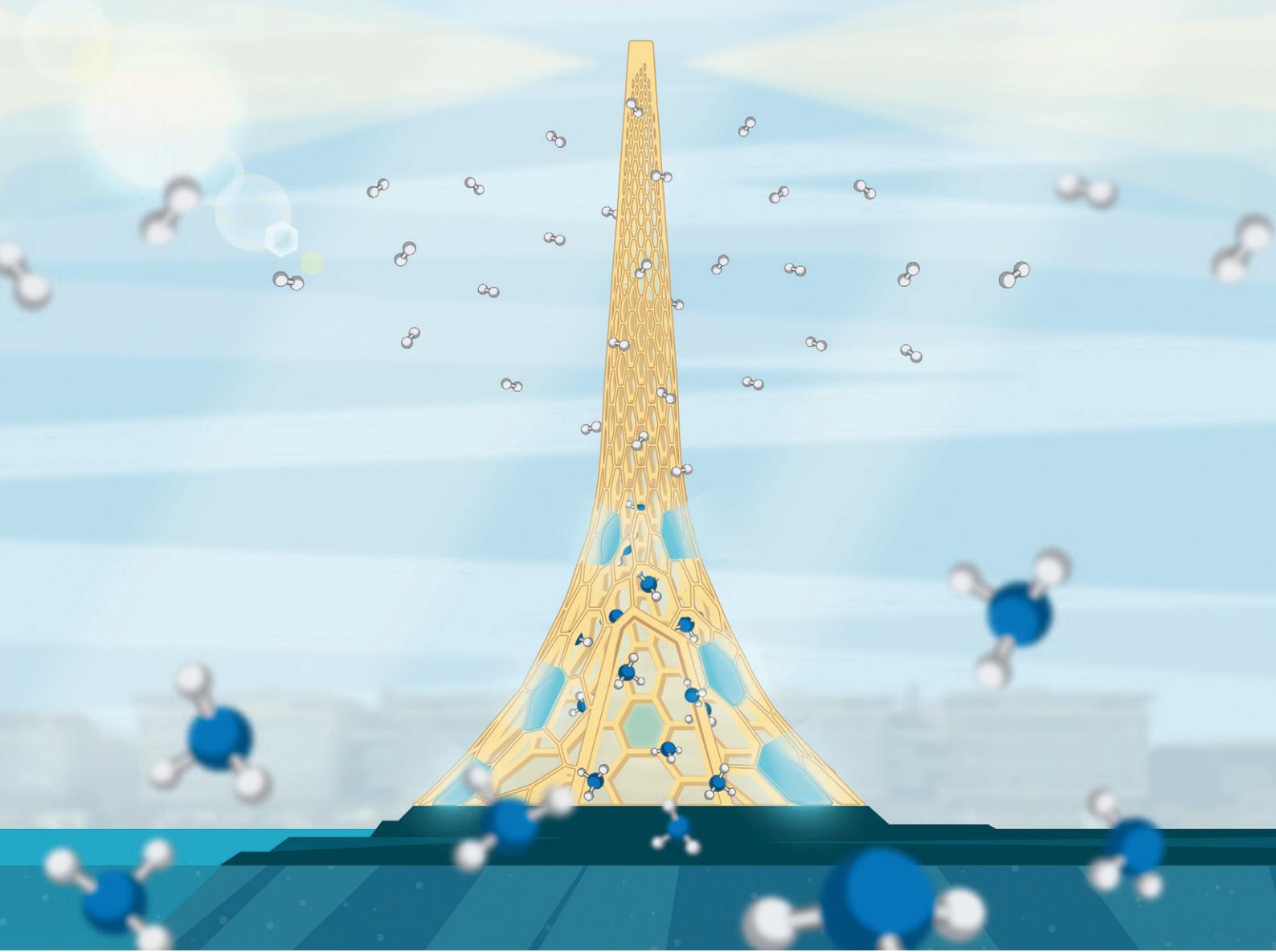


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Modeling-aided coupling of catalysts, conditions, membranes, and reactors for efficient hydrogen production from ammonia†

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The production of high-purity, pressurized hydrogen from ammonia decomposition in a membrane catalytic reactor is a feasible technology. However, because of the multiple coupled parameters involved in the design of this technology, there are extensive opportunities for its intensification. We investigated the coupling between the type of catalyst, process conditions, type of membrane, and reactor operation (isothermal and non-isothermal) in the catalytic decomposition of ammonia. First, we developed an agnostic dimensionless model and calculated the kinetic parameters for a set of lab-made Ru- and Co-based catalysts and the permeation parameters of a Pd-Au membrane. The non-isothermal model for the Pd-Au membrane reactor was validated with the experiments using Co-based catalysts. Finally, we analyzed the coupling conditions based on the model predictions, results obtained in the literature and our experimental results, including several case studies. The thorough analysis led us to identify optimized combinations of catalyst–conditions–membrane–reactor that yield similar or improved results compared to the ones of Ru-based catalyst in a non-membrane reactor. Our results indicate that optimizing a single factor, such as the catalyst, may not lead to the desired outcome and a more holistic approach is necessary to produce pressurized and pure hydrogen efficiently.

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1. Introduction

Hydrogen is assuming a leading role in the upcoming energy-fuel scenario owing to its emission-free combustion.¹ However, it suffers from inherent transportation and storage challenges that limit the potential benefits of its widespread, large-scale implementation as a fuel.² Hydrogen vectors or carriers do not require the transport or storage of large quantities of on-site hydrogen.³ These hydrogen-containing vectors can be organic or inorganic molecules that can be easily converted to pure hydrogen with ease and exhibit a minimal carbon footprint.⁴ Ammonia stands out compared to multiple hydrogen vectors primarily owing to its high

volumetric-energetic density, low liquefaction pressure of *ca.* 8 bar, and a global production and distribution network.^{5–7}

Ammonia decomposition is an endothermic reaction that is thermodynamically favored at low pressures.⁸ However, pressurized hydrogen production is desirable to decrease costs and derived emissions of its post-production compression for its storage and transportation⁹ (*ca.* 6.0 kW h kg^{−1} for compression of H₂ to 70 MPa, which leads to approximately 1.3 kg of CO₂ kg^{−1} of H₂).¹⁰ The energy analysis in our previous work¹¹ allowed us to prove that the efficiency of the whole process (including the post-compression stage up to 350 bar) would drop from 77.7% to ~68–73%, when hydrogen in the permeate is produced at 1–20 bar, respectively, thus highlighting the significance of developing reliable kinetic models at high pressure.

Many active phase-support combinations have been studied in the literature with Ru-based catalysts as the benchmark with remarkable activities at low temperatures.^{12,13} In addition, multiple non-noble metals, such as Fe, Co, and Ni, have been tested for this reaction,^{14–17} albeit displaying lower activities at low temperatures in contrast to Ru-based catalysts. In addition to the development of new catalysts, the use of a catalytic packed bed membrane reactor (denoted hereafter as a membrane

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reactor) has been proposed as an intensification route facilitating the production of high-yield and high-purity hydrogen.^{18–20} For a given catalyst, the comparison of the membrane reactor against the catalytic packed bed reactor (denoted hereafter as a non-membrane reactor) typically leads to improvements in the conversion and a relatively pure hydrogen stream.²¹

The modeling of membrane reactors is tackled by solving fundamental equations of mass balance, transport phenomena, and chemical kinetics in a standard shell-tube geometry.^{22–24} Hydrogen permeation *via* the membrane, driven by the pressure difference between the retentate and permeate sides, has been represented either as a linear pressure difference dependence²¹ or as the Sieverts–Fick law considering the permeation to be proportional to the difference of the square root of retentate and permeate pressures.²⁵ Moreover, hydrogen permeance through the membrane is considered to be temperature-dependent and thereby expressed in the form of Arrhenius law.^{26–28} Pd-based materials have been extensively studied as hydrogen selective layers,^{29–40} although other additional novel materials have been reported, some of which have remarkable hydrogen permeances and selectivities.^{41–57} Most modeling studies are performed for the benchmark Ru-based catalysts, and the kinetics are often represented with the Temkin–Pyzhev model or modifications thereof on account of its simplicity or to express the decomposition kinetics mathematically, thus resulting in an expression with a positive order ammonia and a negative order hydrogen dependence of varying magnitudes.^{58–66} The hydrogen reaction order significantly varies from catalyst to catalyst in the range of -2.5 to -0.25 ; it is commonly regarded as negative. Thus, maintaining a low partial pressure of hydrogen is beneficial for reaction kinetics in the membrane reactor.

Two dimensionless numbers are used to identify the underlying governing mechanisms in the operation of membrane reactors, namely, the Damköhler (Da) and Peclet (Pe) numbers. Using these numbers allows the operation to be separated into several zones, each of which is defined by the associated limiting factors: kinetics, convection, and permeation. The relatively high endothermicity of ammonia decomposition poses challenges to the isothermal operation, particularly for larger reactors.^{22,67} In this non-isothermal scenario, the Stanton (St) number is employed to create a dimensionless assessment of the energy balance⁶⁸ and measures how far the reactor operates from the isothermal condition.

Gómez-García *et al.*⁶⁹ modeled an Fe-catalyzed membrane reactor (high-pressure retentate stream, atmospheric pressure on the permeate side, and with a sweep-gas) and identified four operation zones that are either kinetically or permeation controlled. Furthermore, they compared the relative improvements to equilibrium conversions under various operating conditions. Li *et al.*²¹ performed ammonia decomposition in a membrane reactor using a sweep gas through a concerted experimental and dimensionless

simulation approach. Using Da and the permeation number (the equivalent of Pe), they demonstrated how hydrogen yields and purity could be tuned for silica-based and Pd-based membranes. The former includes permeation selectivities $\alpha_{\text{NH}_3/\text{H}_2}$ and $\alpha_{\text{N}_2/\text{H}_2}$, accounting for the slip of NH_3 and N_2 across the membrane, thereby reducing the purity of separated H_2 for membranes with low selectivity.

Our study focuses on developing an agnostic model in terms of the catalyst, process conditions (temperatures between 350 and 600 °C and pressures from 1 bar to 20 bar), membrane properties, and reactor operation (membrane and non-membrane, or isothermal and non-isothermal) for identify coupling and intensification possibilities between these parameters for the decomposition of ammonia in a catalytic membrane reactor. To validate the developed model, we performed a number of experimental tests in (i) an isothermal catalytic packed-bed reactor using Ru–K/CaO and Co–Ba/CeO₂ catalysts, (ii) a Pd–Au/Al₂O₃ membrane module without a catalyst, and (iii) a non-isothermal catalytic packed-bed membrane reactor with the same Co-based catalyst and membrane. This builds on our recently published promising results¹¹ and expands to other experimental conditions, catalysts, and non-isothermal reactors. The data were used to estimate our lab-scale configuration's kinetic and permeation parameters and embed them within the model. Finally, we use the model and data published in the literature to establish the areas of optimization, coupling, and intensification. Accordingly, we discuss how this holistic model can be used.

2. Methodology

2.1 Reactor model statement

We consider the geometry and conditions of a shell-tube catalytic packed bed membrane reactor (Fig. 1) to derive the numerical model. The porous tube inside is covered by a selective layer allowing H_2 permeation. The catalyst is placed in the shell side, thus forming a packed bed in the annular

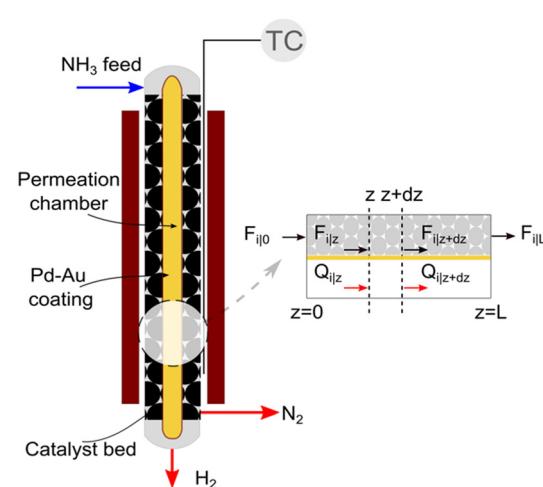


Fig. 1 Schematic and sectional view of the catalytic packed bed membrane reactor used for ammonia decomposition.



space. The feed enters at the top of the reactor, and the generated H₂ permeates through the selective layer towards its center across the reactor length. Assuming a co-current operation of the reactor, permeated H₂ exits at the bottom. The remaining gases (N₂ and unreacted NH₃, if any) flow down the catalytic bed (retentate side).

A pseudo-homogeneous one-dimensional mathematical model is chosen to describe the system, assuming that axial diffusion of mass and heat and radial concentration gradients on both sides are negligible. Based on the experimental observations, the kinetic control regime was proven due to the absence of external and internal mass and heat transfer limitations for all simulated operating conditions. The procedure used to prove that all experiments were obtained under kinetic control is described in the ESI† document of our previous work.⁷⁰ Other assumptions of the model are: (i) steady-state operation, (ii) plug flow in both feed and permeate streams, (iii) co-current operation, (iv) no concentration polarization effect through the membrane, and (v) isothermal operation on the permeate side. Under these assumptions, the following material balance equations are obtained in the axial (z) direction for the three species involved:

$$\frac{dF_i}{dz} = v_i r_{\text{rxn}} \left(\frac{A_c W_{\text{cat}}}{V} \right) - \frac{J_i A_m}{L} ((P x_i)^n - (P_p y_i)^n) \quad (1)$$

$$\frac{dQ_i}{dz} = \frac{J_i A_m}{L} ((P x_i)^n - (P_p y_i)^n) \quad (2)$$

where F_i and Q_i are the molar flow rate of component i in the catalytic bed and permeate side, respectively. Their corresponding mole fractions are designated as x_i and y_i . z is the axial direction along the reactor, A_c and V are the cross-sectional areas and total volume of the bed, respectively, W_{cat} is the total catalyst mass in the bed, A_m/L is the membrane area per unit length of the reactor, P and P_p are the catalytic bed and permeate side pressures respectively, and r_{rxn} is the ammonia decomposition reaction rate. For NH₃ and N₂, the permeation order n is set to one, while for H₂ $0.5 < n < 1$ depending on the phenomenon governing the permeation of H₂ across the membrane. J_i is the permeance of component i , and its temperature dependence is expressed using the Arrhenius expression.

$$J_i = J_{i,0} \exp \left(-\frac{E_{a,J}}{RT} \right) \quad (3)$$

Both energy and momentum balances corresponding to the catalytic bed section are similar derived based on a one-dimensional steady-state plug flow model. Eqn (4) is the energy balance of the catalyst bed:

$$\frac{dT}{dz} = \frac{1}{\sum_{i=1}^{N_s} (F_i C_{p,i})} \left(v_i r_{\text{rxn}} \left(\frac{A_c W_{\text{cat}}}{V} \right) (-\Delta H_{\text{rxn}}) + \frac{A_c A_s U}{V} (T_w - T) \right) \quad (4)$$

where T is the temperature of the catalytic bed, $C_{p,i}$ is the specific heat of component i , A_s is the reactor-oven contact

surface, U is the overall heat transfer coefficient between the catalytic bed and the reactor wall, and T_w is the reactor wall temperature. The total momentum balance to the catalytic bed section is as follows:

$$\frac{dP}{dz} = -\frac{G}{\rho d_p} \frac{(1-\phi)}{\phi^3} \left(\frac{150(1-\phi)\mu}{d_p} + 1.75G \right) \quad (5)$$

where G is the superficial mass velocity of the gas, ϕ is the catalytic bed porosity, d_p is the average catalyst particle diameter, μ is the viscosity, and ρ is the density of the gas. Catalytic bed porosity is assumed to be 0.55 and the particles are assumed to be well-rounded spheres with a diameter of 500 µm.

2.2 Kinetics of ammonia decomposition

In the literature, the Temkin–Pyzhev equation (eqn (6)) has been used to describe the ammonia decomposition reaction rate.^{58–65} This rate law represents a modification of a power law considering the effects of proximity to the equilibrium where nitrogen desorption is assumed as the rate-determining step. Note that the temperature dependence is expressed via the two-term Arrhenius equation.

$$r_{\text{rxn}} = k_0 \exp \left(-\frac{E_a}{RT} \right) \left(\left(\frac{p_{\text{NH}_3}}{p_{\text{H}_2}} \right)^2 - \frac{p_{\text{N}_2}}{K_{\text{eq}}^2} \left(\frac{p_{\text{H}_2}}{p_{\text{NH}_3}} \right)^{1-\beta} \right) \quad (6)$$

where k_0 is the pre-exponential factor, E_a is the activation energy, R is the universal gas constant, p_i is the partial pressure of component i , K_{eq} is the thermodynamic equilibrium constant, and β is the Temkin–Pyzhev empirical constant that correlates NH₃ and H₂ orders of reaction with their stoichiometric coefficients. Other authors, including this work, successfully fitted both orders of the reaction independently (eqn (7)) and reported them over a wide range of pressures and temperatures. This is the case of Ru-based⁷⁰ and Co-based catalysts.^{71,72}

$$r_{\text{rxn}} = k_0 \exp \left(-\frac{E_a}{RT} \right) p_{\text{NH}_3}^a p_{\text{H}_2}^b \left(1 - \frac{1}{K_{\text{eq}}^2} \left(\frac{p_{\text{N}_2} p_{\text{H}_2}^3}{p_{\text{NH}_3}^2} \right) \right) \quad (7)$$

where a and b are the reaction orders of NH₃ and H₂, respectively. The equilibrium constant is computed from eqn (8), and $\Delta G_{\text{rxn}}^{\circ}$ is estimated with the correlation in eqn (9), valid for the 673–1273 K range.⁷³

$$K_{\text{eq}} = \prod_{i=1}^{N_s} (p_i/P^{\circ})^{v_i} = \exp \left(-\frac{\Delta G_{\text{rxn}}^{\circ}}{RT} \right) \quad (8)$$

$$\Delta G_{\text{rxn}}^{\circ} = 95\,117 - 193.67T - 0.035293T^2 + 9.22 \times 10^{-6}T^3 \quad (9)$$

The activity of the catalyst is assumed to be constant due to lack of deactivation proven in our long-term stability tests. No deactivation for at least 100 h was observed for the Ru-based catalyst.⁷⁴ As for the Co-based catalyst stability tests were carried out in the packed bed membrane reactor for over 600 h with no evidence of detrimental effects on ammonia conversion, hydrogen recovery, composition, or flow rates of the permeate and retentate.¹¹



Table 1 Dimensionless variables, numbers, and parameters used in the model development

Dimensionless variables	Dimensionless numbers	Dimensionless parameters
$f_i = \frac{F_i}{F_{\text{NH}_3,0}}$ (10)	$\text{Da} = \frac{k_0 \exp(-\frac{E_a}{RT}) P_f^{(a+b)} W_{\text{cat}}}{F_{\text{NH}_3,0}}$ (15)	$\alpha_{\text{H}_2/i} = \frac{J_{\text{H}_2}}{J_i}$ (19)
$q_i = \frac{Q_i}{F_{\text{NH}_3,0}}$ (11)	$\text{Da}_{\text{III}} = \text{Da} \frac{-\Delta H_{\text{rxn}}}{C_{\text{pNH}_3,0} T_0}$ (16)	$P_r = \frac{P_f}{P_p}$ (20)
$\hat{P} = \frac{P}{P_f}$ (12)	$\text{Pe} = \frac{F_{\text{NH}_3,0}}{J_{\text{H}_2} A_m P_f^n}$ (17)	$\hat{C}_{\text{p},i} = \frac{C_{\text{p}i}}{C_{\text{pNH}_3,0}}$ (21)
$\theta = \frac{T}{T_0}$ (13)	$\text{St} = \frac{U A_s T_w}{C_{\text{pNH}_3,0} F_{\text{NH}_3,0} T_0}$ (18)	
$\zeta = \frac{z}{L}$ (14)		

Table 2 Dimensionless mass, energy, and pressure drop equations

Dimensionless equations
Mass balances
$\frac{df_i}{d\zeta} = v_i \text{Dax}_{\text{NH}_3}^a x_{\text{H}_2}^b \left(1 - \frac{P_f}{K_{\text{eq}}} \left(\frac{x_{\text{N}_2} x_{\text{H}_2}^3}{x_{\text{NH}_3}^2} \right) \right) - \frac{1}{\text{Pe}} \frac{(x_i^n - y_i^n / P_r^n)}{\alpha_{\text{H}_2/i}}$ (22)
$\frac{dq_i}{d\zeta} = \frac{1}{\text{Pe}} \frac{(x_i^n - y_i^n / P_r^n)}{\alpha_{\text{H}_2/i}}$ (23)
Energy balance
$\frac{d\theta}{d\zeta} = \frac{\text{Da}_{\text{III}} x_{\text{NH}_3}^a x_{\text{H}_2}^b \left(1 - \frac{P_f}{K_{\text{eq}}} \left(\frac{x_{\text{N}_2} x_{\text{H}_2}^3}{x_{\text{NH}_3}^2} \right) \right) + \text{St} (1 - \theta T_0 / T_w)}{\sum_{i=1}^{N_s} (f_i \hat{C}_{\text{p},i})}$ (24)
Momentum balance
$\frac{dP}{d\zeta} = \frac{LG(1-\phi)^2}{P_f \phi^2 d_p^2} [150 + 1.75 \text{Re}]$ (25)
Initial conditions
Mass balances
$f_i _{\zeta=0} = \frac{F_{i,0}}{F_{\text{NH}_3,0}}$ (26)
$q_i _{\zeta=0} = 0$ (27)
Energy balance
$\theta _{\zeta=0} = 1$ (28)
Momentum balance
$\hat{P} _{\zeta=0} = 1$ (29)



2.3 Dimensionless model

To evaluate the reactor performance in a wide range of operational conditions unconstrained by the reactor dimensions, a non-dimensional analysis of the mass, energy, and momentum balances was performed. This dimensional model is used to assess the system behavior *via* dimensionless variables and numbers summarized in Table 1.

Four dimensionless numbers are defined: the (i) Damköhler number (Da), comparing the rate of reaction with the rate of convective mass transfer. Da can be increased by increasing the space-time *via* either a high catalyst load or a low NH_3 feed flow; (ii) third Damköhler number (Da_{III}), defined as the ratio of the heat absorbed in the reaction to the convective rate of heat supplied by the feed; (iii) Peclet number (Pe), indicating the correlation between the convective and diffusive mass transports in the catalytic bed. A low Pe represents faster H_2 permeation across the membrane, achieved by decreasing NH_3 molar flow or increasing the driving force or the membrane area; and (iv) Stanton number (St) defined as the ratio between the heat externally transferred to the gaseous reaction medium and the energy content of the reaction medium. Higher St values indicate closeness to the isothermal operation. Dimensionless forms of eqn (1)–(5) are obtained incorporating eqn (10)–(21).

Equations reported in Table 2 were simultaneously solved using inbuilt MATLAB solvers. Stiffness while integrating the initial reaction rates at small lengths arises from the negative H_2 reaction order (absent in the feed). Therefore, an infinitesimal concentration of H_2 ($ca. 10^{-16}$) was defined in the feed for the simulation studies.²¹ The same approach was followed in determining the initial conditions for the permeate side wherein a small amount of an external inert component was assumed. Fractional NH_3 conversion and H_2 recovery are defined as follows:

$$X_{NH_3} = \frac{(f_{NH_3,0} - f_{NH_3,L} - q_{NH_3,L})}{f_{NH_3,0}} \quad (30)$$

$$R_{H_2} = \frac{q_{H_2,L}}{(f_{H_2,L} + q_{H_2,L})} \quad (31)$$

2.4 Experiments and fitting

In our previous studies, we investigated the ammonia decomposition reaction with two catalysts, Ru–K/CaO and Co–Ba/CeO₂.^{11,70,74} The Co-based catalyst with 80/20 molar Co/Ce ratio was synthesized by the co-precipitation method, followed by the addition of Ba (0.5 wt% Ba) as a promoter using the incipient wetness impregnation method. Ru–K/CaO was prepared using the impregnation method, followed by a pyrolysis step and final incorporation of potassium using the same method (10 wt% K). More details about synthesis, optimization and characterization of both catalysts can be found in our previous studies.^{70,74} Herein, we expanded the

experimental database of both catalysts with additional experiments classified into three sets:

i. In an isothermal catalytic packed-bed reactor (Fig. S1 and S2†) using Ru–K/CaO or Co–Ba/CeO₂ catalysts, to obtain a dataset of 95 experimental data points for each catalyst under the following conditions: temperatures from 250 to 550 °C, pressures from 1 to 40 bar, space times from 0.2 to 6 g_{cat} h mol⁻¹, partial pressures of NH_3 from 0.05 to 1 bar, and partial pressures of H_2 from 0 to 0.75 bar. The aim is to use the obtained X_{NH_3} to estimate k_0 , E_a , a , and b for each catalyst.

ii. In an isothermal membrane reactor module (Fig. S3†) using a Pd–Au/Al₂O₃ membrane, without a catalyst, to obtain a dataset of 55 experimental data points for different operating temperatures (350–500 °C), inlet flow rates (100–4000 N mL min⁻¹), and feed pressures (4–9 bar). The aim is to use R_{H_2} to estimate J_0 and $E_{a,J}$.

iii. In a non-isothermal catalytic packed-bed membrane reactor (Fig. S4†) using the Co–Ba/CeO₂ catalyst, to obtain a dataset of 56 experimental data points under the following conditions: temperatures from 250 to 500 °C, space times from 2.6 to 40 g_{cat} h mol⁻¹, and feed pressures from 4 to 16 bar.

Parameters estimated with sets (i) and (ii) were incorporated in the mass balances (eqn (22) and (23)) and solved together with the energy balance to match experimental X_{NH_3} and R_{H_2} , thus estimating the empirical heat transfer coefficient U of the lab-scale setup (set (iii)). T_w measurements were obtained at five points along the reactor for each experiment (Fig. S5†). A parabolic correlation (eqn (32)) was reported to determine the temperature profile along the reactor wall, in addition to a correlation (eqn (33)) to predict the inlet reactor temperature from the setpoint value ($T_{w,sp}$). Hence, unless the T_0/T_w ratio is specified, eqn (32) and (33) determine T_0 and T_w from $T_{w,sp}$.

$$T_w = -100.57z^2 + 95.77z + T_{w,sp} - 22 \quad (32)$$

$$T_0 = 0.9T_{w,sp} + 28 \quad (33)$$

where T_w , $T_{w,sp}$, and T_0 are the reactor wall temperature, reactor wall setpoint temperature, and feed temperature at the reactor inlet, expressed in K.

Insights into the characteristics of experimental reactors used for new data acquisition of each set are available in the ESI.† Parameter estimation for each fitting set was achieved with the *fminsearch* function of MATLAB, employed to determine the minimum of unconstrained multivariable systems using a derivative-free method. The sum of the squared residuals (SSR) of the experimental data and predicted values (*i.e.*, target variables like X_{NH_3} and R_{H_2}) was set as the objective function to minimize (eqn (34)). The 95% confidence intervals for nonlinear least squares parameter estimates were reported using the *nlpaci* MATLAB function given the residuals and the Jacobian matrix at the solution. Error variance was computed from the SSR as per eqn (35).



$$\text{SSR} = \sum_{i=1}^N (\chi^{\text{exp}} - \chi^{\text{cal}})^2 \quad (34)$$

$$\sigma = \sqrt{\frac{\text{SSR}}{N-2}} \quad (35)$$

3. Results and discussion

3.1 Agnostic model results

The model developed up to section 2.3, which is agnostic in terms of catalyst, conditions, and membrane, is used to identify the regimes of potential coupling possibilities based on the understanding of which phenomena governs different operation regions. For this purpose, we investigate the influence of dimensionless numbers (Da, Pe, St) on the process performance, determined by X_{NH_3} and R_{H_2} .

Fig. 2 shows contour plots of NH_3 conversion and H_2 recovery for a representative catalyst, $a = 0.5$ and $b = -0.75$, using different Da and Pe values assuming a highly selective membrane to H_2 ($\alpha_{\text{H}_2/\text{NH}_3} = 10^5$; $\alpha_{\text{H}_2/\text{N}_2} = 10^5$) operating at isothermal conditions. Fig. 2(a) can be divided into three regions: (i) the region comprising high Pe values ($\text{Pe} > 10^0$) where NH_3 conversion is independent of Pe, and the maximum conversion achieved is slightly $>90\%$; (ii) the region bounded by $10^{-2} < \text{Pe} < 10^0$ where the effect of the Pe is most noticeable and conversion of 100% is possible; and (iii) the region with $\text{Pe} < 10^{-2}$ where conversion is practically unaffected by the Pe and Da values. For all regions identified in Fig. 2, the Da value positively affects the NH_3 conversion. Based on its definition, the Da number (Table 1) can be

increased in many ways, *e.g.*, by increasing the catalyst activity (temperature, pressure, and type of catalyst) or amount of catalyst (or space-time). The primary difference between region (i) and the other two is that the convective flow is higher than the permeation term. This region is not affected by Pe numbers because of the high convective flow, poor performance of the membrane, or its absence, thus approaching the behavior of a conventional packed bed (non-membrane) reactor. Therefore, maximum conversion in this region is limited by thermodynamic equilibrium. In region (iii), the effect of Pe on NH_3 conversion is insignificant because of the limited membrane performance enhancement restricted by the maximum achievable driving force (*i.e.*, H_2 partial pressure in the catalytic bed) for permeation. In this region, the reactor performance is strictly controlled by reaction kinetics over permeation. Unlike other regions, in region (ii), conversion can be effectively tuned by modifying Da or Pe, the latter by increasing operating temperature, feed pressure, membrane area, or working with a more permeable material. For this study, this region represents an intensified area of interest. Therefore, these values of Pe and Da are used as a reference for upcoming simulations. The same regions are observed in Fig. 2(b). H_2 recovery is null for region (i) as it is only significant for cases where the H_2 diffusive flux is significantly higher than the convective one. As for region (ii), H_2 recovery is considerably influenced by a change in Pe, thus representing a regime controlled by the membrane characteristics. Region (iii) is affected only by the kinetics. These remarks agree with the previous observations.

For non-isothermal conditions, we examined the effects of different degrees of isothermal character on the reactor by assuming different values of St for $\text{Pe}_0 = 0.05$ (membrane reactor) and $\text{Pe}_0 = 500$ (non-membrane reactor), $T_w/T_0 = 1$, and activation energies of 100 and 25 kJ mol^{-1} for a representative catalyst and membrane, respectively. Because certain dimensionless numbers vary along the reactor (*e.g.*, Da with non-isothermal operation), the subscript 0 (*e.g.*, Da_0) is used to indicate its values at the reactor inlet. Fig. 3(a) shows that there is a maximum difference of 87% between the membrane reactor conversion at $\text{St} = 0$ (adiabatic) and $\text{St} = \infty$ (isothermal) at $\text{Da}_0 \approx 1$. This Da value is the minimum value required to achieve complete conversion in an isothermal membrane reactor. At extremely low Da values ($\text{Da}_0 < 10^{-2}$), conversions are limited, thus making the effect of St negligible. Fig. 3 shows that $\text{St} = 100$ is close to isothermal operation for the entire Da range. At a sufficiently high Da, the closer the operation is to an isothermal reaction, the larger the difference is between the conversion achieved in both reactors.

Furthermore, Fig. 3(a) shows that a membrane reactor not only can go beyond the intrinsic thermodynamic limitations of the reaction, but it also performs better than the non-membrane reactor under less isothermal scenarios (low-St operation). For values of $\text{St} < 100$, the Da requirements for achieving high NH_3 conversion exponentially increase. At $\text{Da}_0 < 0.1$, all curves demonstrate the same conversion for the

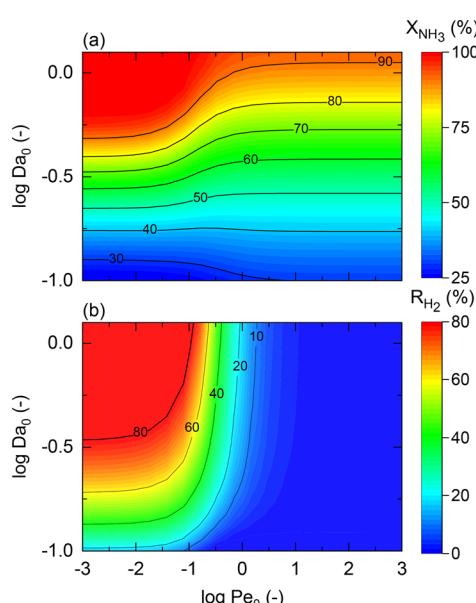


Fig. 2 Effect of Damköhler (Da) and Peclat (Pe) numbers on (a) NH_3 conversion and (b) H_2 recovery. Simulation conditions: $\text{St} = \infty$; $P_r = 4$; $P_p = 1$ bar; $T_w = 400$ $^{\circ}\text{C}$; $T_0/T_w = 1$; $\alpha_{\text{H}_2/\text{NH}_3} = 10^5$; $\alpha_{\text{H}_2/\text{N}_2} = 10^5$; $a = 0.5$; $b = -0.75$; $n = 0.5$. $X_{\text{eq}@400^{\circ}\text{C}} = 99.8\%$.



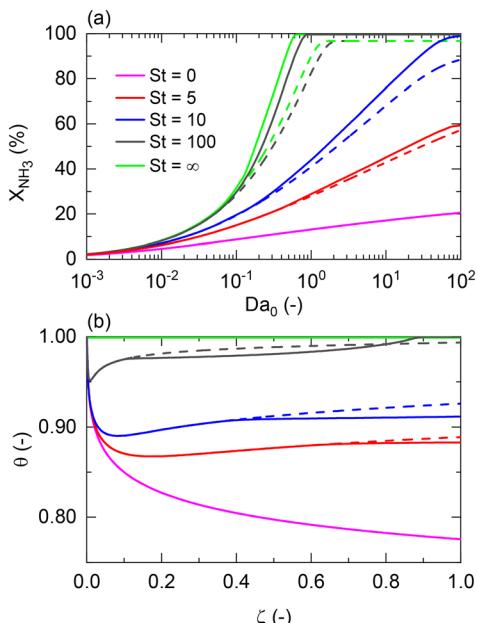


Fig. 3 St effect on (a) NH_3 conversion as a function of the Damköhler number at the reactor inlet (Da_0) and (b) dimensionless temperature profiles, θ , for a membrane reactor (solid lines) and a non-membrane reactor (dashed lines) across the dimensionless length. Simulation conditions: $Da_0 = 1$; $Pe_0 = 0.05$ (membrane reactor); $Pe_0 = 500$ (non-membrane reactor); $P_r = 4$; $P_p = 1$ bar; $T_w = 400$ °C; $T_0/T_w = 1$; $\alpha_{\text{H}_2/\text{NH}_3} = 10^5$; $\alpha_{\text{H}_2/\text{N}_2} = 10^5$; $a = 0.5$; $b = -0.75$; $n = 0.5$; $E_a = 100$ kJ mol⁻¹; $E_{a,J} = 25$ kJ mol⁻¹; $X_{\text{eq},400\text{ °C}} = 99.8\%$.

membrane reactor and the packed bed reactor. Fig. 3(b) shows dimensionless temperature profiles for $Da_0 = 1$ and shows the similarity between $St = 100$ and the isothermal operation, where the feed temperature can be restored after the initial drop characteristic of endothermicity in this reaction. However, the initial temperature drop at lower St is more significant and cannot be neglected. Higher temperature drops in the membrane reactor compared to the non-membrane reactor are explained by higher heat consumption of the reaction (higher conversion) and heat loss of hydrogen permeating to the other side of the membrane.

Returning to the isothermal membrane reactor conditions, pressurized H_2 production is explored by comparing the performance of a membrane reactor at different permeate pressures as a function of the feed and pressure ratio (Fig. 4). Fig. 4(a) shows the expected adverse effect of pressure for both the non-membrane reactor (dashed line) and equilibrium (dotted line) conversions. Feed pressure negatively affects the thermodynamic equilibrium and kinetics. These effects are superposed by the positive impact on H_2 permeation by increasing the driving force across the membrane, enhancing the net ammonia decomposition rate. However, there is a tradeoff between the desired permeate pressure and feed pressure required to obtain complete NH_3 conversion (Fig. 4(a)). For example, a 3 bar feed pressure is required to completely

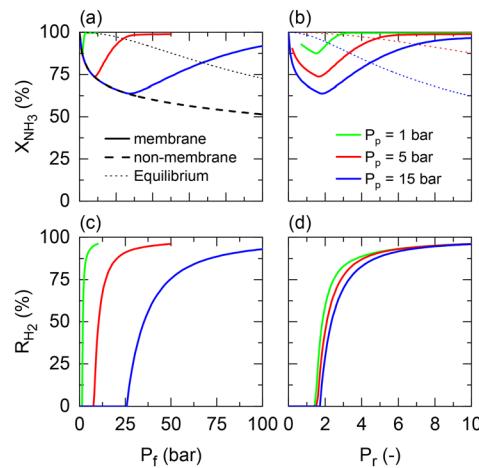


Fig. 4 Permeate pressure effect on NH_3 conversion for different (a) feed pressures (P_f) and (b) pressure ratios (P_r); H_2 recovery for different (c) feed pressures (P_f) and (d) pressure ratios (P_r). Simulation conditions: $Da/P_f^{(a+b)} = 1$; $Pe \times P_f^n = 0.05$; $St = \infty$; $T_w = 400$ °C; $T_0/T_w = 1$; $\alpha_{\text{H}_2/\text{NH}_3} = 10^5$; $\alpha_{\text{H}_2/\text{N}_2} = 10^5$; $a = 0.5$; $b = -0.75$.

convert NH_3 to yield pure H_2 at 1 bar on the permeate side, whereas if an H_2 product pressure of 15 bar is desired, all of the feedstock cannot be converted even at 100 bar.

When representing the former in terms of the pressure ratio rather than feed pressure (Fig. 4(b)), we observe that the pressure ratio required to reach 100% conversion becomes increasingly higher with the permeate pressure. Therefore, as per the model, high-pressure H_2 production with complete feedstock conversion becomes asymptotically more pressure demanding as the desired product pressure increases up to the point where 15 bar H_2 production would require a feed pressure of >150 bar. However, this seems not to be the case with the pressure ratio (ca. $P_r = 1.5\text{--}2$) required to overcome the non-membrane reactor performance, which is approximately $P_r = 2$ irrespective of the permeate pressure, and therefore setting the minimum pressure ratio for a membrane reactor. Fig. 4(c) and (d) show the same effect on H_2 recovery. The similarity between the impact of different permeate pressures on H_2 recovery determines that the minimum pressure ratio to achieve maximum recovery is $\sim P_r = 6$, virtually independent of permeate pressure. These observations agree with the experimental results reported in our previous study.¹¹ The potential areas for the performance improvement are summarized in Table 3. Note that the values identified for Da correspond to the entire evaluated range in Fig. 2, indicating that tuning Da allows achieving any target conversion for a specific membrane reactor configuration.

In all previous simulations, the dimensionless form of the momentum balance (eqn (25)) was used to compute the pressure drop along the catalytic bed. None of the simulated conditions exhibited significant pressure drop values (<1%). Hence, the pressure drop can be considered negligible for all operating Da , Pe , and St ranges addressed in this study.

Table 3 Identified ranges of interest for different dimensionless numbers and studied parameters

Dimensionless number/parameter	Criterion	Value
Pe	Potential area for coupling	$10^{-2} < Pe < 10^0$
Da	Tune reactor performance under specific conditions	$10^{-1} < Da < 10^0$
St	Avoid severe axial temperature gradients	$St > 100$
P_r	Overcome non-membrane reactor performance	$P_r > 2$

Hence, additional references to the pressure drop are omitted when discussing these operating conditions. To support this observation, pressure drop estimations are shown in Fig. S6,† even for the most severe scenarios (*i.e.*, 1000 N mL min⁻¹ NH₃ in the feed).

3.2 Parameter estimation

Kinetic parameters of Ru-K/CaO and Co-Ba/CeO₂ catalysts (set i), Pd-Au/Al₂O₃ membrane parameters (set ii), and the membrane reactor heat transfer coefficient (set iii) were estimated based on the previously described extended experimental sets. Fig. 5 shows the parity plots comparing the experimental and calculated target variables for all experimental conditions and parameter sets: (a) the kinetic model for the Ru-K/CaO and Co-Ba/CeO₂ catalysts developed with experimental set (i), using NH₃ conversion as a fitting parameter; (b) the permeation model of the Pd-Au/Al₂O₃ membrane using experimental data from set (ii) with H₂ recovery as a fitting parameter; and (c) and (d) the non-isothermal membrane reactor model, experimental set (iii), using both NH₃ conversion and H₂ recovery, respectively. Because of the relatively good fitting of all models, we claim that the set of assumptions is adequate for this system, *i.e.*, eqn (7) is a reliable kinetic model for the ammonia decomposition kinetics in the evaluated ranges of operational

conditions for both catalysts and eqn (3) is adequate for modeling the H₂ permeation and the diabatic thermal regime is well represented by a curved wall temperature profile. The statistics for the estimated parameters in Table 4 likewise demonstrate the reliability of the fitting procedure.

Table 4 summarizes the estimated parameters for different experimental datasets. The modified Temkin-Pyzhev parameters obtained for Ru-K/CaO and Co-Ba/CeO₂ catalysts show substantial differences, with the former showing faster overall kinetics with larger NH₃ and smaller H₂ reaction orders. The lower E_a value estimated for the Co-Ba/CeO₂ catalyst indicates that the kinetic constant is less affected by the temperature than the Ru-K/CaO catalyst. However, the fact that the pre-exponential factor of Ru-K/CaO is one order of magnitude larger than that of Co-Ba/CeO₂ agrees with the *a priori* expected higher activity of the former. The following section provides a full assessment of the estimated reaction orders and activation energies in the context of the previously reported values.

As per the membrane parameters, the fitted $J_{H_2,0}$ and $E_{a,J}$ values lie within the range expected for H₂ selective Pd membranes, as shown in section 3.3. The estimated H₂ permeation order is in the lower limit of the acceptable values ($n = 0.5$). Because only H₂ was detected at the permeate side in

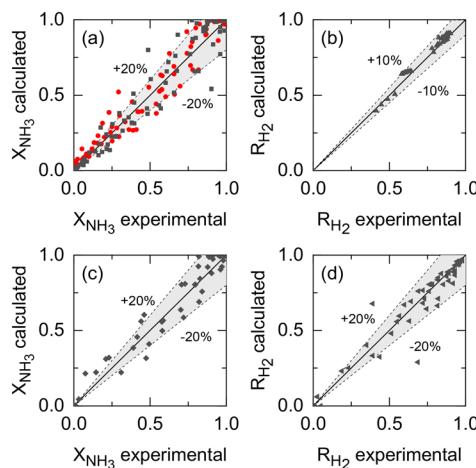


Fig. 5 Comparison of experimental values and simulation predictions after parameter estimation of (a) set (i) (Ru-K/CaO, black squares; Co-Ba/CeO₂, red circles); (b) set (ii); and (c) and (d) set (iii) in terms of NH₃ conversion and H₂ recovery, respectively.

Table 4 Estimated parameters for different experimental datasets

Parameter set i		
Catalyst	Ru-K/CaO	Co-Ba/CeO ₂
k_0 (mol g ⁻¹ h ⁻¹)	$(6.27 \pm 0.01) \times 10^{11}$	$(3.33 \pm 0.29) \times 10^{10}$
E_a (kJ mol ⁻¹)	166.4 ± 0.1	154.1 ± 3.2
a	0.47 ± 0.00	0.37 ± 0.02
b	-1.42 ± 0.00	-1.04 ± 0.02
r^2	0.968	0.984
σ^2	0.096	0.070
Parameter set ii		
J_0 (mol m ⁻² s ⁻¹ Pa ⁻ⁿ)		$(7.85 \pm 0.11) \times 10^{-5}$
$E_{a,J}$ (kJ mol ⁻¹)		6.7 ± 1.6
n		0.5
α_{H_2/N_2}		∞
α_{H_2/NH_3}		∞
r^2		0.991
σ^2		0.040
Parameter set iii		
U (W m ⁻² K ⁻¹)		244 ± 53
r^2		0.945
σ^2		0.057



permeation-exclusive experiments (set (ii)) and membrane reactor experiments (set (iii)), α_{H_2/N_2} and α_{H_2/NH_3} were set to ∞ . Furthermore, experimental Pe_0 (0.03–0.61) and Da_0 (0–20) are within the ranges defined in Table 3, indicating that the fitted parameters are suitable for the reactor–catalyst coupling study presented in the following sections. The value of the overall heat transfer coefficient at the wall, along with the dimensions of the reactor, corresponds to experimental values of $300 < St < 1600$, which based on our previous discussion, are sufficiently high values ($St > 100$) to rule out significant heat-related limitations. These high values of St arise not only from the estimated value of U , which is within the expected range based on the literature,^{75–77} but also due to the annular geometry of the reactor.

3.3 Effect of model parameters

Once the model is fitted to our experimental results, it can be used (along with other experimental results available in the literature) to confirm the realistic coupling regions of the membrane reactor. Considering the effect of membrane properties, in Fig. 6(a) and (b), the impact of H_2/NH_3 membrane selectivity is shown for a wide range of H_2 permeances, representing a region $5 \times 10^{-3} < Pe < 5 \times 10^3$.

Herein, the results show that NH_3 conversion for $J_{H_2} < 10^{-7} \text{ mol s}^{-1} \text{ m}^2 \text{ Pa}^{0.5}$ is <90% and independent of α_{H_2/NH_3} , which for the $F_{NH_3,0}/A_m$ ratio and selected P_r represents $Pe > 10^0$, indicating that the H_2 permeation flux is not sufficient to overcome non-membrane reactor performance. For higher J_{H_2} values and low H_2/NH_3 selectivity ($\alpha_{H_2/NH_3} < 10^2$), NH_3 conversion (based on eqn (30)) can be lower than that obtained in a non-membrane reactor because of NH_3 permeation. With the values of $J_{H_2} > 10^{-5} \text{ mol s}^{-1} \text{ m}^{-2} \text{ Pa}^{-0.5}$ and $\alpha_{H_2/NH_3} > 10^3$, the complete conversion of ammonia can be achieved in the reactor. Minimum α_{H_2/NH_3} requirements to achieve H_2 purity higher than 99.9% are very similar. For most lab-scale studies reported for ammonia decomposition in a Pd-coated membrane reactor, α_{H_2/NH_3} is not reported because no NH_3 is detected in the permeate side,^{22,24,25,78} indicating that although there is a minimum requirement of α_{H_2/NH_3} on the order of 10^3 , in most of the reported cases this value is easily achieved for common H_2 permselective materials. This is the case of the membrane studied in this work, for which pure H_2 is obtained in the permeate in all experiments.

Fig. 6(c) shows the effect of H_2/N_2 membrane selectivity on H_2 purity. The most significant difference between Fig. 6(b) and (c) is the area bounded by $J_{H_2} > 10^5 \text{ mol s}^{-1} \text{ m}^{-2}$

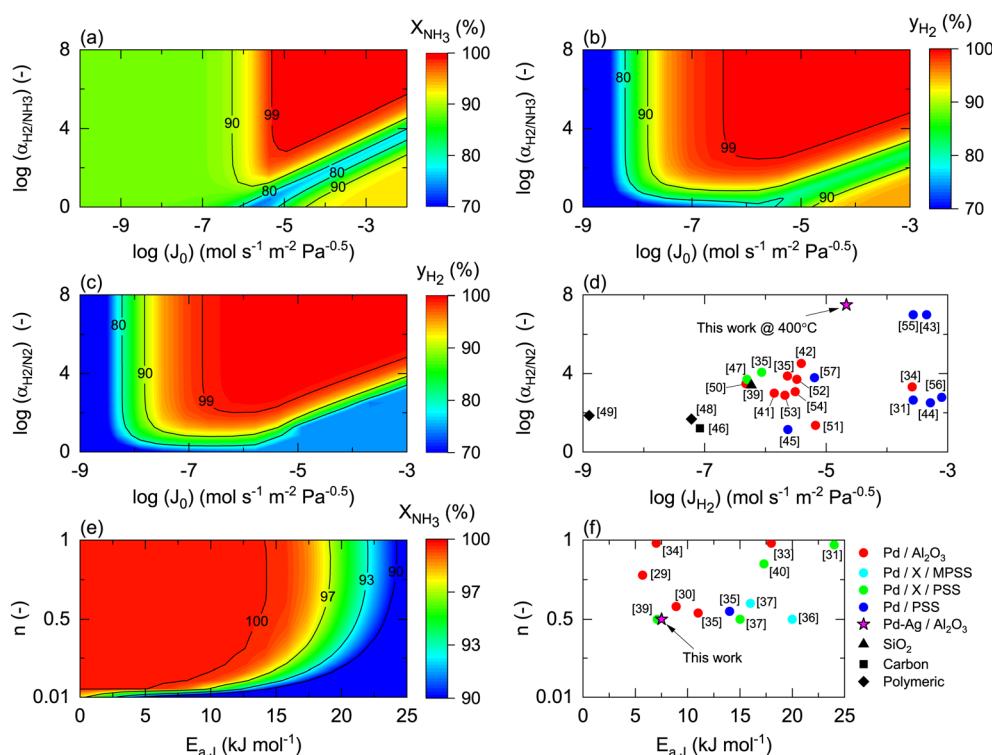


Fig. 6 Membrane properties' influence on membrane reactor performance, expressed as the effect of H_2/NH_3 selectivity (α_{H_2/NH_3}) and H_2 permeance (J_{H_2}) on (a) simulated NH_3 conversion and (b) simulated H_2 purity (y_{H_2}); H_2/N_2 selectivity (α_{H_2/N_2}) and H_2 permeance (J_{H_2}) on (c) simulated H_2 purity (y_{H_2}), and (d) their reported values in the literature; activation energy (E_a) and H_2 permeation order (n) on (e) simulated NH_3 conversion and (f) their reported values in the literature. Simulation conditions: $Da_0 = 1$; $F_{NH_3,0}/A_m = 0.1 \text{ mol m}^{-2} \text{ s}^{-1}$; $St = 100$; $P_r = 4$; $P_p = 1 \text{ bar}$; $T_w = 400 \text{ }^\circ\text{C}$; $T_0/T_w = 1$; $\alpha_{H_2/NH_3} = 10^5$; $\alpha_{H_2/N_2} = 10^5$; $a = 0.5$; $b = -0.75$; $n = 0.5$; $E_a = 100 \text{ kJ mol}^{-1}$; $E_{a,J} = 25 \text{ kJ mol}^{-1}$. For case (e), $J_{H_2,0} = 10^{-4} \text{ mol m}^{-2} \text{ s}^{-1} \text{ Pa}^{-1}$. $X_{eq@400^\circ\text{C}} = 99.8\%$.

$\text{Pa}^{-0.5}$ and $\alpha_{\text{H}_2/\text{N}_2} < 10^2$, where H_2/N_2 membrane selectivity has a more significant impact on H_2 purity. In this region, permeance for NH_3 and N_2 is the same; however, as NH_3 is converted along the axial position of the reactor, the partial pressure of the former decreases as opposed to the latter, directly affecting the permeation flux. Because of the N_2 flux increases with NH_3 conversion, H_2/N_2 membrane selectivity has a more significant effect on H_2 purity than $\alpha_{\text{H}_2/\text{NH}_3}$. Fig. 6(d) shows the reported values of J_{H_2} and $\alpha_{\text{H}_2/\text{N}_2}$ for H_2 selective materials tested in membranes. As per the model, most $\text{Pd}/\text{Al}_2\text{O}_3$ membranes are expected to yield a pure permeate. However, when using supports for Pd that endow extremely high H_2 permeation rates (e.g., PSS), the selectivity is compromised, and consequently so is H_2 purity. The latter highlights the benefits of selecting membranes like $\text{Pd}/\text{Al}_2\text{O}_3$ that exhibit an excellent $J_{\text{H}_2}-\alpha_{\text{H}_2/\text{N}_2}$ balance. The estimated value of J_{H_2} at 400 °C for the $\text{Pd}-\text{Au}/\text{Al}_2\text{O}_3$ membrane used in this study (Table 4) is coherent with the values obtained for most $\text{Pd}/\text{Al}_2\text{O}_3$ membranes and others that are exclusively selective to H_2 . Furthermore, considering values of $\alpha_{\text{H}_2/\text{NH}_3}$ and $\alpha_{\text{H}_2/\text{N}_2}$ on the order of 10³, for a J_{H_2} on the order of 10⁻⁵ as representative of an acceptable membrane performance for the simulation conditions, only Pd -based membranes meet the requirements. In the simulations, the effect of $\alpha_{\text{H}_2/\text{N}_2}$ on NH_3 conversion was reported to be negligible and is not shown in Fig. 6.

Fig. 6(e) and (f) show the effects of the membrane permeation order, activation energy, and experimental values. Several reported H_2 selective materials follow Sieverts–Fick's law ($n = 0.5$), indicating that the limiting step of H_2 transport across the membrane is the diffusion in the film. However, other studies report deviations from this law ($n > 0.5$), indicating that the limiting step is the dissociation in the film or the transport across the porous support.⁷⁹ Based on the experimental data gathered in Fig. 6(f), a direct correlation between the membrane support and value of n seems to be missing, highlighting that other factors in addition to the layer material and support (e.g., preparation method) certainly play a role. Under the simulated conditions, n does significantly contribute to the final performance. The activation energy has a more significant impact with a reduction of NH_3 conversion of 10% when $E_{\text{a},J}$ increases from 10 to 25 kJ mol⁻¹. However, the $E_{\text{a},J}$ value estimated in this work (6.7 kJ mol⁻¹) is similar to the $\text{Pd}/\text{Al}_2\text{O}_3$ membranes in Fig. 6(f), which indicates that their performance is not highly influenced by the temperature. When the pre-exponential factor for permeance is sufficiently high, low activation energy membranes are preferred over higher ones as cold spot formation cannot be avoided nearby the reactor inlet where the reaction rates are the highest.

To explore the effect of catalyst kinetics on both reactor performances, the effects of activation energy (Fig. 7) and reaction orders (Fig. 8) were analyzed. In this simulation, we assumed the same pre-exponential constant ($k_0 = 10^{11}$ mol h⁻¹ g_{cat}⁻¹) provided that a one-to-one comparison of the values reported in the literature is challenging because different

authors define the rate constant based on other terms. However, considering our results of Ru - and Co -based catalysts, there is one order of magnitude difference between them (Table 4). This indicates that the effect of the pre-exponential constant must not be decoupled from the analysis in Fig. 7. The theoretical impact of k_0 and E_{a} shown in Fig. S7† is as expected, namely, higher k_0 values allow complete conversions even if E_{a} is high. Despite the latter, the simulations and experimental results in Fig. 7 enable the comparison between different catalytic systems on the same basis and assess the impact of catalyst activity (E_{a}) on the reactor performance and wall temperature. In this manner, Fig. 7 shows that NH_3 conversion is less dependent on reactor wall temperature for a membrane reactor than for a non-membrane reactor. The activation energy band dividing the reactive zone from the inactive area for a non-membrane reactor ($125 < E_{\text{a}} < 175$ kJ mol⁻¹) is narrower and displaced to the right for the membrane reactor ($137 < E_{\text{a}} < 185$ kJ mol⁻¹), similar to what is observed in Fig. S7.† The simulations predict that for the assumed k_0 , $\text{Co}-\text{Ba}/\text{CeO}_2$ catalysts with $E_{\text{a}} = 154$ kJ mol⁻¹ (Table 4) achieve complete NH_3 conversion at $T_{\text{w}} > 420$ °C in a membrane reactor; however, conversion in a non-membrane reactor using the same catalyst will not achieve complete conversion even at $T_{\text{w}} = 500$ °C. The role of E_{a} is highlighted with Fe -based catalysts, known to require higher temperatures along with higher E_{a} values.

Reaction orders play a central role in the selection of coupling strategies. In Fig. 8, we explore the effect of parameters a and b on the performance of a non-

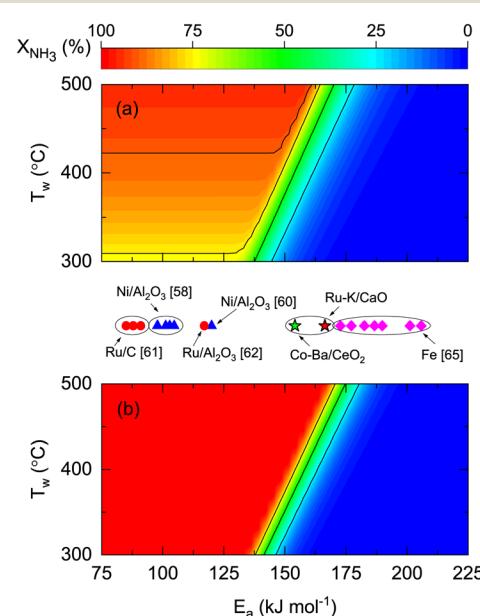


Fig. 7 Contour plots for NH_3 conversion for different reactor wall temperatures (T_{w}) and catalyst activation energies (E_{a}) for (a) non-membrane and (b) membrane reactors. Values reported in the literature are at scale. Simulation conditions: $\text{Pe}_0 = 0.05$ (membrane reactor); $\text{Pe}_0 = 500$ (non-membrane reactor); $W/F_0 = 22$ g_{cat} h mol⁻¹; $\text{St} = 100$; $P_r = 10$; $P_p = 1$ bar; $T_0/T_{\text{w}} = 1$; $\alpha_{\text{H}_2/\text{NH}_3} = 10^5$; $\alpha_{\text{H}_2/\text{N}_2} = 10^5$; $a = 0.5$; $b = -0.75$; $n = 0.5$; $k_0 = 10^{11}$ mol h⁻¹ g_{cat}⁻¹; $E_{\text{a},J} = 25$ kJ mol⁻¹.



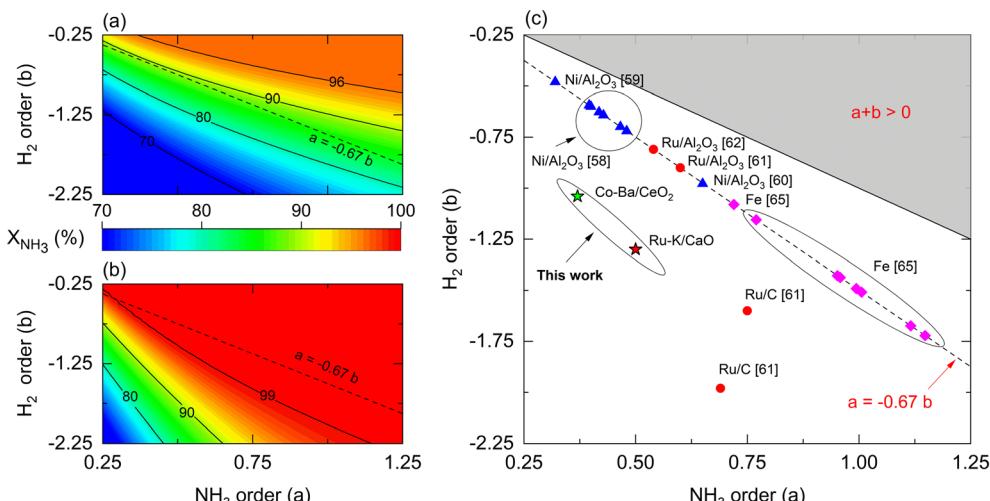


Fig. 8 Rate law parameters effect on simulated NH_3 conversion for (a) membrane reactor, (b) non-membrane reactor, and (c) their reported values for suitable catalysts in the literature. Simulation conditions: $D_{\text{eff}} = 1$; $P_{\text{O}_2} = 0.05$ (membrane reactor); $P_{\text{O}_2} = 500$ (non-membrane reactor); $\text{St} = 100$; $P_r = 4$; $P_{\text{p}} = 1$ bar; $T_w = 400$ °C; $T_0/T_w = 1$; $\alpha_{\text{H}_2/\text{NH}_3} = 10^5$; $\alpha_{\text{H}_2/\text{N}_2} = 10^5$; $n = 0.5$; $E_a = 100$ kJ mol $^{-1}$; $E_{\text{a,J}} = 25$ kJ mol $^{-1}$.

membrane and membrane reactor assuming $E_a = 100$ kJ mol $^{-1}$. Using a membrane reactor decreases the effect of b on NH_3 conversion, thus allowing catalysts with lower H_2 reaction orders to reach higher conversion. In Fig. 8(a), the slope of the contour lines is similar to the one formed by the reported parameters (Fig. 8(c)), indicating a correlation between the a/b ratios and performance. However, this effect changes in a membrane reactor where catalysts with the same a/b ratios belong to a broader spectrum of interpretations. These observations, in addition to the remarks made in Fig. 7, highlight the impact of kinetic parameters on membrane reactor performances, indicating that the intrinsic activity of the catalyst is less significant. For most catalysts shown in Fig. 8(c), β values in the Temkin–Pyzhev model (eqn (6)) are represented by the $a = -0.67 \cdot b$ line. Fe-based catalysts possess higher NH_3 and lower H_2 reaction orders than the Ni-based catalysts. The reaction orders previously estimated in Table 4 do not fall in the line, $a = -0.67 \cdot b$, as the parameters were obtained independently according to eqn (7). Furthermore, according to the values of a and b reported in the literature, the choice of a rate law that independently treats both orders yield lower values compared to the Temkin–Pyzhev law.

However, a and b must not be understood as sole indicators of intrinsic catalyst activity, as the estimation order affects the pre-exponential factor estimation. The latter is exemplified with Ru- and Co-based catalysts of this study, where Co-Ba/CeO₂ has higher reaction orders. Nevertheless, its estimated pre-exponential factor value is an order of magnitude lower than that of Ru-K/CaO.

3.4 Assessment of coupling opportunities

Previous studies discussed the benefits of using membrane reactors over packed bed reactors for ammonia decomposition due to the enhanced reaction rates and thermodynamics

resulting from the selective removal of H_2 from the catalytic bed.^{22,25,80} In this section, the developed and validated model is used as a tool to harness other less-common effects in non-isothermal membrane reactors. This approach enables defining high-performance opportunity windows for catalysts that are moderately active for ammonia decomposition. This is the case of the Co-Ba/CeO₂ catalyst in this study, compared to the benchmark Ru-K/CaO. To explore the potential applicability of the Co-Ba/CeO₂ catalyst, NH_3 conversions for four catalyst/reactor combinations (Table 5) are compared based on the catalysts and reactors used in the experimentation. This analysis is performed for specific operating conditions (temperature, space-time) based on the parameters estimated in section 2.4.

When comparing the performance of the catalyst/reactor couples listed in Table 5, the improvement in NH_3 conversion (ΔX_{NH_3}) is the largest when comparing case 3 vs. 1 in Fig. 9(a). This observation was not only expected due to the higher intrinsic activity (expressed as higher k_0) of the catalyst, but also for the buffering role of the membrane reactor on the H_2 reaction order in the rate law, in line with the conclusion drawn from Fig. 8. The same reasons apply to the enhancement behavior provided by a couple of cases 4 vs. 2, to a lesser extent expressed as a narrower maximum enhancement window in Fig. 9(b), explained by the more reduced activity followed by a higher H_2 reaction order. Nonetheless, it must be noted that a 50% enhancement can be achieved in both comparisons (and hence due to the

Table 5 Potential reactor/catalyst combinations

Pair no.	Reactor	Catalyst
1	Non-membrane	Ru-K/CaO
2	Non-membrane	Co-Ba/CeO ₂
3	Membrane	Ru-K/CaO
4	Membrane	Co-Ba/CeO ₂

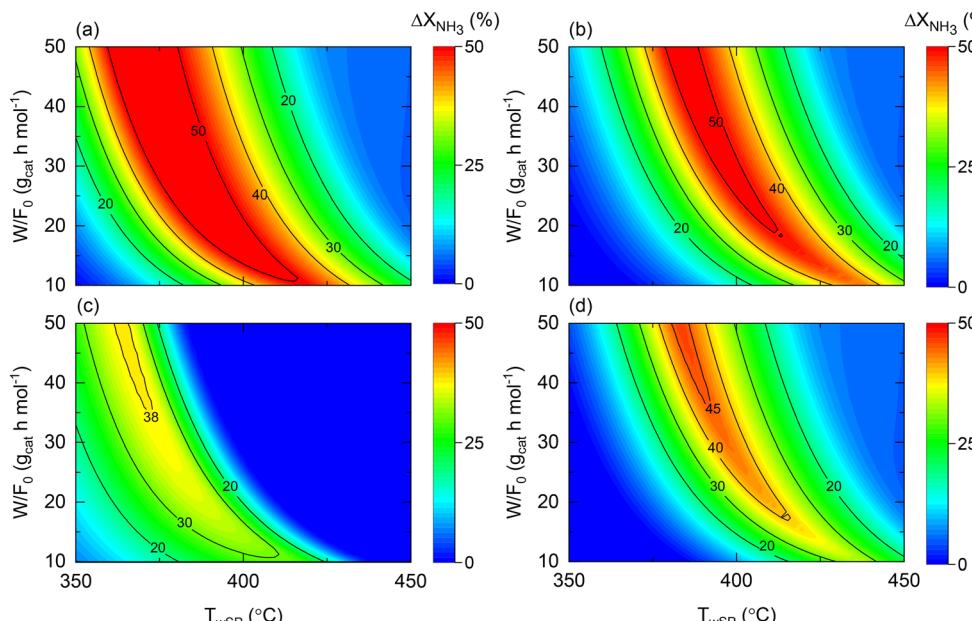


Fig. 9 Contour plots comparing NH_3 conversion enhancement as a function of space-time (W/F_0) and wall setpoint temperature (T_{wSP}) for cases (a) 3 vs. 1; (b) 4 vs. 2; (c) 3 vs. 4; and (d) 4 vs. 1. Simulation conditions: estimated kinetic, membrane, and heat transfer parameters (Table 4) and $W = 20 \text{ g}_{\text{cat}}$; $P_r = 10$; $P_p = 1 \text{ bar}$.

membrane reactor). The temperature dependence of the NH_3 conversion improvement in both analyses similarly can be directly related to the estimated E_a values for each catalyst. When assessing the effect of the catalyst in a membrane reactor (Fig. 9(c)), similar conversions can be obtained ($\Delta X_{\text{NH}_3} \sim 0$) over a wide range of W/F_0 and T_w values, and overall, under more severe conditions with higher NH_3 conversions.

Nonetheless, compared to the performance of both catalysts in a non-membrane reactor (e.g., Fig. S1(b) and S2(b)†), the utilization of a membrane reactor has a shrinking effect on the conditions requiring exact conversions, suggesting that catalyst activity plays a less crucial role when working with a membrane reactor. This argument is explicitly exemplified in Fig. 9(d), wherein remarkable conversion enhancements up to *ca.* 45% can be attained with a Co-based catalyst in a membrane reactor, if careful tuning of the operating conditions is performed. Previous observations reveal that the reactor type can change the catalyst comparison criteria and background. Notably, Fig. 9 results correspond to the following dimensionless number ranges: $0.01 < \text{Da}_0 < 0.5$; $0.03 < \text{Pe}_0 < 0.15$; $130 < \text{St}_0 < 700$.

The role of the wall temperature gradient on the membrane and non-membrane reactor performance is illustrated in Fig. 10, which shows the conversion difference between a constant wall temperature and the experimentally observed profile for both reactors. The wall temperature profile experimentally observed for the membrane reactor is assumed for the non-membrane counterpart as reactors with comparable dimensions and furnaces are considered. In the case of the non-membrane reactor (Fig. 10(a)), NH_3 conversion can decrease up to 8% due to the wall-temperature effect, whereas for the membrane reactor, this effect results in a maximal 4% performance drop. This

observation is in agreement with the previous insights provided by the model, suggesting that the membrane reactor can buffer temperature-related deviations from ideality. Although other important aspects related to heat transfer are not included in the model (e.g., radial gradients), the model proposed in the present study successfully

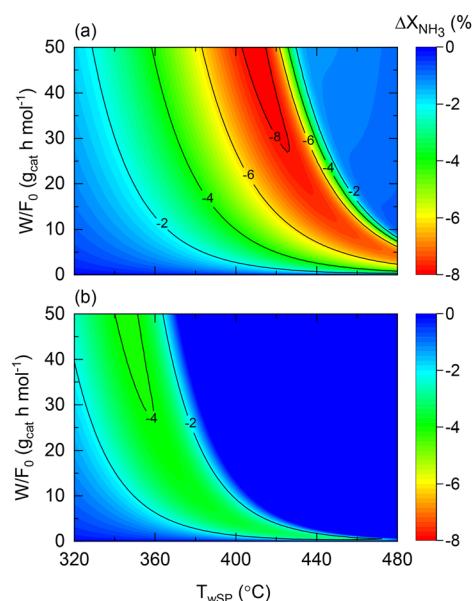


Fig. 10 Contour plots of NH_3 conversion loss using Co-Ba/CeO_2 with variable wall temperature compared to constant wall temperature ($T_w/T_{\text{w}} = 1$) in terms of wall temperature setpoint and space-time for (a) non-membrane reactor and (b) membrane reactor. Simulation conditions: estimated kinetics, membrane reactor, heat transfer parameters (Table 4), and $F_{\text{NH}_3,0} = 1 \text{ N mL min}^{-1}$; $P_r = 10$; $P_p = 1 \text{ bar}$.



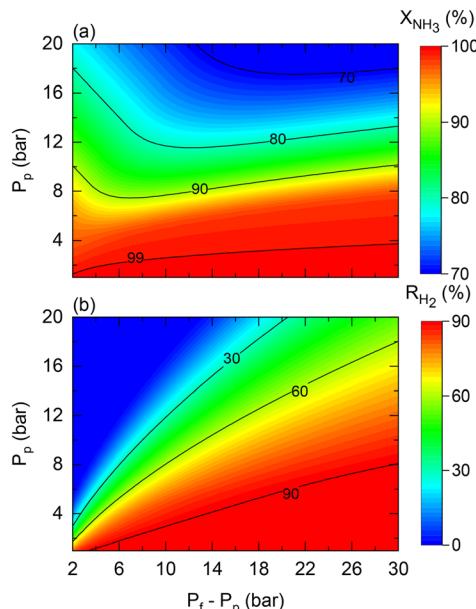


Fig. 11 Contour plots of membrane reactor performance using Co-Ba/CeO₂ as a function of permeate (P_p) and transmembrane differential pressure ($P_f - P_p$) in terms of (a) NH₃ conversion and (b) H₂ recovery. Simulation conditions: estimated kinetic, membrane, and heat transfer parameters (Table 4) and $T_{WSP} = 450$ °C; $W = 20$ g_{cat}; $W/F_0 = 22$ g_{cat} h mol⁻¹.

identifies and quantifies the effect of heat-transfer limitations. This is the case of the experimentally observed wall temperature profile, where process variables and the impact of the perturbations are measurable with confidence. In a larger-scale reactor, the role of such effects will be significantly more pronounced, calling for more detailed heterogeneous models that can account for them. Fig. 10 results correspond to the following dimensionless number ranges: $0 < Da_0 < 7$; $0.05 < Pe_0 < 0.08$; $240 < St_0 < 260$.

As shown in Fig. 4, the theoretical pressure ratio defines the required pressure feed to achieve complete NH₃ conversion at high permeate pressures. However, in practice, regardless of the scale of operation, membrane reactors can encounter restrictions that constrain the required pressure ratios for the production of pressurized H₂ from NH₃. That is the case of the maximum allowed transmembrane differential pressure, which in the case of the commercial Pd-Au/Al₂O₃ membrane used in this study is set to $P_f - P_p < 30$ bar by the manufacturer. For example, when targeting a permeate pressure of 15 bar, the maximum allowable operating pressure ratio is three. This active restriction induces changes in other operating conditions to obtain the desired NH₃ conversion. Fig. 11 shows predictions for (a) NH₃ conversion and (b) H₂ recovery considering the transmembrane differential pressure constraint. The maximum achievable NH₃ conversion decreases as the permeate pressure increases due to the maximum applied driving force limitation.

The fact that the pressure difference is limited (hence, working with pressure ratios to a greater extent) implies that obtaining complete NH₃ conversion at higher permeate pressures becomes increasingly complex under high

permeate pressure, thus requiring tuning of other operating conditions such as the temperature or space-time. For example, the reaction temperature must be increased if the resulting H₂ has to be pressurized. In this case, for a target H₂ pressure of 15 bar using the Co-Ba/CeO₂ catalyst under the conditions shown in Fig. 11, by raising the reaction temperature from 450 to 500 °C, NH₃ conversion and H₂ recovery are predicted to increase from 76% to 97% and 70% to 82%, respectively. Fig. 11 shows the results corresponding to the following dimensionless number ranges: $0.07 < Da_0 < 0.9$; $0.02 < Pe_0 < 0.2$; $St_0 = 300$.

4. Conclusions

We identified coupling opportunities for ammonia decomposition in a membrane reactor using modeling, experimental, and literature-review approaches. First, we developed a dimensionless model to explore the effects of different operating conditions, the impact of the reactor's isothermal or non-isothermal condition on the overall performance, and the implications of aiming for a high pressure and pure hydrogen product. Second, we fit the kinetic parameters for the in-house produced Ru-K/CaO and Co-Ba/CeO₂ catalysts and the permeance parameters of our commercial Pd-Au/Al₂O₃ membrane module. Finally, the developed model enables us to identify opportunities for different catalysts, membranes, and operation condition combinations where the reactor can lead to similar or improved results.

With the agnostic model, a representative catalyst ($a = 0.5$, $b = -0.75$, $E_a = \text{kJ mol}^{-1}$), and a highly H₂ selective membrane ($\alpha_{H_2/NH_3} = 10^5$; $\alpha_{H_2/N_2} = 10^5$), we found that NH₃ conversion enhancements can be achieved at the region bounded by $10^{-2} < Pe < 10^0$; $10^{-1} < Da < 100$ with $St > 100$ and $P_r > 2$. Under these operating conditions, experiments for fitting and parameter estimation were performed, and the values obtained agree with ammonia decomposition kinetics in the literature. Using a thorough parametric study of the kinetics, the buffering role of the membrane reactor in the influence of specific kinetic variables (e.g., reaction orders and activation energy) on the final performance could be identified. In this context, the catalyst with higher intrinsic activity (Ru-K/CaO), expressed as higher k_0 , has the highest ammonia conversion enhancement between a non-membrane and a membrane reactor. Nonetheless, model predictions for the Co-Ba/CeO₂ catalyst, which has lower intrinsic activity but is cost-effective, led to operating areas where enhancements of up to 50% were achievable with a membrane reactor (compared to a non-membrane). The latter reveals that using a membrane reactor alters the paradigm of catalyst design, as enhancements up to *ca.* 40% can be achieved with the Co-based catalyst compared to the Ru-based catalyst in a non-membrane reactor.

Our study provides solid arguments for employing the single-step ammonia decomposition process in a membrane reactor using a moderate activity catalyst, while making further strides towards scaling up the technology and obtaining high pressure, high-purity CO_x-free H₂.



Nomenclature

a	Reaction order for NH_3 , dimensionless	r_{rxn}	Rate of ammonia decomposition reaction, mol $\text{g}_{\text{cat}}^{-1} \text{s}^{-1}$
A_c	Cross-sectional area of the annular space, m^2	St	Stanton number
A_m	Superficial area of the membrane, m^2	T	Temperature of the catalytic bed, K
A_s	Reactor-oven contact surface, m^2	T_0	Temperature of the feed, K
b	Reaction order for H_2 , dimensionless	T_w	Reactor wall temperature, K
$C_{p,i}$	Heat capacity of component i , $\text{J mol}^{-1} \text{K}^{-1}$	T_{wsp}	Reactor wall setpoint temperature, K
$\hat{C}_{p,i}$	Dimensionless heat capacity of component i	U	Overall heat transfer coefficient between the catalytic bed and the reactor wall, $\text{kW m}^{-2} \text{K}^{-1}$
$C_{p,\text{NH}_3,0}$	Heat capacity of NH_3 in the feed, $\text{J mol}^{-1} \text{K}^{-1}$	V	Volume of the annular space, m^3
Da	Damköhler number	W_{cat}	Total catalyst mass, g
Da_{III}	Third Damköhler number	x_i	Molar fraction of component i in the catalytic bed
d_p	Average catalyst particle diameter, μm	X_{NH_3}	Conversion of NH_3 , % or fractional
E_a	Activation energy for ammonia decomposition reaction, kJ mol^{-1}	y_i	Molar fraction of component i in the permeate side
$E_{a,J}$	Activation energy for H_2 permeance, kJ mol^{-1}	z	Axial length of the reactor, m
$f_{\text{H}_2,L}$	Dimensionless molar flow rate of H_2 at the outlet of the catalytic bed		
f_i	Dimensionless molar flow rate in the catalytic bed of component i		
F_i	Molar flow rate in the catalytic bed of component i , mol s^{-1}		
$f_{\text{NH}_3,0}$	Dimensionless molar flow rate of NH_3 in the feed		
$F_{\text{NH}_3,0}$	Molar flow rate of NH_3 in the feed, mol s^{-1}		
$f_{\text{NH}_3,L}$	Dimensionless molar flow rate of NH_3 at the outlet of the catalytic bed		
G	Superficial mass velocity of the gas, $\text{kg m}^{-2} \text{s}^{-1}$		
J_i	Permeance of component i , $\text{mol s}^{-1} \text{m}^{-2} \text{Pa}^{-n}$		
$J_{i,0}$	Pre-exponential factor for permeance of component i , $\text{mol s}^{-1} \text{m}^{-2} \text{Pa}^{-n}$		
k_0	Pre-exponential factor for ammonia decomposition reaction, $\text{mol g}_{\text{cat}}^{-1} \text{s}^{-1} \text{bar}^{-(a+b)}$		
K_{eq}	Thermodynamic equilibrium constant, dimensionless		
L	Total reactor length, m		
n	Permeation order, dimensionless		
N	Total number of experiments		
N_s	Total number of species		
P	Total pressure in the catalytic bed, bar		
\hat{P}	Dimensionless total pressure in the catalytic bed		
P°	Standard pressure, bar		
Pe	Peclet number		
P_f	Total pressure in the feed, bar		
p_i	Partial pressure of component i , bar		
P_p	Total pressure in the permeate side, bar		
P_r	Pressure ratio, dimensionless		
$q_{\text{H}_2,L}$	Dimensionless molar flow rate of H_2 at the outlet of the permeate side		
q_i	Dimensionless molar flow rate of component i in the permeate side		
Q_i	Molar flow rate of component i in the permeate side, mol s^{-1}		
$q_{\text{NH}_3,L}$	Dimensionless molar flow rate of NH_3 at the outlet of the permeate side		
R	Universal gas constant, $\text{kJ mol}^{-1} \text{K}^{-1}$		
R_{H_2}	Hydrogen recovery, % or fractional		

Abbreviations

SSR Sum of square residuals

Greek symbols

$\alpha_{\text{H}_2/i}$	Permeation selectivity of component i relative to H_2 , dimensionless
β	Temkin–Pyzhev empirical constant, dimensionless
$\Delta G_{\text{rxn}}^{\circ}$	Standard Gibbs free energy of ammonia decomposition reaction, kJ mol^{-1}
$\Delta H_{\text{rxn}}^{\circ}$	Standard heat of ammonia decomposition reaction, kJ mol^{-1}
ΔX_{NH_3}	Ammonia conversion enhancement, %
ζ	Dimensionless axial reactor length
θ	Dimensionless temperature in the catalytic bed
μ	Viscosity of the gas, $\text{kg m}^{-1} \text{s}^{-1}$
v_i	Stoichiometric number of component i
ρ	Density of the gas, kg m^{-3}
σ	Standard deviation
τ	Space time, $\text{g}_{\text{cat}} \text{h mol}^{-1}$
ϕ	Catalytic bed porosity, $\text{m}_{\text{gas}}^{-3} \text{m}_{\text{bed}}^{-3}$
χ	Fractional target variable

Subscripts

H_2	Hydrogen
N_2	Nitrogen
NH_3	Ammonia
0	At the reactor inlet

Superscripts

cal	Calculated
exp	Experimental

Author contributions

Natalia Realpe: investigation; data curation; software; formal analysis; visualization; writing – original draft. Shekhar R. Kulkarni: methodology; software; formal analysis; visualization; writing – original draft. Jose L. Cerrillo: investigation; validation; formal analysis; writing – original draft. Natalia



Morlanes: investigation; validation; formal analysis; writing – review & editing. Gontzal Lezcano: conceptualization and formal analysis. Sai P. Katikaneni: conceptualization; formal analysis; resources; supervision. Stephen N. Paglieri: conceptualization; resources. Mohammad Rakib: resources; supervision. Bandar Solami: resources; supervision. Jorge Gascon: conceptualization; funding acquisition; supervision; writing – review and editing. Pedro Castaño: conceptualization; methodology; funding acquisition; supervision; project administration; writing – review and editing.

Conflicts of interest

There are no conflicts to declare.

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