RSC Mechanochemistry



PAPER

View Article Online



Cite this: RSC Mechanochem., 2025, 2,

Rational screening of milling parameters for Ru-Na/Al₂O₃ dual-function materials for integrated CO₂ capture and methanation†

Andrea Braga, D Maila Danielis, D * Sara Colussi D and Alessandro Trovarelli

The interest in the use of mechanochemistry as a green alternative to conventional solution-based synthesis methods has been steadily growing in recent years. Recently, Dual-Function Materials (DFMs) have been explored for the preparation of multicomponent systems which combine a sorbent and a catalytic phase co-supported on a support oxide for the capture of CO2 from flue gases and its subsequent conversion into added-value products when exposed to H2 (or CH4) in a chemical-loopingtype reaction. However, the complexity of setting the right milling parameters, which are interconnected and strongly dependent on the precursor materials, is exacerbated in the multi-component system. In this work, we address this issue by employing a Design of Experiments (DoE) statistical approach for the screening of the most relevant milling parameters for the synthesis of Ru-Na/Al₂O₃ DFMs for integrated CO₂ capture and methanation (ICCU-MET). The milling intensity and the organic precursors proved to be the key factors positively affecting the DFMs' capture capacity and CH₄ conversion, respectively.

Received 23rd December 2024 Accepted 3rd May 2025

DOI: 10.1039/d4mr00147h rsc li/RSCMechanochem

Introduction

Among the several strategies available in the field of CO₂ emission abatement,1 the recent implementation of the socalled integrated CO2 capture and utilisation (ICCU)2,3 technologies is being studied as an alternative to classical processes such as amine sorption4 and CaO carbonation,5 which are very efficient in CO₂ capture but require an additional process for its valorisation. The ICCU process, instead, is based on cyclic CO₂ capture alternated with a CO2 conversion step on the same reactive bed, performed by switching the reaction atmosphere between the CO2-containing flue gas and the reactive gas.6

The key components of this technology are the materials used to capture and valorise CO2, called Dual-Function Materials (DFMs).7 These materials belong to the family of heterogeneous catalysts and are solid nanomaterials composed of two functionalities dispersed on a support oxide: the sorbent function, usually linked to elements with basic characteristics such as alkaline metals, and the catalytic function, normally associated with transition metals.^{2,8} The two functions need to work in synergy to be able to first capture CO2 and then efficiently convert it into useful products such as CH4, CO and H2 (in syngas).9 Indeed, one of the most interesting uses of CO2 is to recycle it as a feedstock for e-fuels.10 In principle, renewable energy can be used to produce green hydrogen by electrolysis,

Dipartimento Politecnico e INSTM, Università degli Studi di Udine, Via del Cotonificio 108, 33100 Udine, Italy. E-mail: maila.danielis@uniud.it

available. See DOI: † Electronic supplementary information (ESI) https://doi.org/10.1039/d4mr00147h

which is then combined with CO₂ for the synthesis of CH₄ to store energy and be used as a sustainable fuel.

The main advantage of the ICCU process for DFMs is that, after the capture step directly from the emission source, the CO₂ valorisation to useful products (such as CH4, CO + H2, and CH₃OH) is performed in the same reactor, thus removing the necessity to regenerate the sorbent material and transport the CO2 to another facility. Hence, energy and transport costs can

In this regard, the most commonly studied DFM formulations combine Al₂O₃ as the support oxide, since it has a relatively high surface area and good mechanical and thermal stability, Na, Ca, K, and Ba oxides as sorbent phases due to their basic properties, and Ni and Ru as the elements for the catalytic phase. 2,3,8,11 The most suitable DFMs for the methanation of CO2 are composed of Ru or Ni, and Ca and Na as sorbent phases or a mixture of them to optimise the CO2 capture and release at different temperatures.3,12-14 The most commonly used methods for the preparation of DFMs are based on wet chemistry, usually by impregnating metal solutions on support oxides, followed by co-precipitation and sol-gel methods.2 These standard methods are suitable for producing materials with strong interactions among the different components, although they often require several impregnation and drying steps, long thermal treatments, and calcination steps at high temperatures to decompose remove the element precursors. mechanochemical methods are a more sustainable alternative for the preparation of heterogeneous catalysts for several reasons: 15,16 they are solventless methods, they can be scaled up to industrial production,17 and in principle, they can reduce or

remove the need for thermal treatments yielding the final working materials with quick syntheses at room temperature and reduced energy consumption. 18-23 Successful attempts have been reported in several fields, 16,24 but limited research has been conducted on the application of dry milling to the synthesis of dual-function materials.

Our previous work25 explored the feasibility of mechanochemically preparing DFMs based on RuNi/(Na2O or CaO)/ CeO₂-Al₂O₃ for CO₂ capture and conversion to CO by the reverse water-gas shift reaction. In this work, we expand the use of dry ball milling with a rational investigation of the effect of the milling parameters on the CO2 capture and methanation performance in the integrated carbon capture and methanation process (ICCU-MET). The DFMs prepared in this study are composed of Ru + Na₂O/Al₂O₃, which is a reference composition^{2,3,12,26,27} mostly suitable for medium-temperature (250–400 $^{\circ}$ C) CO₂ capture and methanation. 28,29 The simple composition allows us to focus our attention on the milling parameters and their effects on the materials' performance. The capture properties of Na-based DFMs are influenced by the nature of the Na precursor,29 and, similarly, different precursors used in dry ball milling can affect the final catalytic properties. 30,31 For these reasons, the choice of Ru and Na precursors and three different milling regimes were thoroughly studied to account for the different nature of the precursors and the effect of milling on the obtained DFMs. A full factorial experimental design was used to efficiently combine these parameters and to analyse the results, highlighting the most relevant parameters affecting the performance of DFMs.

Experimental

Synthesis of dual-function materials (DFMs) by dry ball milling

The DFMs were prepared by mixing both Na and Ru precursors with 5%La₂O₃-γ-Al₂O₃ (MI386, Grace) in a single-step dry ball milling process using a Fritsch Pulverisette 23 Mini-mill. The MI386 support was previously calcined at 550 °C for 2 h in air. With 890 mg of MI386, appropriate amounts of Ru acetylacetonate (RuAc, Strem Chemicals), metallic Ru black powder (RuM, Strem Chemicals), Na₂CO₃ (NaC, Carlo Erba) and NaNO₃

(NaN, VWR Chemicals) were mixed to obtain respectively 1 wt% metallic Ru and 10 wt% Na₂O equivalent. The powders were loaded into a 15 mL ZrO₂ jar with a single ZrO₂ ball ($m = 10 \text{ g}, \emptyset$ = 15 mm), and the ball-to-powder mass ratio varied between 9 and 10 depending on the different precursors used (Table 1).

The milling intensity was set at low, medium, and high intensity regimes by coupling the milling time and milling frequency: the low-intensity samples were milled for 15 min at 15 Hz, the medium-intensity ones were milled for 27 min at 27 Hz, and the high-intensity samples were milled at 35 Hz for 35 min. The milling was paused after 15 min to clean the jar and ball surfaces and manually mix the powders. The loose powder was first removed from the jar, and then the milling medium was scratched with a brush to avoid the formation of large amounts of unmixed powders on the walls of the jar. The samples were ready and used directly after the milling procedure.

A full factorial experimental design, 32,33 generated using Minitab 21.4, was used to create the sample series to study the combination of different parameters and their combined effects: the milling intensity, the Ru precursor and the Na precursor. Given the categorical nature of both Ru and Na precursors, a total of 12 samples were generated. The sample series contained all the combinations of the three parameters to study the effect of each one on different DFM characteristics, such as the CO2 adsorption capacity and the CH4 production rates. The samples were prepared and tested in a random order to minimise and distribute the environmental effects.31

The prepared DFMs, as reported in Table 1, were named x-Ru(M,Ac)Na(C,N)/MI386-(15,27,35), where x represents the identification number, (M,Ac) represents the Ru precursor, (C,N) represents the Na precursor, and the last number represents both the milling time and frequency. Several Na/MI386 samples were also prepared by milling different loadings of the two Na precursors with MI386 at the three different milling regimes as preliminary samples to find the experimental design ranges.

Characterisation

Powder X-ray diffraction (XRD) measurements were carried out with a Philips X'Pert diffractometer with a Cu cathode (Cu K α λ

Table 1 List of DFMs prepared with the different combinations of milling parameters and Na and Ru precursors

#	Sample	Time (min)	Frequency (Hz)	Ru precursor	Na precursor	BPR
1	RuMNaC/MI386-15	15	15	Ru metal	Na_2CO_3	10
2	RuMNaC/MI386-35	35	35	Ru metal	Na_2CO_3	10
3	RuMNaN/MI386-15	15	15	Ru metal	$NaNO_3$	9
4	RuMNaN/MI386-35	35	35	Ru metal	NaNO ₃	9
5	RuAcNaC/MI386-15	15	15	RuAc	Na_2CO_3	9
6	RuAcNaC/MI386-35	35	35	RuAc	Na_2CO_3	9
7	RuAcNaN/MI386-15	15	15	RuAc	NaNO ₃	9
8	RuAcNaN/MI386-35	35	35	RuAc	$NaNO_3$	9
9	RuMNaC/MI386-27	27	27	Ru metal	Na_2CO_3	10
10	RuMNaN/MI386-27	27	27	Ru metal	NaNO ₃	9
11	RuAcNaC/MI386-27	27	27	RuAc	Na_2CO_3	9
12	RuAcNaN/MI386-27	27	27	RuAc	NaNO ₃	9

= 1.5406 Å, 40 kV and 40 mA) collecting the diffractograms with a step of 0.02° and 40 s per step; higher resolution patterns were collected on the spent DFMs from 30 to 50° with a step of 0.01° and 320 s per step. The patterns were fitted with pseudo-Voigt functions in OriginLab 2021 to isolate the metallic Ru peak FWHM at 44° and estimate the crystallite size with Scherrer's equation:³⁴

$$d = \frac{0.9\lambda}{\beta \cos(\theta)}$$

where $\lambda = 1.5406$ Å is the X-ray wavelength and

$$\beta = \sqrt{\text{FWHM}_{\text{obs}}^2 - \text{FWHM}_{\text{instr}}^2}$$

is the peak broadening corrected for the instrumental broadening, measured on a Si standard, at a given 2θ .

Thermogravimetric analysis (TGA) was conducted using a TGA Q5500 (TA Instruments) on as-prepared samples in 25 mL min $^{-1}$ of synthetic air (*i.e.*, dry air in a commercial cylinder, with no moisture or CO₂) to observe the precursors' decomposition patterns by loading about 10 mg of powders on platinum pans and heating from room temperature to 900 °C (10 °C min $^{-1}$). Additionally, TGA was used to study the activation of the DFMs in 5% $\rm H_2/N_2$ (25 mL min $^{-1}$) by loading about 10 mg of DFM on platinum pans and heating from room temperature to 400 °C and holding for 30 min (10 °C min $^{-1}$).

Specific surface area and pore size distribution values were obtained using a Nova 800 analyser (Anton Paar) to record physisorption isotherms at $-196~^{\circ}\mathrm{C}$ in N_2 . The specific surface area and pore size were estimated using the Brunauer–Emmett–Teller (BET) and Barrett–Joyner–Halenda (BJH) methods, respectively. Prior to the analysis, all samples were outgassed in a vacuum at 150 $^{\circ}\mathrm{C}$ for 90 min. Selected samples were reduced ex~situ for 1 h at 400 $^{\circ}\mathrm{C}$ in 10% $\mathrm{H_2/N_2}$ (50 mL min $^{-1}$, 10 $^{\circ}\mathrm{C}$ min $^{-1}$) to measure the surface area and pore distribution of the activated DFMs.

Temperature Programmed Reduction (H_2 -TPR) measurements were performed from the activation step inside the reactor used for the ICCU-MET tests. Briefly, 250 mg of DFM

were placed inside a fixed bed quartz reactor (12 mm internal diameter) and activated in 50 mL min $^{-1}$ of 10% $\rm H_2/N_2$ from room temperature to 400 °C and holding for 1 h (10 °C min $^{-1}$). The evolution of $\rm H_2$, CO $_2$, CO and CH $_4$ was monitored with an ABB AO2020 online gas analyser recording data every 5 s and integrated with OriginLab.

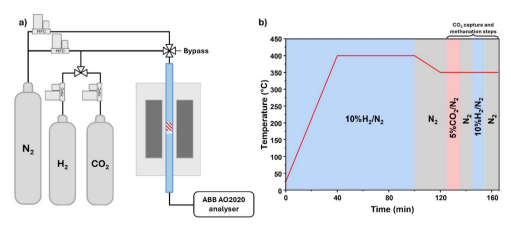
Integrated CO₂ capture and methanation testing

The DFMs were tested for the cyclic capture and methanation of CO₂ (ICCU-MET) in a fixed-bed quartz reactor with a 12 mm internal diameter. The reaction apparatus and the testing procedure are reported in Scheme 1. The DFM powders were supported on a quartz wool bed and a thermocouple was placed above the catalytic bed. 250 mg of DFM was activated in situ with 50 mL min $^{-1}$ of 10% H₂/N₂ from room temperature to 400 °C for 1 h (10 °C min⁻¹) and cooled to 350 °C in N_2 . After the temperature was stable, ICCU-MET cyclic testing was conducted as follows: (i) 10 min of CO₂ capture with 50 mL min⁻¹ of 5% CO_2/N_2 ; (ii) a 10 min purge with 100 mL min⁻¹ of N_2 ; (iii) 10 min of methanation with 50 mL min⁻¹ of 10% H₂/N₂; (iv) a final 10 min purge with 100 mL min⁻¹ of N₂. In total, five cycles were performed for each test. The evolution of H2, CO2, CO and CH4 was followed online with an ABB AO2020 analyser recording data points every 5 s. The molar flows were integrated with OriginLab to obtain the moles of CO, CO2 and CH4:

$$n_i = \frac{1}{m_{\rm DFM}} \int_{t_0}^{t_1} \dot{F}_i \mathrm{d}t$$

where $i = (CO, CO_2, CH_4)$, m_{DFM} is the DFM mass loaded in the reactor, and F_i is the molar flow.

To estimate the amount of $\rm CO_2$ captured, blank measurements were taken at 350 °C in a reactor loaded with 250 mg of coarse quartz grains previously calcined at 1200 °C. Three injections of 10 min of 50 mL min⁻¹ of 5% $\rm CO_2/N_2$ were performed, similar to the $\rm CO_2$ capture steps to obtain the reference area. The blank measurement was repeated after the testing campaign to check the reproducibility and stability of the reaction system. The $\rm CO_2$ capture capacity at each step was



Scheme 1 (a) Reaction apparatus scheme composed of independently controlled mass flow controllers connected to a 4-way valve leading to the reactor or a bypass; (b) ICCU-MET reaction procedure. The CO_2 capture and methanation steps represent one cycle; 5 cycles are performed per experiment.

estimated by subtracting the area of the integrated CO2 signal from the reference CO2 area (see Fig. S1†). In addition, considering that all the DFMs produced some amounts of CO during the capture steps and that during the purge some weakly physisorbed CO₂ was released, the captured CO₂ was corrected by subtracting these two contributions as well:

$$n_{CO_2}^{captured} = n_{CO_2}^{blank} - n_{CO_2}^{capture} - n_{CO_2}^{purge} - n_{CO}^{capture}$$

The CO₂ conversion to CH₄ during the methanation step was calculated considering the release of CO2 during the methanation step:

$$X_{\text{CO}_2} = \frac{n_{\text{CH}_4}}{n_{\text{CO}_2}^{\text{captured}} - n_{\text{CO}_2}^{\text{released}}} \times 100$$

The methanation selectivity was calculated considering the production of CH₄ and CO and the release of CO₂:

$$S_{\text{CH}_4} = \frac{n_{\text{CH}_4}}{n_{\text{CH}_4} + n_{\text{CO}} + n_{\text{CO}_2}} \times 100$$

Results and discussion

Evaluation of alkali-supported oxide binary systems

A preliminary investigation by BET, XRD, and TG analyses was carried out on metal-free Na(C,N)/Al2O3 systems to ensure significant variability among the different milling parameters and precursor choices for the factorial design of experiments. The results so obtained, reported in Fig. 1, also provided reference samples for the fully formulated DFMs investigated in the following.

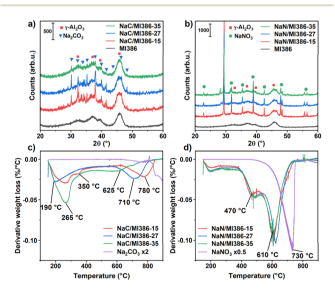


Fig. 1 XRD diffractograms of the (a) Na₂CO₃ and (b) NaNO₃ precursors milled with MI386 at different intensities. Derivative mass loss signals during the TGA in air of (c) NaC/MI386 and (d) NaN/MI386. Derivative signals for the bare salts are reported either doubled $(Na_2CO_3, \times 2)$ or halved $(NaNO_3, \times 0.5)$ in intensity for clarity.

The crystalline structure of the as-prepared Na/MI386 samples was studied by XRD to observe the modifications induced on the Na salts by milling at different regimes. The XRD patterns of NaC/MI386 and NaN/MI386 are reported in Fig. 1a and b. The identified phases are Al₂O₃ (PDF: 00-010-0425), Na₂CO₃ (PDF: 00-037-0451 and 00-019-1130), and NaNO₃ (PDF: 01-085-0850). The intensity of the Na₂CO₃ pattern decreased with increasing milling intensity, indicating a progressive reduction in the crystallinity degree. Remarkably, a significant loss in crystallinity was observed in the sample milled at 35 Hz for 35 min, while less striking differences were observed between the sample milled at 15 Hz for 15 min and the one milled at 27 Hz for 27 min. Regarding the NaN/MI386 samples, the NaNO3 pattern was less affected by the increasing milling intensity and large crystallites were still present in the sample NaN/MI386-35. The NaNO₃ crystallinity decreased gradually by about 33% from the lowest to the highest milling intensity. The peak shape and FWHM did not change, suggesting minor modifications to the NaNO3 structure. In all samples, the Al₂O₃ crystalline structure did not change upon increasing the milling intensity.

In parallel, the BET specific surface areas, the BJH pore diameters and volumes are reported in Table S1,† while the N2 isotherms and the pore size distributions are shown in Fig. S2.† In general, adding both precursors to MI386 by milling decreased the support surface area and pore volume; at higher milling intensities, the BET area decreased further, and the pore structure collapsed. The use of NaNO₃ led to a lower BET surface area and pore volume compared to Na2CO3, likely because of the higher amount (23 vs. 16 wt%, respectively) to result in 10 wt% Na₂O. However, similar trends were observed for the pores' diameter and volume, with significant losses, especially at the highest milling intensity.

To investigate whether the changes in crystallinity would affect the decomposition behaviour, the weight of the Na precursors was followed in TGA measurements by calcining the samples from room temperature to 900 °C (10 °C min⁻¹) in air. The derivative weight loss profiles of the NaC/MI386 and NaN/ MI386 samples are reported in Fig. 1c and d and the respective weight loss signals are in Fig. S3,† together with the pure Na2CO3 and NaNO3 samples reported as references. It clearly appears that the weight loss of the Na(C,N)/MI386 samples was influenced by the presence of Al2O3, which shifted the salt reduction at lower temperatures. Indeed, pure Na2CO3 decomposition started after 850 °C, as it is a very stable compound, 28,35 while pure NaNO3 decomposed in a single step peaking at 730 $^{\circ}$ C. When supported on Al₂O₃, the characteristic decomposition occurred at much lower temperatures, most noticeably for NaC/ MI386. On the latter, the milling intensity also plays a relevant role, while NaN/MI386 samples were mainly unaffected. In fact, the shift towards lower temperatures of the Na₂CO₃ decomposition steps observed with increasing milling intensity can be ascribed to a larger fraction of Na₂CO₃ species in close contact with Al₂O₃, promoted by milling, which is reported to favour the decomposition of the carbonate. 35,36

Overall, the TGA results were in good accordance with the XRD patterns: for Na₂CO₃, higher milling intensity resulted in more amorphous Na_2CO_3 and enhanced decomposition at low temperatures (Fig. 1a and c);^{35,36} for $NaNO_3$, the change in milling parameters barely affected its crystalline structure as well as the decomposition rate (Fig. 1b and d).

Phase composition and thermal stability of the DFMs

First, the morphological and redox properties of the 12 DFM samples of the statistical design were investigated. A focus of the XRD patterns of the as-prepared DFMs is reported in Fig. 2, while full-range patterns are reported in Fig. S4.† In Fig. S4e,† the XRD diffractograms of the four different precursors are also reported in the range 35-50°. In agreement with what observed for NaC/MI386 samples (Fig. 1a), on Ru(M,Ac)NaC/MI386 samples, the intensity of the Na₂CO₃ pattern decreased with increasing milling intensity (Fig. 2a and S4a†) and the peaks were broader; this could be related to the gradual amorphisation of the Na₂CO₃ crystallites with increasing milling intensity. No Ru-related patterns could be identified for both RuM and RuAc DFMs; in particular, the metallic Ru peak at 44.0° was not observed, indicating a high dispersion for the RuM precursor. The NaNO₃ pattern was less affected by the increasing milling intensity: the area of the main peak, centred at 29.3° 2θ (Fig. 1b), decreased by 34% on the samples milled at 35 Hz for 35 min compared to the ones milled at 15 Hz for 15 min, and no differences were found by milling with RuM or RuAc, confirming the low influence of the milling intensity on the NaNO₃ crystallinity. Similarly to the Na₂CO₃-based DFMs, no patterns

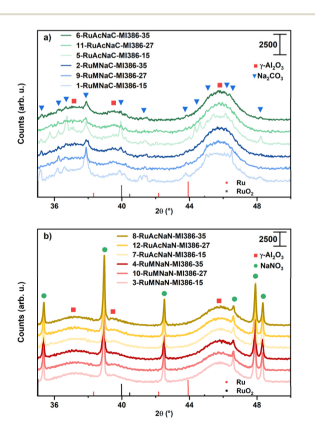


Fig. 2 Details of the XRD patterns of the as-prepared (a) Na_2CO_3 -based DFMs and (b) $NaNO_3$ -based DFMs.

related to Ru species were observed suggesting a high dispersion of both Ru precursors (Fig. 2). On all samples, the $\gamma\text{-Al}_2O_3$ crystalline phase did not show any modification compared with the bare support and at the different milling intensities. Overall, the as-prepared DFMs were similar to the binary Na(C,N)/MI386 samples reported in Fig. 1 from a structural point of view, suggesting that the presence of ruthenium has a negligible effect on the structural properties of the materials.

However, the presence of Ru influenced the decomposition behaviour of the Na precursors in air, as shown in Fig. S5 and S6,† compared with the Ru-free supports shown in Fig. 1c and d. In Na2CO3-based DFMs, the samples milled at medium and high intensity displayed similar weight loss signals occurring earlier than the respective DFMs milled at 15 Hz and 15 min. The presence of Ru promoted the low-temperature decomposition in both sample subgroups, reducing the decomposition temperature by 20-70 °C compared with the metal-free supports (compare Fig. 1c with S5c and d†). DFMs based on RuAc showed a sharper and more intense decomposition at 190-240 °C associated with the RuAc decomposition, which accelerated the Na₂CO₃ decomposition. At high temperatures, low-intensity milled samples displayed higher weight loss than metal-free NaC/MI386-15, with peak temperatures reduced by 30-50 °C due to Ru. The samples milled at higher intensities showed minor differences compared to the metal-free supports at high temperatures.

Regarding the NaNO₃ samples, as observed in XRD (Fig. 1 and S4†) and the TGA analysis of the metal-free supports (Fig. S1d†), even the presence of Ru has a minimal impact on the decomposition properties observed across milling intensities. RuM-based DFMs showed the same weight loss profiles; compared with the metal-free supports, the peak at 480 °C was only slightly anticipated at 320 °C, while the peak at about 600 °C remained unchanged. The RuAc-based DFMs showed a sharp peak at 280 °C with a shoulder at 190 °C corresponding to the acetylacetonate and part of the NO₃ decomposition. The high-temperature decomposition was unaffected by both the milling intensity and the Ru precursor for the samples milled at medium-high regimes, while only for the sample 7-RuAcNaN/MI386-15 it shifted to 560 °C.

However, in the current application, no prior calcination treatment in air was needed on the DFM samples, and their activation was carried out directly in the reactor in H2 up to 400 °C to decompose the Ru and Na precursors. Hence, additional TGA measurements were used to study their reactivity under H₂ atmosphere. The DFMs' weight profiles in H₂ and the derivative signals are reported in Fig. 3 and S7,† respectively. The four samples milled at low intensity showed a higher residual weight after the activation compared to the samples milled at medium-high intensity, indicating a lower degree of Na salt decomposition. For comparison, in Fig. S8,† the weight loss at 400 °C under H2 is reported alongside the weight loss values measured in air at 400 °C and 900 °C, where Na precursor decomposition was completed for all the samples (from Fig. S5 and S6†). The samples milled at low intensity showed a smaller weight loss in H2 compared with the weight loss at 900 °C in air, while the samples milled at medium and high intensity showed

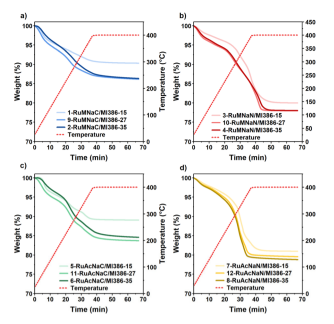


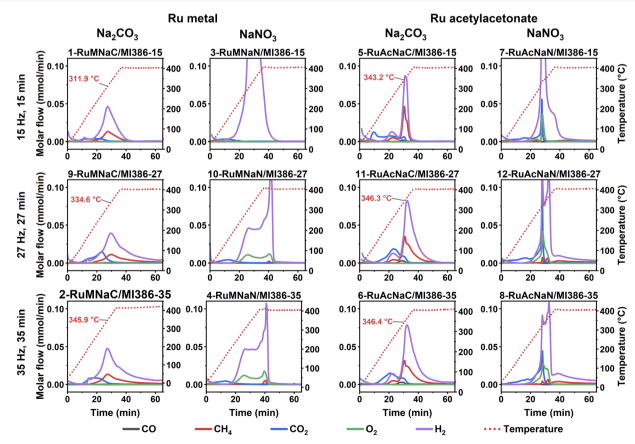
Fig. 3 TGA weight profiles in 5% H₂/N₂ up to 400 °C of DFMs based on (a) $RuM + Na_2CO_3$, (b) $RuM + NaNO_3$, (c) $RuAc + Na_2CO_3$ and (d) RuAc+ NaNO₃.

comparable weight losses suggesting the complete decomposition at 400 °C in H₂. For comparison, all the samples showed 1.5-3 times smaller weight losses at 400 °C in air, highlighting the role of Ru in activating H₂ to enhance the low-temperature Na precursors' decomposition. These results show that medium-high milling intensities and H2 activated by Ru allowed the complete decomposition of the Na precursors at 400 °C.

Redox behaviour of the DFMs

The derivative signals in Fig. S7[†] can be compared to the H₂-TPR signals obtained during the in situ activation in the reactor reported in Fig. 4. The H2-TPR step carried out in the reactor allowed us to follow the gas evolution of the RuNa/MI386 samples, which also provides insights into the passive Direct Air Capture (DAC) capacity of the DFMs.25

On Na₂CO₃-based DFMs, as the temperature increased the CO₂ adsorbed from the atmosphere was released and gradually converted to CH₄ starting from 250 °C. The samples based on RuAc showed a more intense and sharper CH₄ release peak around 350 °C suggesting the formation of more active sites, while the samples based on RuM showed a longer tail at 400 °C, indicating the presence of less active Ru species. Furthermore, the CH₄ production slowed with increasing milling intensity. The maximum CH₄ production rate was achieved at 340-350 °C, which was consequently taken as the testing temperature. Similarly, the NaNO₃ samples showed comparable profiles in both the TGA and the reactor (see Fig. S7b, d and 4), corresponding to the decomposition of NaNO3. The samples based



H₂-TPR signals measured during the activation procedure inside the reactor with 50 mL min⁻¹ of 10% H₂/N₂

on RuM showed a broader temperature range for NaNO₃ decomposition, starting at 200–250 °C and ending with a sharp event after 5 min at 400 °C in both the TGA and in the reactor. The two samples milled at medium-high intensity showed a very similar profile and during the H₂-TPR the O₂ signal was recorded, which probably originated from the NO_x species detected from the O₂ sensor. Regarding the RuAcNaN/MI386 samples, the decomposition of NaNO₃ started at about 280 °C and a quick release of both CO₂ and O₂ was observed at about 320 °C with an exothermic reaction, followed by a sharp consumption of H₂ at 350–360 °C. Also in this sample subgroup, the two DFMs milled at medium-high intensity showed the same decomposition pattern, suggesting that similar structures and reactivity were obtained already by milling the Ru and Na precursors for 27 min at 27 Hz.

In Table 2, the quantitative analysis of the evolution of CH₄, CO₂ and CO released during the in situ DFM activation shown in Fig. 4 is reported. The evolution of carbon species can be related both to the presence of the precursors, as in the case of the Na₂CO₃-based and RuAc-based DFMs, and to the atmospheric CO₂ captured at room temperature.²⁵ Overall, all the samples showed higher total C species than the amount loaded with the precursors, suggesting that they are all characterised by some passive CO2 capture capacity. The evolution of the carbon species by passive Direct Air Capture (DAC) and the CH₄/CO₂ ratio are also summarised in Fig. S9.† The samples based on NaC and RuAc release the highest amount of carbon, both due to the precursors and DAC, followed by the RuMNaC/MI386 and RuAcNaN/MI386 samples. The least performing were the ones with no carbon contribution from the precursors (RuMNaN/ MI386), which also exhibited the lowest CH₄ production. Conversely, the RuAc-based DFMs displayed the highest methane release, suggesting higher reactivity.

All the samples showed the same trend of both the total $\rm CO_2$ release as a function of the milling intensity: the lowest values were related to the samples milled for 15 min at 15 Hz, while the highest amounts of $\rm CO_2$ released and $\rm CH_4$ production were achieved respectively at medium and high intensities.

ICCU-MET performance and DoE analysis

Finally, the ICCU-MET performance of the prepared DFMs was evaluated by performing five cycles of CO₂ capture and methanation at 350 °C (5% CO₂/N₂ and 10% H₂/N₂, 50 mL min⁻¹), in order to investigate the effects of the Ru and Na precursors and the milling parameters. The CO₂ capture capacity and the CH₄ production for all the cycles are reported in Fig. 5, while the transient profiles of CO, CO₂ and CH₄ during the 5th methanation step are shown in Fig. 6. The summary of the DFMs' performance quantitative analyses is reported in Table S2.† The reactions involved in the ICCU-MET process can be described by the following chemical equations:³⁷

CO₂ capture: Na₂O + CO₂
$$\rightleftharpoons$$
 Na₂CO₃
NaOH + CO₂ \rightleftharpoons NaHCO₃
CH₄ formation: Na₂CO₃ \rightleftharpoons Na₂O + CO₂
CO₂ + 4H₂ \rightleftharpoons CH₄ + 2H₂O
Na₂O + H₂O \rightleftharpoons 2NaOH

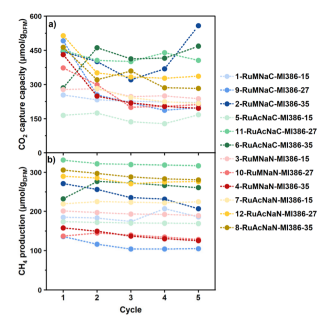
where the methanation of CO₂ requires an available neighbouring catalytic phase, such as Ru. From Fig. 5, in general, the samples based on RuAc (yellow and green lines) captured higher amounts of CO₂ than the DFMs based on RuM. The latter captured high amounts of CO₂ only during the first capture step and then rapidly lost the initial capture capacity in the subsequent cycles, with the exception of the sample 2-RuMNaC/MI386-35, which increased the capture capacity.

In parallel, the RuAc samples produced more CH₄ during the methanation step. A table comparing the activity of the best performing dry milled samples compared to wet-based DFMs from the literature is reported in the ESI (Table S3).† The samples milled at medium and low intensity showed a more stable performance within five cycles, while the highest milling regime resulted in a constant decrease in the CH₄ production.

At the beginning of each methanation step, on all samples some CO₂ and CO were released together with CH₄, as can be

Table 2 Evolution of CO₂, CH₄ and CO gases during the H₂-TPR activation step in 50 mL min⁻¹ of 10% H₂/N₂ up to 400 °C for 1 h. The values are reported as μmol q_{DFM}^{-1} . The samples are grouped by Ru and Na precursors, respectively

#	Sample	CO_2	CH_4	CO	Total C	C from precursors	C from passive DAC
1	RuMNaC/MI386-15	293	598	0	891	151	740
9	RuMNaC/MI386-27	709	776	21	1506	151	1355
2	RuMNaC/MI386-35	534	931	4	1468	151	1317
3	RuMNaN/MI386-15	182	49	0	232	0	232
10	RuMNaN/MI386-27	289	50	0	339	0	339
4	RuMNaN/MI386-35	287	78	0	365	0	365
5	RuAcNaC/MI386-15	724	813	24	1561	328	1233
11	RuAcNaC/MI386-27	916	1224	57	2196	328	1868
6	RuAcNaC/MI386-35	872	1287	59	2218	328	1890
7	RuAcNaN/MI386-15	640	239	0	879	165	714
12	RuAcNaN/MI386-27	1029	372	10	1412	165	1247
8	RuAcNaN/MI386-35	958	436	12	1406	165	1241



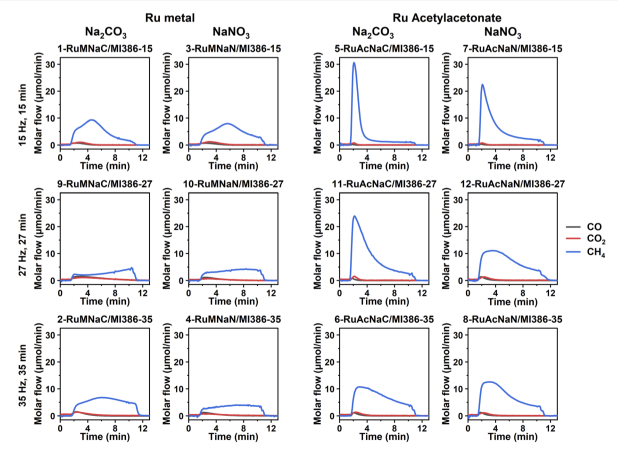
(a) CO₂ capture capacity and (b) CH₄ production for each cycle of the DFMs tested at 350 °C under 5% CO₂/N₂ and 10% H₂/N₂ (50 $mL min^{-1}$, 250 mg, and 10 min each cycle).

observed in Fig. 6. The CO₂ release is likely due to the presence of weakly bound CO₂ or the decomposition of carbonate sites far away from the Ru catalytic sites. Another CO2 release

mechanism can be triggered by the local increase in temperature due to the reduction of oxidised surface species.³⁸

The Ru precursor played a major role in the DFMs' performance considering both the CH₄ production and the release of unwanted products like CO₂ and CO. The RuM DFMs showed higher amounts of CO₂ and CO released at the beginning of the methanation step, and the transient evolution reached the maximum CH₄ production only after both CO and CO₂ release decreased to zero. In addition, the higher the milling intensity, the slower the CH₄ production rate. On the other side, the samples based on RuAc released very small amounts of CO and CO2 and the CH4 production was rapid (see Table S2†), characterised by a quick production of CH₄ at the beginning of the methanation step. Similar to the RuM DFMs, by increasing the milling intensity the transients flattened, and the profile showed a longer tail, suggesting the presence of slower active sites formed at higher milling regimes in line with the CH₄ profiles during the H2-TPR in Fig. 4.

The cumulative CH₄ production during the 5th methanation cycle is shown in Fig. S10† and is helpful to highlight both the total CH₄ production and the kinetics of the methane release. It is possible to appreciate the separation of the two groups based on the Ru precursor, where RuAc samples not only produced more CH₄ but also showed the steepest increase, as highlighted by the difference in the slope at the beginning of the methanation step. Indeed, it is possible to estimate the initial



Transient gas profiles of CO, CO₂ and CH₄ during the 5th methanation step (350 °C, 10% H₂/N₂, 50 mL min⁻¹, and 250 mg).

methanation release rate by integrating the amount of CH_4 produced in the first 60 s of the 5th methanation step, considering that the cumulative release of CH_4 is almost linear.

In addition to the most evident results, statistical analysis of the experimental design results was carried out to reveal additional correlations between milling parameters, DFM precursors, and performance. Several performance indicators (CO₂ capture capacity, total CH₄ production, CO₂ conversion, CH₄ selectivity, CH₄ release rate, CO release during the capture step, and CO₂ release during the methanation step) were analysed to evaluate the influence of each parameter. Details of the catalytic DFMs' performance are reported in full in Table S2,† while the statistical analysis of the experimental design results is reported as ANOVA tables in Tables S4–10.† The most influential milling parameter for each performance indicator is the one with the lowest p-value.

From the statistical analysis, it clearly appears that the $\rm CO_2$ capture capacity was mostly influenced by the milling intensity: the p-value associated with the milling intensity is 0.05, while the Na and Ru precursor choice showed p-values of 0.27 and 0.17 respectively (see Table S4†), suggesting a limited effect of the precursors on the capture capacity.

To better understand the effect of each parameter on the capture capacity, for each sample the captured ${\rm CO_2}$ amount was

grouped and plotted against the different milling parameter values, as reported in Fig. 7a. These plots are built by plotting the median value of a given response versus a given factor at different levels. In other words, the CO2 capture capacity of all the samples is reported grouped for each milling parameter depending on the parameter's values. Therefore, the effect of the milling intensity on the CO₂ capture capacity appears very clear: the higher the milling intensity, the higher the capture capacity. Regarding the Na precursor, it seems that on average the Na2CO3-based DFMs performed slightly better than the NaNO₃ ones, although the wide distribution of values around the median limits the significance. A similar conclusion can be drawn for the effect of the Ru precursor. In Fig. 7a, the response values at different milling intensities are also grouped by the Ru precursor; in fact, the positive effect of increasing the milling intensity was mostly associated with the combination of higher intensity and the RuAc precursor. This is also evident from the response values in the Ru precursor plot, where the values are grouped by milling intensity.

On the other hand, the average CH₄ production was solely affected by the choice of Ru precursor; in particular, high CH₄ production was obtained for the samples based on RuAc. In Table S5† the ANOVA table is reported: the Ru precursor *p*-value is the smallest at 0.08, and the other *p*-values are much higher,

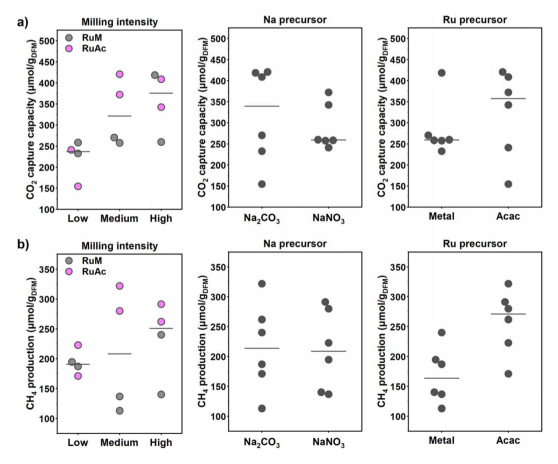


Fig. 7 Main effect plots of the (a) CO_2 capture capacity and (b) average CH_4 production at the different milling parameters. Each dot represents the response of one sample prepared with the corresponding milling parameter value.

indicating no significant effect on the methane production. The main effect plots in Fig. 7b show the positive effect related to the use of RuAc, while the type of Na precursor clearly had no effect on the production of CH4. Interestingly, there was an interaction between the milling intensity and the Ru precursor, with RuM samples showing a negative interaction upon increasing the milling intensity and the RuAc samples instead displaying a positive interaction.

Other relevant metrics to evaluate the DFMs' performance are the CO2 conversion, the CH4 selectivity and release rate, the CO produced during the capture step and the CO₂ released during the methanation (see Table S2†).

The median CO₂ conversion for the RuM samples was 61% and for the RuAc DFMs it was 86%, showing a higher ability of the latter to convert the CO₂ captured in the previous steps (Fig. S11†). The most influential parameter was the Ru precursor with a p-value of 0.07, followed by the milling intensity with a p-value of 0.11, associated with a higher CO₂ conversion at lower milling intensities (Table S6†). For both RuM and RuAc samples, there was no difference between the subgroups of samples based on Na₂CO₃ and NaNO₃ on average, indicating that the conversion was only affected by the Ru active sites. Thus, the spillover of the adsorbed CO2 on different Na sorbent sites formed from different Na precursors is not a limiting factor. Higher CO2 conversions were achieved at the lowest milling regimes, although this was mostly due to the lower CO_2 capture capacity, i.e., a lower amount of reacting CO_2 , as shown in Fig. 7a. The sample 5-RuAcNaC/MI386-15 stood out as an outlier with a CO₂ conversion of 115%: this was probably due to the conversion of residual Na₂CO₃ precursor species, which were not decomposed during the DFM activation.

Similar results were obtained for the CH₄ selectivity, considering both the production of CO as a byproduct and the release of CO2 from weak basic sites or from sorbent sites that might be too far from the Ru active sites to be converted. The only parameter with a low p-value of 0.11 was the Ru precursor (Table S7†). The RuAc DFMs showed higher selectivity with an average of 95.6% of CH₄, 1.0% of CO and 3.4% of CO₂ released at the beginning of the methanation step. No differences were observed between the Na precursors, and a slightly higher selectivity towards methane was associated with lower milling intensity. The CH₄ selectivity is closely related to the CO₂ released during the methanation step. A higher amount of CO₂ was released for the samples based on RuM, and considering the CO2 captured, the RuM samples lost about 5-10% of the CO₂ adsorbed, while the RuAc DFMs lost just 3% of it. This was probably linked to a better dispersion of the Ru active sites which covered more efficiently the surface of the DFMs intercepting the release of CO₂ from the sorbent sites.

Regarding the CH₄ release rate, high values are important because faster DFMs can allow for shorter methanation steps and more efficient capture-conversion cycles. The related main effect plots are shown in Fig. S12.† The RuAc samples were twice as fast compared to the RuM DFMs, with a median of 38 and 18 $\mu mol~CH_4~g_{\rm DFM}^{-1}~min^{-1},~respectively.$ Furthermore, faster DFMs were obtained by milling at lower intensities, although this effect is only related to RuAc DFMs. This can be clearly seen

in Fig. 6 with the shape of the CH₄ transient that flattens in the RuM samples and by increasing the milling regimes. The best performing samples were the ones prepared with RuAc and NaC (5-RuAcNaC/MI386-15, 7-RuAcNaC/MI386-15 and 11-RuAcNaC/ MI386-27); the first one, despite producing the lowest amount of CH₄ for the RuAc samples (170.8 μ mol g_{DFM}^{-1}), released 50% of the total CH₄ in the first minute.

The last metric to consider is the unwanted production of CO during the CO2 capture step. This phenomenon is often linked to the presence of reducing species (either H2 or oxygen vacancies, O_v) on the DFM's surface coming from the prior reduction step. A low amount of CO is desirable, as this molecule is a pollutant itself. Statistically, low significance was obtained for this metric for each milling parameter (Table S9†), although the smallest p-value of 0.20 was associated with both the Ru precursor and the two-way interaction between Na and Ru precursors. Indeed, by looking at Fig. S13,† the use of RuAc resulted in lower amounts of CO during the capture step compared to the use of RuM. As the ANOVA analysis suggested, there is a combination between Ru and Na precursors, in particular coupling RuAc and Na₂CO₃, resulting in 25% less CO release compared to samples based on RuAc and NaNO3, while no differences are observed in the RuM samples.

Overall, the best performing DFM was the sample 11-RuAcNaC/MI386-27. This sample showed both the highest CO₂ capture capacity and CH4 production (respectively 420.6 and 321.8 μ mol g_{DFM}^{-1}), with a rather fast and sustained CH₄ production rate. The combination of RuAc and Na₂CO₃ also resulted in lower amounts of CO produced during the capture phase and the least CO₂ lost at the beginning of the methanation step.

To compare the overall performance of the DFMs studied here, radar plots summarising all main activity indicators are represented in Fig. 8, where a larger area corresponds to better global performance. The F and G axes, corresponding to the CO and CO2 releases, are inverted, as lower values are preferred. The green dashed line represents the sample 11-RuAcNaC/ MI386-27 as a comparison with the other DFM subgroups.

Post-test characterisation and activity correlation

The textural and morphological properties of the post-reaction samples were analysed in order to understand the most important factors leading to the differences in reactivity observed during ICCU-MET tests and thus their correlation with the milling parameters.

First, BET and XRD analyses were carried out to investigate changes in the DFM structure. The evolution of the surface area and the pore structure at different stages of the study is reported in Table 3 for the samples based on RuAc, which are also considered representative of the ones prepared with RuM. As previously observed (Table S1†), with increasing milling intensity a lower surface area was obtained for the samples based on both Na₂CO₃ and NaNO₃ due to the structural modifications induced by stronger mechanical forces. After reduction, the surface area of 5-RuAcNaC/MI386-15 remained unchanged, while it decreased for the two Na2CO3-based DFMs milled at

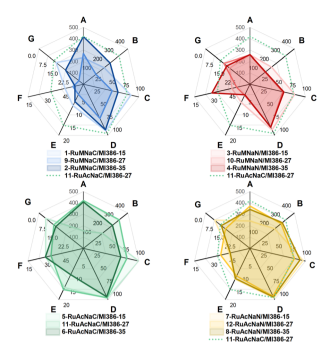


Fig. 8 Radar plots showing the performance indicators used to evaluate the DFMs. A = CO_2 capture capacity (μ mol g_{DFM}^{-1}), B = CH_4 production (μ mol g_{DFM}^{-1}), C = CO_2 conversion (%), D = CH_4 selectivity (%), E = CH_4 release rate (μ mol g_{DFM}^{-1} min⁻¹), F = CO released during the capture (μ mol g_{DFM}^{-1}), and G = CO_2 released during the methanation step. The green dashed line represents the sample 11-RuAc-NaC/MI386-27 as a comparison with the other DFM subgroups.

Table 3 BET specific surface area values ($\rm m^2~g_{DFM}^{-1}$) of the RuAc-based DFMs at the indicated experiment steps: as-prepared (directly after milling, no further treatments), reduced (after 10% $\rm H_2/N_2$ treatment at 400 °C for 1 h), and spent (after 5 cycles of ICCU-MET)

Sample	As-prepared ^a	Reduced	Spent
5-RuAcNaC/MI386-15	138	139	137
11-RuAcNaC/MI386-27	120	102	80
6-RuAcNaC/MI386-35	73	70	61
7-RuAcNaN/MI386-15	120	138	112
12-RuAcNaN/MI386-27	77	89	72
8-RuAcNaN/MI386-35	34	78	65

^a reported BET values of the NaC/MI386 and NaN/MI386 samples, to avoid RuAc decomposition during the outgassing pretreatment.

higher intensities (samples 11-RuAcNaC/MI386-27 and 6-RuAcNaC/MI386-35). This can be explained by the lower degree of $\mathrm{Na_2CO_3}$ decomposition observed at the low milling intensity, which resulted in smaller modifications of the sample. Regarding the NaNO₃ samples, for all the DFMs the area increased after the reduction treatment, likely due to the increase in the size and volume of the pores resulting from the evolution of NO_x species. After ICCU-cycling tests, all the spent DFMs showed a lower surface area compared to the activated (reduced) DFMs, probably due to the dynamic surface reorganisation occurring under switching oxidising/reducing atmospheres.

It is interesting to notice that the samples with higher CO₂ capture capacity, namely those milled at higher intensities, exhibited lower surface areas, suggesting that the chemical interaction between Na and the Al₂O₃ surface plays a key role in the CO₂ capture rather than the surface area itself. Furthermore, the decrease in surface area can partially explain the slower reactivity of the Ru species observed with increasing milling intensity, as the Ru species are probably less dispersed.

Thus, XRD analysis was carried out to investigate the structure of the spent DFMs, including the sorbent, support, and ruthenium sites. The full diffractograms are reported in Fig. S14,† while the detailed 35–50° 2θ range is reported in Fig. 9. Compared with the XRD of the as-prepared DFMs in Fig. 2 and S4,† in general for the Na₂CO₃-based DFMs the intensity of the Na₂CO₃ pattern decreased after the ICCU-MET cyclic testing, especially for the samples milled at mediumhigh intensity, while the two samples milled at the lowest intensity still showed a similar intensity compared to the asprepared ones. This is in agreement with the TGA results shown in Fig. 4 where the samples milled at the lowest intensity experienced a lower weight loss at 400 °C in H2, possibly due to an insufficient mechanical activation and decomposition of the Na₂CO₃ precursor. Furthermore, the Na₂CO₃ species observed in the spent NaC-DFMs correspond to natrite, a monoclinic dehydrated form of Na₂CO₃ (PDF: 00-037-0451). The samples based on NaNO₃ instead showed the complete decomposition of the nitrate precursor and the appearance of a weak peak at 20.5°, which could be attributed to the formation of NaAlO_r species, although other peaks could not be clearly detected at these concentrations. For some NaN-DFMs, it was also possible

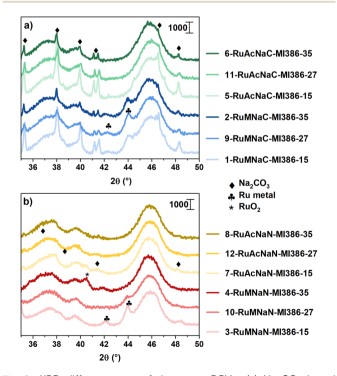


Fig. 9 XRD diffractograms of the spent DFMs: (a) Na_2CO_3 -based DFMs and (b) $NaNO_3$ -based DFMs.

to see weak peaks corresponding to residual Na₂CO₃ formed during the CO₂ capture.

Considering the Ru species, on the as-prepared samples no metallic Ru or RuO2 reflections could be identified, inferring the presence of highly dispersed Ru species (Fig. 2). After the ICCU-MET testing, all six RuAc samples did not show any Rurelated phases suggesting that they maintained a highly dispersed Ru phase. The samples based on RuM, instead, showed a clear peak centred at 44.0°, which corresponds to the main peak of metallic Ru (PDF: 01-088-1734). The Ru crystallite size, estimated with the Scherrer equation after the deconvolution of the complex peak structure (see Fig. S15†), ranged from 11 to 17 nm. Within this subgroup, no correlation was found between the Ru crystallite and the performance of the DFMs (CO production and CH₄ selectivity). However, the absence of the Ru metal peak in the as-prepared DFMs and the rather large crystallite size identified after the ICCU-MET testing suggest that the RuM precursor is not stable on the surface of the DFMs, regardless of the milling intensity, leading to the sintering of the Ru active sites. This could explain the better performance of the RuAc DFMs, which in turn showed high Ru dispersion even after the ICCU-MET tests. The high Ru dispersion coupled with the enhanced CO2 sorption capacity of Na₂CO₃-based samples is likely the key for the peak in activity exhibited by RuAcNaC/MI386 samples, especially when milled under intermediate energy conditions (sample 11).

Conclusions

In this work, the mechanochemical preparation of RuNa/Al₂O₃ dual-function materials for integrated CO2 capture and conversion to CH4 was investigated based on a full factorial design, with the aim of rationally analysing the effect of different milling parameters, such as the milling intensity and the choice of Ru and Na precursors, on the DFM performance.

Seven performance indicators were chosen as representatives of the complex DFM behaviour, i.e., the CO2 capture capacity, the CH₄ production, the CH₄ release rate and selectivity, and the CO and CO2 release during the CO2 capture and the methanation step. The statistical analysis of the obtained results highlighted that all the milling parameters affected the DFMs' properties to a certain extent, but the strongest effects were observed specifically for the milling intensity on CO2 capture capacity and Na and Ru precursors on methanation.

In fact, higher milling intensities were associated with improved CO₂ capture capacity by creating stronger chemical interactions between Na and Al2O3, while the combination of Na₂CO₃ and Ru acetylacetonate resulted in faster DFMs with higher CH₄ production and selectivity and a decreased CO release during the CO2 capture step. The use of RuAc also resulted in smaller and more stable Ru nanoparticles, likely contributing to the improved CO2 conversion performance compared to the Ru metal precursor. It clearly appears that an interplay between all the components of the DFMs is crucial, with a balance between the easier Na2CO3 formation and decomposition, achieved at medium-high milling regimes, and the formation of rapid and active Ru sites obtained under

milder milling conditions. The DoE approach allowed the isolation of the main effects of the chosen milling parameters on optimal DFM performance, paving the way for the rational preparation of DFMs by mechanochemistry. Nonetheless, further characterisation will be needed to understand in detail the structure and reactivity of these DFMs, as the complex structure is not easily characterised and the structure-activity relationships are not straightforward, as evinced by the fact that higher CO2 capture is associated with a lower surface area.

Data availability

Data for this article, including original data for all main and supplementary figures, are available at Zenodo at https:// doi.org/10.5281/zenodo.14415918.

Author contributions

A. B.: conceptualization; investigation; methodology; data curation; formal analysis; validation; visualization; writing original draft. M. D.: conceptualization; investigation; methodology; validation, visualization; supervision; writing - review & editing. S. C.: funding acquisition; resources; supervision; writing - review & editing. A. T.: funding acquisition; resources; supervision; project administration; writing - review & editing.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This study was carried out within the Interconnected Nord-Est Innovation Ecosystem (iNEST) and received funding from the European Union Next-GenerationEU (PIANO NAZIONALE DI RIPRESA E RESILIENZA (PNRR) - MISSIONE 4 COMPONENTE 2, INVESTIMENTO 1.5 - D.D. 1058 23/06/2022, ECS00000043, CUP: G23C22001130006). M.D. is grateful for funding under the REACT EU Italian PON 2014-2020 Program - Action IV.4 Innovation (DM 1062, 10/08/2021, CUP G51B21006040007).

Notes and references

- 1 W. Gao, S. Liang, R. Wang, Q. Jiang, Y. Zhang, Q. Zheng, B. Xie, C. Y. Toe, X. Zhu, J. Wang, L. Huang, Y. Gao, Z. Wang, C. Jo, Q. Wang, L. Wang, Y. Liu, B. Louis, J. Scott, A. C. Roger, R. Amal, H. He and S. E. Park, Chem. Soc. Rev., 2020, 49, 8584-8686.
- 2 I. S. Omodolor, H. O. Otor, J. A. Andonegui, B. J. Allen and A. C. Alba-Rubio, Ind. Eng. Chem. Res., 2020, 59, 17612-17631.
- 3 X. G. Zhang, Buthiyappan, A. J. Jewaratnam, H. S. C. Metselaar and A. A. Raman, J. Environ. Chem. Eng., 2024, 12, 111799.
- 4 D. J. Heldebrant, J. Kothandaraman, N. Mac Dowell and L. Brickett, Chem. Sci., 2022, 13, 6445-6456.

- 5 Y. Hu, W. Liu, H. Chen, Z. Zhou, W. Wang, J. Sun, X. Yang, X. Li and M. Xu, Fuel, 2016, 181, 199–206.
- 6 S. Sun, H. Sun, P. T. Williams and C. Wu, *Sustainable Energy Fuels*, 2021, 5, 4546–4559.
- 7 P. Melo Bravo and D. P. Debecker, *Waste Disposal Sustainable Energy*, 2019, 1, 53–65.
- 8 A. Porta, R. Matarrese, C. G. Visconti, L. Castoldi and L. Lietti, *Ind. Eng. Chem. Res.*, 2021, **60**, 6706–6718.
- 9 Y. Zhang, S. Zhao, L. Li, J. Feng, K. Li, Z. Huang and H. Lin, Catal. Sci. Technol., 2024, 14, 790–819.
- 10 L. P. Merkouri, T. Ramirez Reina and M. S. Duyar, *Nanoscale*, 2022, 14, 12620–12637.
- 11 A. Bermejo-López, B. Pereda-Ayo, J. A. González-Marcos and J. R. González-Velasco, Sustainable Energy Fuels, 2021, 5, 1194–1210
- 12 A. Bermejo-López, B. Pereda-Ayo, J. A. Onrubia-Calvo, J. A. González-Marcos and J. R. González-Velasco, *J. CO2 Util.*, 2022, 58, 1–11.
- 13 S. Cimino, E. M. Cepollaro and L. Lisi, Appl. Catal., B, 2022, 317, 121705.
- 14 M. A. Arellano-Treviño, N. Kanani, C. W. Jeong-Potter and R. J. Farrauto, *Chem. Eng. J.*, 2019, 375, 121953.
- 15 K. J. Ardila-Fierro and J. G. Hernández, ChemSusChem, 2021, 14, 2145–2162.
- 16 P. Baláž, M. Achimovicová, M. Baláž, P. Billik, C. Z. Zara, J. M. Criado, F. Delogu, E. Dutková, E. Gaffet, F. J. Gotor, R. Kumar, I. Mitov, T. Rojac, M. Senna, A. Streletskii and W. C. Krystyna, *Chem. Soc. Rev.*, 2013, 42, 7571–7637.
- 17 E. Colacino, V. Isoni, D. Crawford and F. García, *Trends Chem.*, 2021, 3, 335-339.
- 18 A. P. Amrute, J. De Bellis, M. Felderhoff and F. Schüth, *Chem.–Eur. J.*, 2021, 27, 6819–6847.
- M. Danielis, S. Colussi, C. de Leitenburg, L. Soler, J. Llorca,
 A. Trovarelli, C. de Leitenburg, L. Soler, J. Llorca and
 A. Trovarelli, Angew. Chem., Int. Ed., 2018, 57, 10212–10216.
- 20 A. Braga, M. Armengol-Profitós, L. Pascua-Solé, X. Vendrell, L. Soler, I. Serrano, I. J. Villar-Garcia, V. Pérez-Dieste, N. J. Divins and J. Llorca, ACS Appl. Nano Mater., 2023, 6, 7173-7185.

- 21 N. J. Divins, A. Braga, X. Vendrell, I. Serrano, X. Garcia, L. Soler, I. Lucentini, M. Danielis, A. Mussio, S. Colussi, I. J. Villar-Garcia, C. Escudero, A. Trovarelli and J. Llorca, *Nat. Commun.*, 2022, 13, 1–11.
- 22 J. De Bellis, H. Petersen, J. Ternieden, N. Pfänder, C. Weidenthaler and F. Schüth, Angew. Chem., Int. Ed., 2022, 61, e202208016.
- 23 X. Liu, Y. Li, L. Zeng, X. Li, N. Chen, S. Bai, H. He, Q. Wang and C. Zhang, *Adv. Mater.*, 2022, 34, 1–30.
- 24 L. Yang, Z. Pan and Z. Tian, ChemCatChem, 2024, 16, e202301519.
- 25 M. Danielis, L. P. Merkouri, A. Braga, A. Trovarelli, M. S. Duyar and S. Colussi, *I. CO2 Util.*, 2024, 86, 102895.
- 26 S. Wang, E. T. Schrunk, H. Mahajan and R. J. Farrauto, *Catalysts*, 2017, 7, 1–13.
- 27 H. Liu, L. Cen, X. Xie, L. Liu, Z. Sun and Z. Sun, J. Energy Chem., 2025, 100, 779–791.
- 28 M. T. Dunstan, F. Donat, A. H. Bork, C. P. Grey and C. R. Müller, *Chem. Rev.*, 2021, 121, 12681–12745.
- 29 A. I. Tsiotsias, A. G. Georgiadis, N. D. Charisiou, A. G. S. Hussien, A. A. Dabbawala, K. Polychronopoulou and M. A. Goula, ACS Omega, 2024, 9, 11305–11320.
- 30 M. Danielis, S. Colussi, C. de Leitenburg and A. Trovarelli, *Catal. Commun.*, 2020, **135**, 105899.
- 31 M. Danielis, A. Braga, N. J. Divins, J. Llorca, A. Trovarelli and S. Colussi, *Crystals*, 2023, 13, 1–16.
- 32 R. Leardi, Anal. Chim. Acta, 2009, 652, 161-172.
- 33 M. Gorbounov, J. Taylor, B. Petrovic and S. Masoudi Soltani, S. Afr. J. Chem. Eng., 2022, 41, 111–128.
- 34 A. L. Patterson, Phys. Rev., 1939, 56, 978-982.
- 35 T. S. Nguyen, L. Lefferts, K. B. Saisankargupta and K. Seshan, *ChemCatChem*, 2015, 7, 1833–1840.
- 36 C. J. Keturakis, F. Ni, M. Spicer, M. G. Beaver, H. S. Caram and I. E. Wachs, *ChemSusChem*, 2014, 7, 3459–3466.
- 37 A. Bermejo-López, B. Pereda-Ayo, J. A. González-Marcos and J. R. González-Velasco, *Appl. Catal.*, *B*, 2019, **256**, 117845.
- 38 S. Sun, Y. Zhang, C. Li, Y. Wang, C. Zhang, X. Zhao, H. Sun and C. Wu, *Sep. Purif. Technol.*, 2023, **308**, 122956.