal support interaction, leading to  
210 lower  $\text{Ru}^0/\text{Ru}_{\text{(total)}}$  ratios.



211 **Fig. 2.** XP Spectra of 3d core level of ruthenium of the catalyst used in reaction: (a)  
 212 Ru(NN)/Al/M; (b) Ru(Cl)/Al/M and (c) Ru(acac)/Al/M.  
 213

214  
 215 The passivated catalysts after reaction were also studied by temperature programmed  
 216 reduction (Fig. 3). The TPR profile of Ru(NN)/Al/M shows two peaks, one intense peak  
 217 at low temperature (ca. 380 K) which can be attributed to surface oxidized large Ru  
 218 nanoparticles and a broad peak at higher temperatures which can be attributed to  
 219 oxidized smaller Ru particles, interacting strongly with the support. The other two  
 220 catalysts, Ru(Cl)/Al/M and Ru(acac)/Al/M, exhibit similar TPR profiles with two  
 221 shoulders, one at ca.430 K and other in the range of 550-750 K which correspond to  
 222 RuO<sub>2</sub> particles with some kind of interaction with support. The peak showed at low

223 temperatures (380 K) is absent in these two latter catalysts, suggesting that the smaller  
224 particles, present in these catalysts, have stronger interaction with the support. On the  
225 other hand, the larger particles found for Ru(NN)/Al/M are oxidized only on the  
226 outermost surface but the inner bulk metal remains in reduced state in agreement with  
227 XPS results [21].

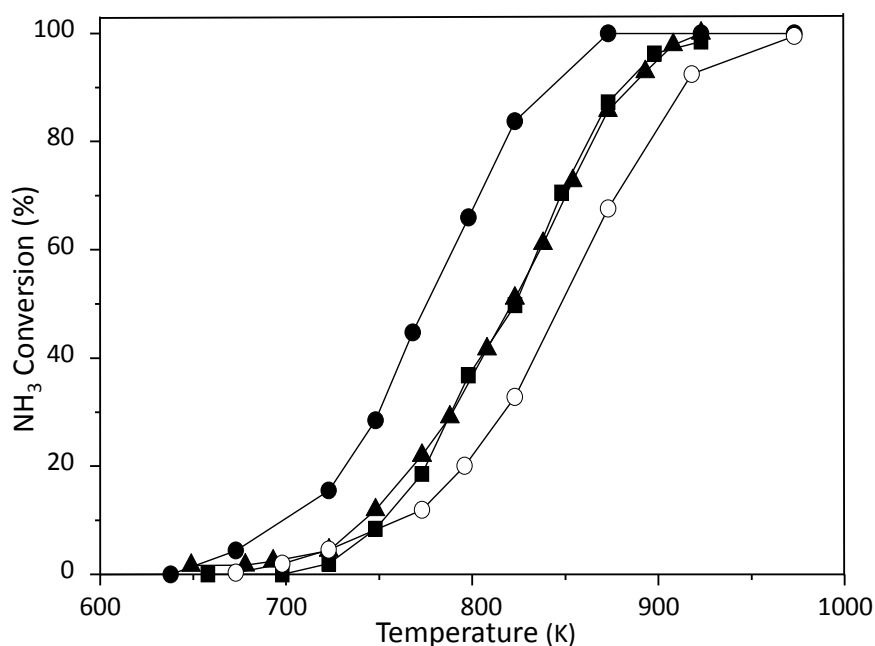


228 **Fig. 3.** TPR profile of catalysts passivated after reaction for Ru catalysts (a)  
229 Ru(NN)/Al/M; (b) Ru(Cl)/Al/M and (c) Ru(acac)/Al/M.  
230  
231

### 232 3.3. Catalytic testing in NH<sub>3</sub> decomposition

233 Fig. 4 shows the NH<sub>3</sub> conversion vs. temperature plot for the Ru-based catalysts. This  
234 Figure also includes the curve obtained for the catalyst that contains a 15% Ni on  
235 Alumina coated monolith. The preparation and characterisation of this Ni/Al/M catalyst  
236 is reported in a previous publication [44]. All Ru-based catalyst exhibit higher  
237 conversion than Ni-based catalyst although the metal loading is five-fold higher for the  
238 later, ca. 3 and 15 wt%/Al<sub>2</sub>O<sub>3</sub>, respectively. Noticeably, the mean particle size of Ni was

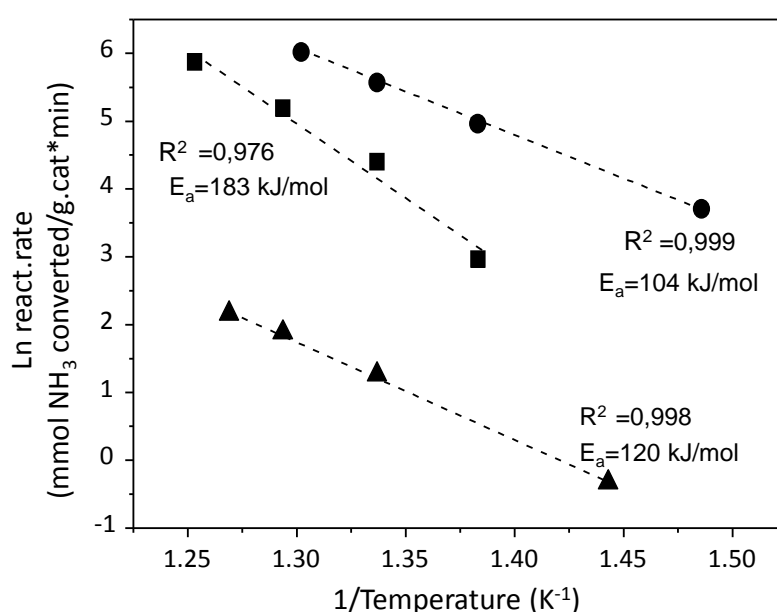
239 larger than for Ru catalysts, 6 nm vs. 2.5 nm for Ru(NN)/Al/M. This seems to indicate a  
240 larger intrinsic activity of Ru metal, in agreement with the literature [21].



241 **Fig. 4.** NH<sub>3</sub> conversion at steady state as a function of the reaction temperature for  
242 different catalysts. (●) Ru(NN)/Al/M; (■) Ru(Cl)/Al/M; (▲) Ru(acac)/Al/M and (○)  
243 Ni/Al<sub>2</sub>O<sub>3</sub>/Monolith. Conditions: 100 ml/min pure NH<sub>3</sub>, space velocity: 35000 h<sup>-1</sup>/g  
244 Ni/Al<sub>2</sub>O<sub>3</sub>.

245  
246 Comparing among Ru-based catalysts, the one prepared with nitrosyl nitrate  
247 (Ru(NN)/Al/M) exhibits higher conversion, for all temperatures studied, than those  
248 prepared with other precursors. Since the main difference between Ru(NN)/Al/M and  
249 the other catalysts is the particle size, it is reasonable to think that this fact may be the  
250 responsible of the different catalytic behaviour. Jacobsen and cols. reported that B<sub>5</sub>-type  
251 sites have the adequate geometry for the ammonia decomposition [45]. The number of  
252 these sites decreases as the particle size diminishes. This fact has been experimentally  
253 confirmed by several authors, who observed an increase of the turnover frequency as the  
254 metal particle size increased [46–48]. Xu and cols. [49] reported that there was an  
255 optimum Ru particle size of 2.2 nm that maximized turnover frequency. They also  
256 found that the apparent activation energy increased from 79 kJ mol<sup>-1</sup> to 122 kJ mol<sup>-1</sup> as  
257 the mean Ru particle size decreased from 4.6 to 1.9 nm. The apparent activation  
258

259 energies of the different catalysts prepared were calculated from the Arrhenius equation  
 260 (Fig. 5) at low conversion (<10%), in order to operate into differential condition. The  
 261 apparent activation energy for Ru(NN)/Al/M was 104 kJ mol<sup>-1</sup>, which was smaller than  
 262 those determined for Ru(acac)/Al/M and Ru(Cl)/Al/M, 120 kJ mol<sup>-1</sup> and 183 kJ mol<sup>-1</sup>,  
 263 respectively. It is possible that presence of chlorine ions may act as inhibitor in the  
 264 catalyst prepared with ruthenium chloride, contributing to the lower conversion and  
 265 higher apparent activation energy for this catalyst.



266 **Fig. 5.** Arrhenius plot and estimation of the apparent activation energies for the Ru  
 267 catalysts from different precursors. (●) Ru(NN)/Al/M; (■) Ru(Cl)/Al/M; (▲)  
 268 Ru(acac)/Al/M.  
 269  
 270

### 271 3.4. Kinetic Model

272 The development of an efficient reactor requires necessarily a depth understanding of  
 273 kinetic aspects. Many researches related to ammonia decomposition reaction have been  
 274 carried out under high vacuum condition and with model catalysts. These conditions are  
 275 not comparable to fuel cell systems fed by ammonia decomposition reaction where  
 276 usually high ammonia concentration, atmospheric pressure and high temperature have.

277 Regarding to ammonia decomposition kinetics, many author have fitted the  
 278 experimental data using the Themkyn-Pyzhev model:

$$(-r_{NH_3}) = k \left[ \left( \frac{PNH_3^2}{PH_2^3} \right)^m - \frac{PN_2^2}{Keq^2} \left( \frac{PH_2^3}{PNH_3^2} \right) \right] \quad (1)$$

279 Where  $m$  is a constant related to the non-uniformity surface, and the second term  
 280 corresponds to the contribution of the reversible equation (i.e. que driving force to the  
 281 equilibrium). According with the experimental conditions (673 K -1023 K), and to the  
 282 results previously reported [50], this term is virtually zero, indicating that the reaction is  
 283 not limited by equilibrium, and therefore this contribution can be neglected.

284 The Temkin-Pyzhev model may be rewrite to the power law rate expression:

$$(-r_{NH_3}) = k' PNH_3^\alpha PH_2^\beta \quad (2)$$

285 where  $k$  and  $k'$  follows an Arrhenius dependence with the temperature, [32,51,52] and  $\alpha$   
 286 and  $\beta$  are the kinetic orders with respect ammonia and hydrogen, respectively.

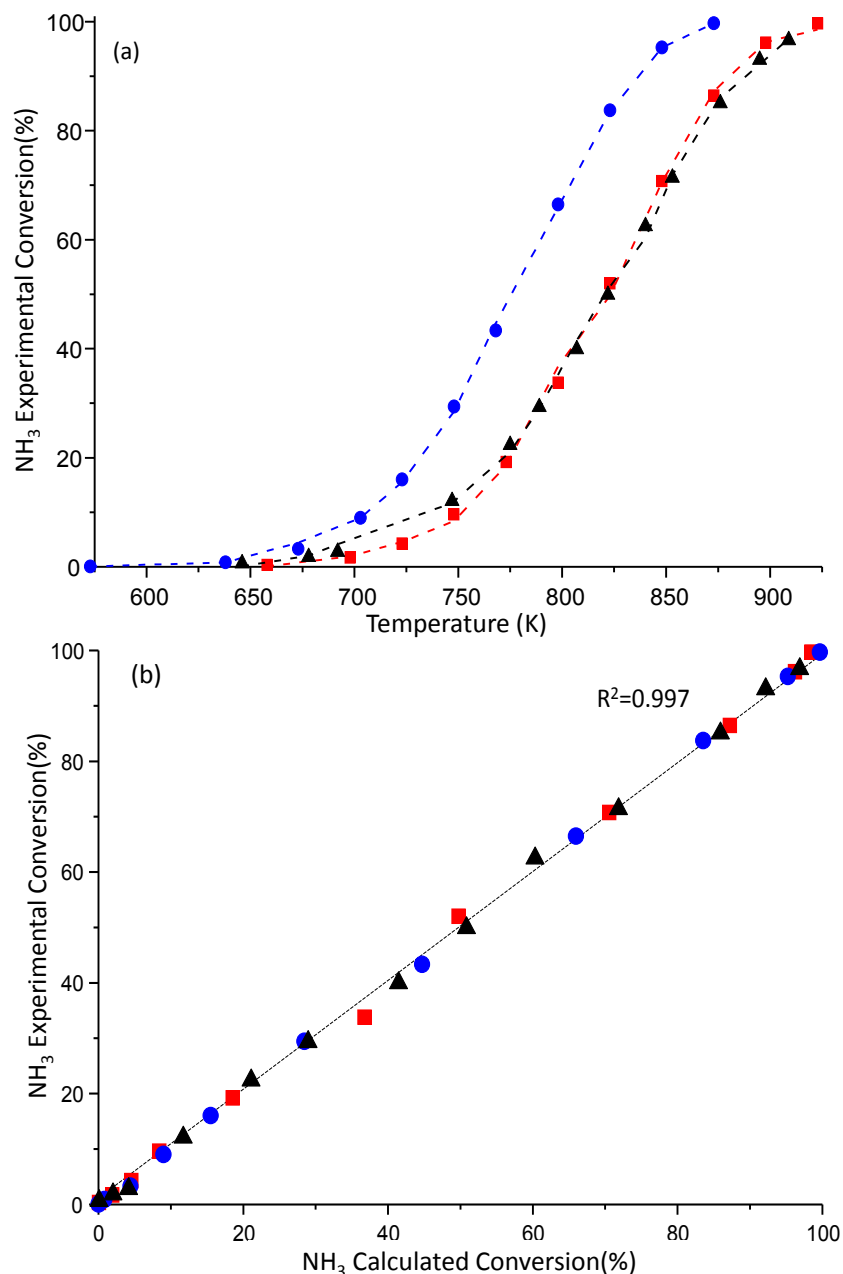
287 The kinetic parameters were calculated by non-linear multivariable regression using  
 288 Levenberg–Marquardt algorithm coupled to a Runge–Kutta–Fehlberg routine, using the  
 289 steady-state ammonia mass balance in plug-flow reactor, [50]:

$$(-r_{NH_3}) = \frac{dX_{NH_3}}{d(W_{cat}/F_{NH_3}^0)} \quad (3)$$

290  $X_{NH_3}$  is the ammonia conversion,  $W_{cat} / F_{NH_3}^0$  is the spatial time ( $g_{cat} \text{ s/mol } NH_3$ ) and  
 291  $(-r_{NH_3})$  is given by the power-law rate expression (2). The numerical solution of eqn  
 292 (3), allows the calculation of the sum of the squared residuals (SSR), used as objective  
 293 function minimized by the Levenberg–Marquardt algorithm:

$$SSR = \min \sum_{i=1}^{i=n} (X_{NH_3}^{exp} - X_{NH_3}^{calc})^2 \quad (4)$$

295



296  
 297 **Fig. 6.** (a) Dependence of NH<sub>3</sub> conversion with reaction temperature ( symbols:  
 298 experimental data; dotted line: model simulation, eqn. 2) (●) Ru(NN)/Al/M; (■)  
 299 Ru(Cl)/Al/M; (▲) Ru(acac)/Al/M; (b) Parity plot of the NH<sub>3</sub> conversion data.

300  
 301 Fig.6 a-b, shows the results the experimental and model prediction of ammonia  
 302 conversion vs. temperature data; and their excellent correlation in the parity plot (Fig. 6,  
 303 b). As is in these figures the homoscedastic distribution of the errors along the entire  
 304 range of conversions, indicates that power law model is an excellent approximation for  
 305 the modelling of the experimental data and, according to the main hypothesis of the



306 Temkin-Pyzhev model, the rate determining step of the reaction mechanism is the  
 307 associative adsorption of N-adsorbed species.

308 In the Table 2 are shown the kinetic parameters of the Temkin-Pyzhev model. The low  
 309 values of the standard errors obtained in all the cases, indicate again the excellent fitting  
 310 achieved, and therefore the validity of the model. Thus, the valued of the apparent  
 311 activation energies presented in the Table 2 have the same tendency (i.e. Ru(NN)/Al/M  
 312 < Ru(Cl)/Al/M < Ru(acac)/Al/M) that the one obtaiend by direct aplication of the  
 313 Arrhenius equation to the low conversion data (Fig. 5). The good coincidence between  
 314 the results obtaiend using the diferential method of the low conversion data (Figure 5)  
 315 and the integral method of whole range of conversion data, eqn. 3 and Figure 6, if an  
 316 additional proof of the confidence of the aproach used here.

317 Table 2: Kinetic parameters of Temkin-Pyzhev model.

Precursor Parameter	Ru(NN)/Al <sub>2</sub> O <sub>3</sub> /M Value	Ru(acac)/Al <sub>2</sub> O <sub>3</sub> /M Value	Ru(Cl)/Al <sub>2</sub> O <sub>3</sub> /M Value
k <sup>(*)</sup>	6.99 x10 <sup>-04</sup> ±error estandar	1.94 x10 <sup>-04</sup> ±error estandar	1.04 x10 <sup>-02</sup> ± estandar
E(kJ/mol)	76.63 ±error estandar	154.81 ±error estandar	89.32 ±error estandar
α	0.72 ±error estandar	1.046 ±error estandar	0.781 ±error estandar
β	0.45 ±error estandar	-0.278 ±error estandar	0.439 ±error estandar
-(α/β)	-1.57 ±error estandar	3.75 ±error estandar	-1.78 ±error estandar

318 \* units: mol NH<sub>3</sub>·atm<sup>(α+β)</sup>/g<sub>cat</sub>·s  
 319

320 According to the “Temkin formalism”, the ratio α/β should be between 0.4 and 0.5,  
 321 however the result calculated by the fitting of the data on Figure 2 shows large  
 322 differences. In fact, the kinetic orders of ammonia and hydorgen show a great variability  
 323 each other, which could be associated to the different catalyst intrinsinc nature, mainly  
 324 the average and distribution of particle sizez, the Ru content, and even the shape of the  
 325 particles, [33,53,54].

326 The lower activation energy and higher conversion of catalyst Ru(NN)/Al/M, compared  
327 to those prepared from the other precursors, can be attributed to its larger mean particle  
328 size (3.5 nm) which entails higher density of B<sub>5</sub>-type sites. This behavior has been  
329 explained by compensatory effects [49,55], derived from the relationship between the  
330 activation energy of the rate-limiting step, and the stability of the reaction intermediates  
331 on the surface of the catalyst.

332 In spite of that the Temkin-Pyzhev model provide an reasonable acknowledge of the  
333 kinetic reaction mechanism, and a simple way to analyze the data, in some cases it is  
334 necessary to use a more robust model, which can predict the kinetic order evolutions  
335 during ammonia decomposition reaction. The kinetic orders and the apparent energies  
336 of activation shown in Table 2 are an average of all the individual values which could  
337 be calculated as the gas composition changes along the reactor length. In fact, we have  
338 previously demonstrated [50], that the variable kinetic orders and apparent activation  
339 energies frequently reported in the literature are consequence of misleading data  
340 analysis, and not a consequence of the any change in the controlling step of the reaction  
341 mechanism.

#### 342 **4. Conclusions**

343 The results of this work show that the Ru catalyst in the form of honeycomb monoliths  
344 is a robust an active catalyst for H<sub>2</sub> generation form ammonia decomposition, thus being  
345 an enabling technology for the use of ammonia as practical H<sub>2</sub> storage media.

346 All the Ru-based catalysts used exhibited higher conversion in the decomposition of  
347 pure ammonia than Ni-based one, despite of ca. 5-fold larger metal loading of the latter.

348 Among all the Ru-based catalysts, the highest conversion and lowest apparent activation  
349 energy correspond to that prepared using ruthenium-nitrosyl-nitrate as precursor. This  
350 catalyst exhibited the largest mean particle size and the highest ratio of Ru<sup>0</sup>/Ru<sub>(total)</sub> in

351 reduced state after the passivation step. Thus, the highest activity of Ru prepared from  
352 nitrosyl-nitrate could be explained by the higher proportion of B<sub>5</sub>-type sites exposed,  
353 which have been related to particles with the same order of size.

354 The Temkin-Pyzhev (Power-Law) kinetic model provide an reasonable acknowledge of  
355 the reaction mechanism, and a simple way to analyze the data, in the whole range of  
356 ammonai conversions. The values of the kinetic orders, and the apparent energies of  
357 activation shown in Table 2, are valid to discriminate the intrinsic differences observed  
358 between the catalysts, and to develop a complete model of the monolithic reactor.

359

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