# Reaction **Chemistry &** Engineering

# PAPER

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Cite this: React. Chem. Eng., 2025, 10, 576

Received 2nd October 2024, Accepted 28th November 2024

DOI: 10.1039/d4re00480a

rsc.li/reaction-engineering

## 1. Introduction

Demand for low-cost and abundant fossil energy resources has exacerbated greenhouse gas emissions (GHGs) leading to detrimental impacts on climate and the environment.<sup>1</sup> To mitigate the detrimental impacts of GHG emissions and realize a carbon-neutral future, it is essential to develop novel chemical pathways to produce clean energy carriers such as H<sub>2</sub> while capturing and storing CO<sub>2</sub> emissions.<sup>2,3</sup> As an alternative to conventional multi-step processes to capture, compress, and store CO<sub>2</sub> emissions, intensified approaches that simultaneously inherently capture and store CO<sub>2</sub> emissions in a single step via reactive separation are needed.<sup>4,5</sup> The abundance of renewable energy resources such as biomethane serves as a viable feedstock for H<sub>2</sub> production as an alternative to fossil-derived methane. H<sub>2</sub> has emerged as a preferred energy carrier given the high energy density of  $H_2$  (122 MJ kg<sup>-1</sup>) compared to alternative fossil fuel-derived resources<sup>5</sup> and the absence of CO<sub>2</sub> emissions on combustion.6

# Enhanced H<sub>2</sub> recovery by coupling the water-gas shift reaction with in situ CO<sub>2</sub> capture and mineralization using earth abundant Ca- and Mgsilicates and hydroxides<sup>†</sup>

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Decarbonization of clean energy carriers such as H<sub>2</sub> by coherent integration of multiphase chemical pathways with inherent carbon mineralization is a thermodynamically downhill pathway designed for a sustainable climate, energy, and environmental future. In this effort, a low-temperature water-gas shift reaction (WGSR) with Pt/Al<sub>2</sub>O<sub>3</sub> catalysts is integrated with in situ carbon mineralization in a multiphase reaction environment. The hypothesis that Pt-based catalysts favor selective formation of H<sub>2</sub> over CH<sub>4</sub> has been investigated. H<sub>2</sub> yields increased by 30.8%, 9.5%, 8.3%, and 1.7% in the presence of Ca(OH)<sub>2</sub>, Mg(OH)<sub>2</sub>, Mg<sub>2</sub>SiO<sub>4</sub>, and CaSiO<sub>3</sub> relative to the blank experiment without the sorbent at constant experimental conditions of 250 °C and reaction time of 12 hours in the presence of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst with initial CO and N<sub>2</sub> pressures of 8 bar and 12 bar, respectively. These studies unlock the feasibility of advancing single-step multiphase pathways for enhancing H<sub>2</sub> yields with inherent CO<sub>2</sub> capture and mineralization for a low carbon and sustainable energy and resource future.

> Current commercial H<sub>2</sub> production involves energyintensive hydrocarbon reforming, by which the nonrenewable hydrocarbon fuel is reformed into carbon monoxide (CO) to release H<sub>2</sub> over metal catalysts.<sup>7</sup> As one of the primary reforming techniques, steam reforming (SR) is widely studied due to its relatively low operating temperature and high purity of the H<sub>2</sub> product.<sup>8</sup> The general reaction for steam reforming can be represented as follows:

$$C_m H_n + m H_2 O = m CO + \left(m + \frac{1}{2}n\right) H_2 \quad \Delta H > 0$$
 (1)

In this expression, when m = 1 and n = 4, the reforming reaction corresponds to steam methane reforming (SMR), which is the most extensively applied approach for industrial H<sub>2</sub> production with thermal efficiencies of up to 85%.<sup>8</sup> SMR is typically coupled with subsequent water-gas shift reaction (WGSR) and methanation to further enhance H<sub>2</sub> yield and purity.9 As an effective supplement to SMR, the WGSR generates additional H2 while simultaneously reducing the concentrations of CO at intermediate temperatures (200-450 °C).10 The literature review by Wagner on catalytic advancements<sup>11</sup> and fundamental insights on kinetic rates and challenges towards water gas shift are pioneering studies focused on optimizing this reaction.<sup>9,12</sup> Thermodynamically, WGSR is favored at low temperatures because it is a reversible exothermic process using steam as the hydrogen source as eqn (2) indicates:



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<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d4re00480a

$$CO + H_2O \rightarrow CO_2 + H_2 \quad \Delta H_{298K} = -41.1 \text{ kJ mol}^{-1}$$
 (2)

However, in the low-temperature regime, WGSR is challenged by sluggish kinetics and low CO conversions. The limiting kinetics of WGSR can be overcome by harnessing catalysts to enhance CO conversion kinetics and yields at low temperatures, for which various noble (Pt, Pd, Ru) and nonmetal (Cu, Zn, Fe, Cr)-based supported catalysts have been reported in precedent literature.<sup>13–15</sup> Especially, Pt-based materials have been reported to be promising metals for WGSR in the medium to low temperature (250–350 °C) regime due to their high stability in oxidizing environments.<sup>16</sup>

Even though steam methane reforming and WGSR are widely deployed for  $H_2$  production, challenges associated with producing high-purity  $H_2$  and achieving high selectivity remain.<sup>17</sup> The competing co-presence of CO and CO<sub>2</sub> along with  $H_2$  from gasification and WGSR at the effluent limits  $H_2$  selectivity and its yield. Consequently, the reactor effluent gases must be purified by the pressure swing adsorption process operating at high pressures (1 MPa) to produce a significant amount of high-purity  $H_2$ .

This challenge necessitates the development of sustainable multiphase chemical pathways for selective and energy-efficient reactive separation of H<sub>2</sub> from CO<sub>2</sub> and CO. Harnessing alkaline materials such as Ca- and Mg-bearing hydroxides and silicates is a thermodynamically favorable pathway to enhance H2 yield. The proposed enhanced WGSR concept works based on the Le Chatelier's principle, in which the reversible gas phase WGSR when integrated with a slurry bearing Ca- and Mg-bearing hydroxides and silicates enhances H<sub>2</sub> yield via in situ CO<sub>2</sub> capture and mineralization.<sup>18</sup> This integrated concept has the potential to significantly improve reaction efficiencies, minimize capital cost, and simplify the process configuration from the twostep conventional catalytic process to a single - step process. This approach also circumvents the thermodynamic limitations of conventional WGSR thereby enhancing the rate of forward reaction for  $H_2$  generation, driving the reaction towards the product side by shifting the equilibrium, and allowing high conversions of CO and steam to  $H_2$  and solid carbonates.

The approach of coupling carbon mineralization as a pathway to capture and crystallize  $CO_2$  emissions as solid carbonates is a significant departure from conventional carbon capture and storage (CCS) technologies in which  $CO_2$  is first captured using solvents, sorbents, and membranes and is then injected into geologic reservoirs for storage.<sup>19–23</sup> The permanence and stability of solid Ca- and Mg-bearing carbonates<sup>24</sup> motivate the coupling of carbon mineralization as a pathway to separate  $CO_2$  emissions and enhance  $H_2$  yields. Table 1 summarizes the state-of-the-art literature reports focused on the enhancement towards WGS performance and H<sub>2</sub> yields.<sup>18,25–33</sup>

Carbon mineralization involves the reaction between silicate, hydroxide, and oxide minerals bearing divalent cations (e.g.,  $Ca^{2+}$ ,  $Mg^{2+}$  ions) and  $CO_2$  to produce the respective carbonates. The negative free energy change of the reaction indicates the spontaneity of the process and the chemical stability of the carbonate products.<sup>34</sup> Apart from the spontaneity of the reaction, carbon mineralization has several advantages. First, the natural abundance of starting materials guarantees easy accessibility and affordable prices. Besides, the alkaline minerals have a large CO2 storage capacity, providing an enormous CO<sub>2</sub> storage potential worldwide. Additionally, the typical products from carbon mineralization, including silica, carbonates, and metal oxides, usually have multiple industrial applications after purification.

For carbon mineralization, the starting materials could be magnesium silicate minerals such as  $(Mg,Fe)_2SiO_4$  (olivine),  $Mg_2SiO_4$  (forsterite), MgO (periclase), CaO (lime), CaSiO\_3 (wollastonite), and Ca<sup>2+</sup>- and Fe<sup>2+</sup>-rich silicates (*e.g.*, larnite). These minerals are known to spontaneously react with atmospheric CO<sub>2</sub> to produce solid carbonates.<sup>35–37</sup> These geoinspired mineralization pathways motivated advances in

Table 1      Comparative table of enhanced WGS with different sorbents and catalytic system	stems
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Feedstock	Sorbents	Catalyst	Temperature [°C]	Pressure [bar]	Sorbents amount [g]	Catalyst amount [g]	H <sub>2</sub> yields [%]	CO <sub>x</sub> yields [%]	Ref.
$CO_2$ , CO, steam,	LDHs	Iron-chromium	400	28	891	434	99.83	0.17%	18
$H_2, N_2$									
CO, steam, Ar	LDHs	Cu/ZnO/Al <sub>2</sub> O <sub>3</sub>	400	1	48.45	48.45	99.99	0.01%	25
$CO$ , steam, $N_2$	CaO/Al <sub>2</sub> O <sub>3</sub>	Ni nanoparticle	400	NA	0.25	0.0075	76-98	NA	26
CO, steam, $N_2$	CaO/Al <sub>2</sub> O <sub>3</sub>	$Pt/Al_2O_3$	350	NA	NA	NA	High purity	NA	27
CO, steam, Ar	CaO/Ca <sub>12</sub> Al <sub>14</sub> O <sub>33</sub>	Fe/Mn	400-700	1	0.88	0.12	88-95	5-12	28
CO, steam	CaO-based	Pd/Ni/Co	425-550	1	20	1	98-99	0.35	29
$CO$ , steam, $H_2$ , He	CaO/NaOH	Pt/ZDC	300-600	1-11	NA	NA	51-100	0-49	30
CO, steam	AMS/MgO/CaCO <sub>3</sub>	$Cu/Ce_{0.6}Zr_{0.4}O_2$	275-375	12	2.625	2.625	86-99	1-14	31
H <sub>2</sub> , CO, steam, N <sub>2</sub>	Na-Mg based	Cu/ZnO/Al <sub>2</sub> O <sub>3</sub>	375	1	5	5	High purity	NA	32
$CO_2, N_2, H_2$	CaO	MO <sub>x</sub> (Al, Ce, Ti, Zr)	600-750	NA	0.2	0.1	NA	NA	33
H <sub>2</sub> , CO, N <sub>2</sub>	Mg/Ca hydroxide and silicate	Pt/Al <sub>2</sub> O <sub>3</sub>	250-300	20	3	3	72.67	27.33	This wor

As an alternative to these hydrothermal routes, direct gas - solid mineralization routes<sup>42,43</sup> are explored but challenged by slow kinetics when Mg- or Ca-bearing silicates are used. In this context, sorbent-enhanced water gas shift reactions (WGSR) in direct gas-solid modes were extensively studied.44-47 Compared with the widely used membrane separation approach to separate CO<sub>2</sub> in WGSR, the sorbent-enhanced WGSR possesses multiple advantages, including more favorable and uniform kinetics, greater material durability, and larger CO2 capture capacity.<sup>30,48</sup> Conventional sorbent-enhanced water-gas shift reactions occur in fluidized bed reactors with steam-saturated CO<sub>2</sub> flow.<sup>32,49</sup> The selected sorbents are typically metal oxides (e.g., CaO) or layered double hydroxide (LDH) (e.g., hydrotalcite) with high CO<sub>2</sub> capture capacity.<sup>18,25,26</sup> However, direct gas-solid carbonation is very kinetically limited and tends to be greatly enhanced with the participation of aqueous media, which dissolves the alkaline resource to release metal cations that capture CO<sub>2</sub> to form carbonate species. 43,50-52

While Ca-bearing oxides have been reported to be effective for enhanced water gas shift reaction in a direct gas-solid reaction mode,<sup>27,29,30</sup> the direct use of earth abundant Mgand Ca-bearing silicates in this mode is significantly challenged by mass transfer limitations and slow kinetics of reaction.<sup>39-41</sup> The direct use of Mg- and Ca-bearing silicates circumvents the need to use additional reagents to produce more reactive Ca- and Mg-bearing hydroxides. To accelerate mass transfer and the kinetics of CO<sub>2</sub> capture using earthabundant Mg- and Ca-bearing silicates, aqueous routes which involve the dissolution of silicates to release Mg<sup>2+</sup> or Ca<sup>2+</sup> ions for capturing CO<sub>2</sub> produced from the WGSR to produce the respective carbonates are investigated in this work.53 Prior studies have reported that the dissolution of silicate minerals facilitates more robust mass transfer and accelerates carbon mineralization.54-56 Prior studies have demonstrated enhanced H<sub>2</sub> production using Mg(OH)<sub>2</sub> for in situ CO<sub>2</sub> capture and mineralization.<sup>57</sup> However, the influence of directly using of Mg- and Ca-bearing silicates for the enhanced water gas shift reaction has not been reported to date. The conventional hypothesis is that silica precipitated as Mg- and Ca-bearing silicates limit mass transfer and thus lower reactivity with CO2 to produce the respective Ca- and Mg-bearing carbonates. However, this hypothesis has not been evaluated in the context of harnessing earth-abundant Ca- and Mg-bearing silicates to enhance the WGSR with in situ CO<sub>2</sub> capture and mineralization, and is therefore, the focus of the proposed investigations.

Several favorable multiphase chemical interactions are necessary for this approach to be feasible. First, the reactivity of CO and steam in the presence of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst needs to be robust in producing CO2 and H2 as the products in sufficient quantities. Second, CO<sub>2</sub> needs to be soluble in the aqueous phase to facilitate the forward WGSR for enhancing H<sub>2</sub> production. Third, the dissolved CO<sub>2</sub> needs to react with Mg- and Ca-bearing silicates to produce the respective carbonates. Fourth, the precipitated silica should not limit mass transfer and the subsequent carbon mineralization behavior. To unlock the full potential of enhanced H<sub>2</sub> production coupled with carbon mineralization, it is essential to identify the rate limiting factor and advance strategies to overcome these limitations. To this end, the multiphase reaction pathways are assembled such that the gas phase catalytic WGSR is coupled with slurry phase carbon mineralization as shown in Fig. 1. The associated CO<sub>2</sub> from WGSR reacts with water to produce carbonate species, which subsequently react with dissolved calcium or magnesium ions obtained from the sorbents for in situ carbon mineralization. To identify and address the factors limiting H<sub>2</sub> production coupled with carbon mineralization, several key research questions are addressed: (i) what are the chemical mechanisms underlying enhanced H<sub>2</sub> conversion with in situ carbon mineralization using Mg- and Ca-bearing

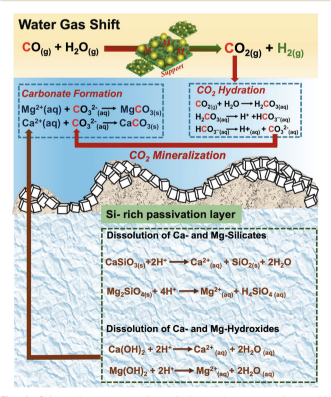


Fig. 1 Schematic representation of the pathway to enhance  $H_2$  conversion by integrating the water gas shift reaction integrated with carbon mineralization of Ca- and Mg-bearing hydroxides and silicates.

hydroxides and silicates? (ii) What is the influence of the silicate vs. hydroxide reactivity on enhancing  $H_2$  production with carbon mineralization? (iii) In this multi-step reaction pathway, what is the rate limiting step and how can this step be accelerated?

Addressing these questions unlock new insights into the mechanisms and the scientific feasibility of enhancing  $H_2$  conversion with *in situ* CO<sub>2</sub> capture and mineralization using Mg- and Ca-bearing silicates and contrasting with that of the respective hydroxides.

## 2. Experimental methods

#### 2.1 Materials and methods

All experiments to investigate enhancement in H<sub>2</sub> conversion with the WGSR coupled with in situ CO2 capture and mineralization are conducted in the presence of platinum on alumina support (Pt/Al<sub>2</sub>O<sub>3</sub>) catalyst.<sup>27</sup> The Pt/Al<sub>2</sub>O<sub>3</sub> catalyst is selected due to its high activity, superior thermal stability, and high product specificity to  $H_2$  and  $CO_2$ .<sup>58-60</sup> Specifically, Pt/Al<sub>2</sub>O<sub>3</sub> catalyst is sized as 3.2 mm pellets with a composition of 0.5 wt% Pt and is procured from Sigma Aldrich, USA. Mg-bearing hydroxide (brucite  $(Mg(OH)_2)$ ) and silicate (forsterite (Mg<sub>2</sub>SiO<sub>4</sub>)) used for in situ CO<sub>2</sub> capture and mineralization are obtained from Fisher Chemical and Xi'An Function Material Group Co. Ltd, respectively. Ca-bearing hydroxide (portlandite (Ca(OH)<sub>2</sub>)) and silicate (wollastonite  $(CaSiO_3)$ ) also used for *in situ* CO<sub>2</sub> capture and mineralization are acquired from Thermo Fischer Scientific Co., USA. These Mg- and Ca-bearing silicates and hydroxides are ball-milled in a bench-top mixer mill (8000 M Mixer/Mill® by SPEX® Sample Prep) to obtain raw materials with comparable mean particle sizes (5.57  $\mu$ m to 10.38  $\mu$ m). The particle sizes of the milled samples are determined using laser diffraction particle size analyzer (Anton Paar). Deionized water (18.2 M $\Omega$  cm, Millipore) is used throughout all the experiments as required. Ultra-high purity CO gas (99.999% purity) is supplied by Airgas Co., USA.

# 2.2 Experimental setup for enhanced WGSR and carbon mineralization

Experiments to investigate enhanced H<sub>2</sub> conversion with in situ CO<sub>2</sub> capture and mineralization are conducted in a 50 mL stainless steel high-pressure stirred reactor (Micro Bench Top Reactor, Parr Instruments Co., USA) in a multiphase reaction environment. A schematic representation of the experimental setup is shown in Fig. 2. The reactor is a stirred cylindrical system with an inner diameter of 33 mm and a height of 57 mm. It features a 4-blade Rushton turbine stirrer with a 20 mm diameter, positioned 9 mm above the bottom along the central axis of the reactor. Given the stirring rate and the dimensions of the stirrer, efficient mixing and mass transfer in the aqueous phase is expected during the reaction. Simultaneously, CO and N2 are introduced through a gas inlet pipe (inner diameter of 3 mm) into the headspace for gas-phase reactions and to build internal pressure. Elevated internal pressures are preferred for these multiphase reactions, as the increased CO partial pressure shifts the equilibrium of the WGSR towards higher H<sub>2</sub> and CO<sub>2</sub> production, enhancing the subsequent dissolution and diffusion of CO2. Catalytic conversion of CO and steam occurs in the gas phase in the presence of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst. The surface area, pore volume, and average pore diameter of  $Pt/Al_2O_3$  catalyst are 163.13 m<sup>2</sup> g<sup>-1</sup>, 0.303 cc g<sup>-1</sup>, and 6.282 nm, respectively (see Table S1 in ESI<sup>+</sup>). Prior to CO gas injection, 3 g (0.5 wt%) of the commercial Pt/Al<sub>2</sub>O<sub>3</sub> catalyst is placed in an aluminum basket and suspended in the

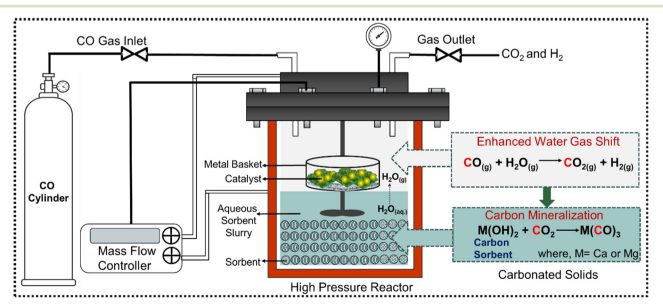


Fig. 2 Schematic representation of the experimental setup for coupling the water gas shift reaction (WGSR) with *in situ*  $CO_2$  capture and mineralization for enhanced H<sub>2</sub> conversion.

headspace of the reactor ensuring the basket is not in contact with the sorbent slurry. CO flow rate from the cylinder is controlled by using a mass flow controller (MFC).  $CO_2$ capture and mineralization occurs in a slurry phase bearing 3 g of Mg- and Ca-bearing hydroxides and silicates, mixed with 10 mL of deionized water. The initial solid-to-liquid weight ratio is set at 3:10, resulting in mole percentages of 8.48% for Mg(OH)<sub>2</sub>, 3.70% for Mg<sub>2</sub>SiO<sub>4</sub>, 6.80% for Ca(OH)<sub>2</sub>, and 4.45% for CaSiO<sub>3</sub>, respectively. The sorbent-to-catalyst ratio is maintained at a consistent weight ratio of 1:1 across all cases.

The headspace of the reactor is purged with CO gas for 15 minutes to evacuate gaseous impurities before it is pressurized and sealed tightly. After reaching the desired pressure set point for CO at 20 bar, the heater is switched on to achieve a setpoint of 250 °C with a constant stirring rate of 300 rpm, signaling the start of the experiment. Steam is generated by the vaporization of water in the slurry. The corresponding partial pressure of steam is determined by the temperature in the steam table (Table S2.1 and 2.2<sup>+</sup>). The reactor outlet is sealed with heating tape to avoid the possibility of steam condensation and is connected directly to a gas chromatograph to analyze the concentrations of gaseous products. The pH of the slurry increased to 9.95 for Mg(OH)<sub>2</sub>, 8.69 for Mg<sub>2</sub>SiO<sub>4</sub>, 12.37 for Ca(OH)<sub>2</sub>, and 8.98 for CaSiO<sub>3</sub>, respectively. The experiments are performed at different reaction times ranging from 3 to 12 hours. After the completion of the reaction, the reactor is cooled down to ambient temperature. Following this, the outlet valve is slowly opened to purge the gaseous products in the micro-GC and the carbonate-bearing solid products are collected for further analysis and characterization. Additionally, enhancement in H<sub>2</sub> yield resulting from CO<sub>2</sub> capture and mineralization is investigated as a function of reaction time and the alkaline sorbent type.

Additional experiments are conducted to determine the exact moles of each gas in the reactor. The experimental setup is the same as described previously. In this approach, 8 bar of CO is injected into the reactor with 12 bar of  $N_2$ .  $N_2$  is used as a reference due to its inert nature and unchanged pressure over the course of the reaction. This approach enables the calculation of the moles for the other gaseous components based on the initial  $N_2$  content. The temperature is increased to a setpoint of 300 °C, and the reaction is performed for 12 hours, while keeping other reaction conditions unchanged, to investigate the influence of Mgand Ca-bearing hydroxides and silicates on enhancing  $H_2$ conversions.

#### 2.3 Product analysis and characterization

After the reaction, the compositions of effluent gases are analyzed by micro gas chromatography (GC) with the calibration data shown in Fig. S1.† The changes in the composition and structure of the Mg- and Ca-bearing hydroxides and silicates are investigated using infrared (IR) spectroscopy collected in an attenuated total reflection (ATR) mode using an attenuated total reflection-Fourier transform infrared spectrometer (ATR-FTIR, Nicolet™ iS50, Waltham, MA), and X-ray diffractometry (XRD, Bruker D8 Advance ECO powder diffractometer, Bruker), respectively. Furthermore, the morphology and particle sizes are determined using the field emission scanning electron microscope (FESEM, LEO 1550 FESEM, Bruker) and particle size analyzer (Anton Paar). The surface area and the pore size distributions of the sorbents before and after the experiments were determined by N<sub>2</sub> adsorption-desorption isotherms using the Brunauer-Emmett-Teller technique (BET) (Quantachrome Autosorb iQ Analyzer, Boynton Beach, FL) with the BJH model. Prior to measuring the adsorption-desorption isotherms, the samples were outgassed at 140 °C for 12 hours. A ±5% BET instrument error was accounted for all the analysed samples. The concentration of formate in the aqueous phase was measured using a 500 MHz <sup>1</sup>H NMR spectrometer (Bruker AVIII 500) using a solvent suppression method. The calibration line for formate concentration and its corresponding slope, determined using DMSO as internal standards, are shown in Fig. S2.<sup>†</sup>

### 3. Results and discussion

# 3.1 Reaction pathways involved in coupled WGSR with *in situ* CO<sub>2</sub> capture and mineralization for enhanced H<sub>2</sub> conversion

The multiphase reactions involved in coupling the WGSR with *in situ* CO<sub>2</sub> capture and mineralization for enhanced H<sub>2</sub> conversion include: (i) low-temperature conversion of CO and steam over Pt/Al<sub>2</sub>O<sub>3</sub> catalyst to produce CO<sub>2</sub> and H<sub>2</sub>; (ii) dissolution of Mg- and Ca-bearing hydroxides and silicates to release Mg<sup>2+</sup> and Ca<sup>2+</sup> ions, respectively and promote alkalinity; (iii) dissolution of the produced gaseous CO<sub>2</sub> in the alkaline aqueous phase; (iv) reactivity of the hydrated CO<sub>2</sub> with Mg<sup>2+</sup> and Ca<sup>2+</sup> ions to mineralize into Mg and Ca-carbonates; and (v) generation of high-purity H<sub>2</sub> due to equilibrium shift due to CO<sub>2</sub> capture and mineralization.

The motivation for using Pt/Al<sub>2</sub>O<sub>3</sub> catalyst stems from the need to direct the formation of H<sub>2</sub> as opposed to CH<sub>4</sub>.<sup>61</sup> Prior analogous studies showed that Pt or Ni/Al2O3 catalysts are effective in reforming aqueous biomass oxygenates such as ethylene glycol and glycerol to produce  $H_2$  and  $CO_2$ preferentially over CH<sub>4</sub>.<sup>62</sup> Therefore, in this study, Pt/Al<sub>2</sub>O<sub>3</sub> catalysts are used to lower the activation energy barrier, accelerate reaction kinetics, suppress the formation of CH<sub>4</sub>, and promote the formation of H<sub>2</sub> and CO<sub>2</sub>. Furthermore, the in situ capture and conversion of gaseous CO<sub>2</sub> in the presence of alkaline Mg- and Ca-bearing hydroxides and silicates to produce the respective carbonates favors the forward equilibrium shift towards more CO conversion and H<sub>2</sub>. Dissolution of Mg - or Ca - bearing hydroxides and silicates releases Mg<sup>2+</sup> and Ca<sup>2+</sup> cations which capture and solubilize CO<sub>2</sub> to produce the respective solid carbonates. For example, in this work, the pH of the alkaline slurry is observed to be in the range of 8.69-12.37 for the Mg- and Ca-bearing hydroxides and silicates. This pH is ideal to capture and solubilize  $CO_2$  from the gas phase to produce the respective (bi)carbonate species.

To probe the mechanisms underlying enhanced  $H_2$ conversion by coupling the WGSR with in situ CO2 capture and mineralization, the first set of experiments are conducted by injecting CO gas. Steam required for the WGSR is provided by the slurry. The ratio of CO to steam is calculated and quantified from the steam table (Table S2.1 and 2.2, ESI<sup>†</sup>). In this approach, the effluent gas compositions are measured using micro gas chromatography (GC), and the molar percentage of each phase is normalized by the total amount of H<sub>2</sub>, CO, and CO<sub>2</sub>. This approach provides insights into the relative abundance of the gas phase compositions. In this scenario, the relative gas phase composition of H<sub>2</sub> can be elevated if CO<sub>2</sub> is absorbed by the liquid phase, even if CO conversion is unchanged. Therefore, CO conversion is determined for quantifying the enhancement in WGSR using the following expression:

$$R_{\rm CO} = \frac{M\%_{\rm H_2}}{M\%_{\rm H_2} + M\%_{\rm CO}} \tag{3}$$

In this expression,  $R_{\rm CO}$  is the conversion of CO,  $M\%_{\rm H_2}$ , and  $M\%_{\rm CO}$  are the mole percentages of H<sub>2</sub> and CO in the measured gas samples, respectively. In this equation, the mole percentage of H<sub>2</sub> is applied as the substitute for the mole percentage of CO<sub>2</sub> because a large amount of generated CO<sub>2</sub> partitions into the aqueous phase as the reactant for carbon mineralization, making it inaccurate to quantify CO<sub>2</sub> only based on the gas phase compositions.

Additionally, the exact number of moles of each gas is necessary to determine the theoretical maximum yield of  $H_2$  and gaseous  $CO_2$  compositions. To this end, additional experiments are conducted with a moderate amount of  $N_2$  in the system as the baseline. In this approach, the absolute number of moles of  $N_2$  is constant throughout the experiment.

This approach enables the determination of the exact moles of  $H_2$ , CO, and  $CO_2$  by the van der Waals equation (as shown below) from the normalized mole percentages after the multi-step reactions.

$$\left(P + \frac{n^2 \cdot a}{V^2}\right)(V - n \cdot b) = n \cdot R \cdot T \tag{4}$$

In the expression above, P = pressure, V = volume, a, b = van der Waal's constants, n = number of moles, R = universal gas constant, T = temperature.

# 3.2 Thermodynamic considerations for enhanced $H_2$ conversion by coupling the WGSR with *in situ* CO<sub>2</sub> capture and mineralization

The reaction conditions for enhanced  $H_2$  conversion by coupling the WGSR with *in situ*  $CO_2$  capture and mineralization is equilibrium-limited and dictated by the thermodynamics for WGSR to form  $CO_2$  and  $H_2$  and  $CO_2$  capture and mineralization to form Mg- and Ca-carbonates.<sup>48</sup> The detailed reactions associated with coupling the WGSR with *in situ* CO<sub>2</sub> capture and mineralization are below:

$$Mg(OH)_{2(s)} + CO_{(g)} + H_2O_{(g)} \rightarrow MgCO_{3(s)} + H_{2(g)} + H_2O_{(l)}$$
 (5)

$$Mg_{2}SiO_{4(s)} + 2CO_{(g)} + 2H_{2}O_{(g)} \rightarrow 2MgCO_{3(s)} + SiO_{2(s)} + 2H_{2(g)} \quad (6)$$

$$Ca(OH)_{2(s)} + CO_{(g)} + H_2O_{(g)} \rightarrow CaCO_{3(s)} + H_{2(g)} + H_2O_{(l)}$$
 (7)

$$CaSiO_{3(s)} + CO_{(g)} + H_2O_{(g)} \rightarrow CaCO_{3(s)} + SiO_{2(s)} + H_{2(g)}$$
 (8)

The overall reaction functions reveal the stoichiometric ratio of the reactants and products, indicating a correlation between equilibrium constants ( $K_{eq}$ ) and gas species partial pressure (P) at the equilibrium state, respectively:

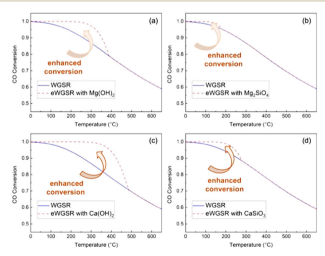
$$K_{eq}(Mg(OH)_2) = \frac{P_{H_2}}{P_{CO} \times P_{H_2O}}$$
(9)

$$K_{\rm eq}({\rm Mg}_{2}{\rm SiO}_{4}) = \frac{P_{{\rm H}_{2}}^{2}}{P_{\rm CO}^{2} \times P_{{\rm H}_{2}{\rm O}}^{2}}$$
(10)

$$K_{\rm eq} \left( {\rm Ca}({\rm OH})_2 \right) = \frac{P_{\rm H_2}}{P_{\rm CO} \times P_{\rm H_2O}}$$
(11)

$$K_{\rm eq}({\rm CaSiO}_3) = \frac{P_{\rm H_2}}{P_{\rm CO} \times P_{\rm H_2O}}$$
(12)

The concentration of the different components involved in enhancing  $H_2$  conversion coupled with *in situ* CO<sub>2</sub> capture and mineralization can be determined by their equilibrium constants using thermodynamic properties and relationships as shown by the equation below:<sup>63</sup>



**Fig. 3** Enhancement in WGSR as determined from CO conversions based on thermodynamic evaluations in gas-solid reaction mode using (a) Mg(OH)<sub>2</sub>, (b) Mg<sub>2</sub>SiO<sub>4</sub>, (c) Ca(OH)<sub>2</sub>, and (d) CaSiO<sub>3</sub> for the *in situ* CO<sub>2</sub> capture. The initial steam-to-CO ratio is 1:1 with adequate sorbents for carbon mineralization. Results for Mg(OH)<sub>2</sub> agree with prior studies.<sup>64</sup>

$$K_{\rm eq} = \exp\left(\frac{5693.5}{T} + 1.077\ln T + 5.44 \times 10^{-4} T - 1.125 \times 10^{-7} T^2 - \frac{49170}{T^2} - 13.148\right)$$
(13)

The equilibrium constants determined using expression (13) are used to calculate the theoretical CO conversions as a function of temperature using expressions (9)–(12). Thermodynamic simulations with steam-to-CO ratio of 1:1 are used to determine theoretical CO conversions using Mg(OH)<sub>2</sub>, Mg<sub>2</sub>SiO<sub>4</sub>, Ca(OH)<sub>2</sub>, and CaSiO<sub>3</sub> as alkaline sources (Fig. 3).

Notably, the WGSR and enhanced WGSR (or eWGSR) conversions define the theoretical percentage of CO converted to  $CO_2$  at the equilibrium.  $CO_2$  conversions are calculated from the partial pressure of  $H_2$ , CO, and steam. The theoretical CO conversion of  $Mg(OH)_2$  confirms prior reported calculations.<sup>64</sup> CO conversions at equilibrium decrease with increasing temperature due to the exothermic nature of the reactions (Fig. 3). Compared with conventional WGSR, a significant enhancement in CO conversion is noted at temperatures below 350 °C, indicating that Mg- and Cabearing hydroxides and silicates are effective in aiding CO (and  $H_2$ ) conversion by capturing  $CO_2$ . It is interesting to note that at temperatures below 350 °C,  $Mg(OH)_2$  and  $Ca(OH)_2$  enhance CO conversion at 250 °C and 300 °C due to *in situ* capture of gaseous  $CO_2$ .

In contrast, the enhancement in CO conversion realized using Mg- and Ca-bearing silicates is achieved below 200 °C. Based on these thermodynamic analyses, it is evident that CO conversion can be enhanced using Ca- and Mg-bearing silicates in gas-solid reaction modes. To overcome the mass transfer limitations associated with gas-solid reactions, multiphase gas-liquid-solid reactions are proposed to accelerate the kinetics of WGSR when coupled with aqueous  $CO_2$  capture and mineralization pathways at temperatures below 300 °C. These experimental conditions are a significant departure from conventional sorbent-enhanced WGSR operating in a gas-solid mode to capture  $CO_2$ .

#### 3.3 Enhancement in WGSR using Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub>

To enhance  $H_2$  conversion with *in situ*  $CO_2$  capture, it is essential to use a suitable catalyst that will convert CO and steam to produce  $H_2$  and  $CO_2$  and an alkaline source to capture  $CO_2$ .<sup>65</sup> Prior studies have reported enhancement in  $H_2$  conversion *via* sorbent – enhanced WGSR using Cabearing sorbents.<sup>46</sup> Enhancement in  $H_2$  yields were also reported by coupling the WGSR with slurry phase  $CO_2$ capture and mineralization using  $Mg(OH)_2$ .<sup>66</sup> However,  $Mg(OH)_2$  needs to be directly mined or extracted from earth – abundant Mg-silicates. Eliminating this additional mining or extraction step and directly using earth – abundant Mgsilicate minerals for enhancing  $H_2$  conversion while capturing  $CO_2$  is less explored but highly transformative. To address this knowledge gap, experiments are conducted to contrast the influence of  $Mg(OH)_2$  *vs.*  $Mg_2SiO_4$  on enhancing  $H_2$  yield with *in situ* CO<sub>2</sub> capture. The abundance of olivine (Mg,Fe)<sub>2</sub>SiO<sub>4</sub> bearing forsterite (Mg<sub>2</sub>SiO<sub>4</sub>) in the natural environment motivates the direct use of these materials for CO<sub>2</sub> capture.<sup>39</sup> To investigate the enhancement in  $H_2$  yield with CO<sub>2</sub> capture, the gaseous products at the effluent are analyzed. Following this, the structures and morphologies of the carbonate-bearing products are characterized separately.

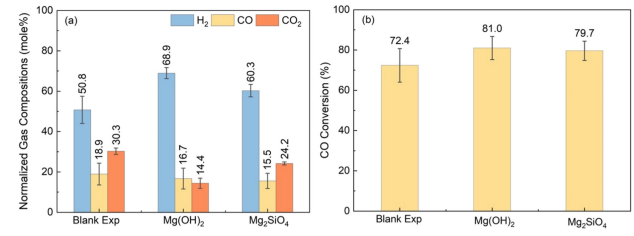
Initially, a blank experiment is performed by considering only the WGSR over Pt/Al<sub>2</sub>O<sub>3</sub> catalyst, to investigate its catalytic activity in the low-temperature regime. Fig. 4(a) represents the gas mole percentages when feeding only 20 bar CO at 250 °C for 9 hours. Fig. 4(b) shows the conversion of CO for the blank experiment and the cases with Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub>. In the absence of the alkaline resource, CO conversion of 72.4% is noted (Fig. 4(b)). Gas analyses did not indicate any CH<sub>4</sub> formation demonstrating that the Pt/Al<sub>2</sub>O<sub>3</sub> catalyst is effective in converting CO and steam to CO<sub>2</sub> and H<sub>2</sub>. The observed higher selectivity towards H<sub>2</sub> using lowtemperature Pt/Al<sub>2</sub>O<sub>3</sub> catalyst is consistent with previous literature on WGSR.<sup>27</sup> Compared with the blank experiment without any alkaline source, WGSR reactions in the presence of a slurry bearing Mg(OH)2 or Mg2SiO4 enhanced CO conversion and H<sub>2</sub> yield and lowered the concentration of gaseous CO2 due to in situ CO2 capture. In contrast to the blank experiment without the alkaline source, CO conversions in the presence of  $Mg(OH)_2$  and  $Mg_2SiO_4$  at 250 °C are 81% and 79.7%, respectively, as seen in Fig. 4(b). The molar compositions of H<sub>2</sub> and CO<sub>2</sub> in the gas phase are 50.8% and 30.3% for H<sub>2</sub> and CO<sub>2</sub> for the blank experiment. In contrast, H<sub>2</sub> concentrations increase by 18% and 9.5%, and CO<sub>2</sub> concentrations decrease by 15.9% and 6.1% in the presence of Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub> compared to the blank experiment (Fig. 4(a)).

The relative composition of H<sub>2</sub> in the gas phase is 8.6% higher when  $Mg(OH)_2$  is used for  $CO_2$  capture as opposed to Mg<sub>2</sub>SiO<sub>4</sub>. CO<sub>2</sub> concentrations are nearly 10% lower when Mg(OH)<sub>2</sub> is used as opposed to Mg<sub>2</sub>SiO<sub>4</sub>. Higher H<sub>2</sub> concentrations and lower CO<sub>2</sub> concentrations in the presence of  $Mg(OH)_2$  versus  $Mg_2SiO_4$  are attributed to the faster dissolution kinetics of Mg(OH)2 relative to Mg2- $SiO_4$ .<sup>67</sup> The enhanced release of  $Mg^{2+}$  ions favors  $CO_2$ capture and the forward WGSR reaction which results in  $H_2$ enhanced conversions and CO consumption (Fig. 4(a) and (b)).

#### 3.4 Enhancement in $H_2$ conversion with *in situ* CO<sub>2</sub> capture as a function of time using Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub> materials

One of the key uncertainties in enhancing  $H_2$  conversions with *in situ*  $CO_2$  capture is the influence of the reactivity of the alkaline resources in hydrothermal environments. In this context, even though it is known that  $Mg(OH)_2$  is



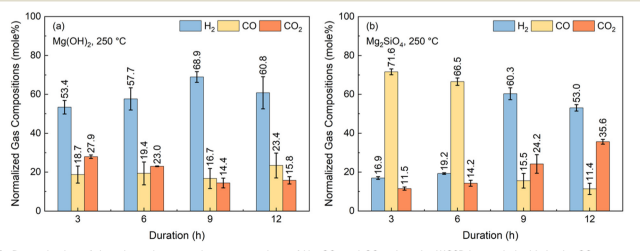


**Fig. 4** Contrasting the influence of  $Mg(OH)_2$  and  $Mg_2SiO_4$  on the water gas shift reaction (WGSR) as determined from  $H_2$ , CO, and  $CO_2$  compositions in (a) and CO conversions in (b). Experiments are conducted at 250 °C, 20 atm for 9 h. The blank experiment is conducted over 3 g of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst without alkaline source. The  $Mg(OH)_2$  and  $Mg_2SiO_4$  cases are conducted over 3 g of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst with a sorbent-to-liquid weight ratio of 3 : 10, leading to mole percentages of 8.48% for  $Mg(OH)_2$  and 3.70% for  $Mg_2SiO_4$ , respectively.

more soluble and has faster kinetics of dissolution compared to  $Mg_2SiO_4$ , the kinetics associated with accelerating WSGR are not well understood. Several factors could be competing in this integrated and enhanced WGSR approach.

First, the gas phase conversion of CO and steam to  $H_2$ and CO<sub>2</sub> can be limiting. Second, the solubility of CO<sub>2</sub> and reactivity of Ca- and Mg-bearing silicates and hydroxide to capture CO<sub>2</sub> can be slow. Based on the changes in the compositions of the gas phase over time, the factors limiting the conversion of CO and H<sub>2</sub>O to H<sub>2</sub> with *in situ* CO<sub>2</sub> capture can be determined. To this end, experiments are conducted with Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub> at 250 °C, 20 atm as a function of reaction time. The normalized gas phase compositions for reactions conducted at 3, 6, 9, and 12 hours are shown in Fig. 5(a) and (b) in the presence of Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub>, respectively.

In the first three hours of the reaction with  $Mg(OH)_2$  as the alkaline source, H<sub>2</sub> composition is 53.4% while that of CO and CO<sub>2</sub> are 18.7 and 27.9%, respectively. An increase in H<sub>2</sub> composition to 68.9% and a decrease in CO and CO<sub>2</sub> compositions to 16.7% and 14.4%, respectively are noted in the 9-hour cases. These results indicate that  $Mg(OH)_2$  is effectively enhancing CO conversions and H<sub>2</sub> yields with inherent CO2 capture. Therefore, the WGSR is not the limiting factor in H<sub>2</sub> conversion in the first nine hours of the reaction. However, increasing the reaction time from 9 hours to 12 hours, lowered H<sub>2</sub> compositions from 68.9% to 60.9%. CO and CO<sub>2</sub> compositions increased from 16.7 to 23.4% and 14.4 to 15.8%, respectively, when the reaction time is increased from 9 hours to 12 hours. These results suggest that CO<sub>2</sub> capture is the likely limiting factor as the reaction time increases from 9 to 12 hours. This observation can be attributed to the fact that at increased reaction times, the



**Fig. 5** Determination of time dependence on the concentrations of  $H_2$ , CO, and CO<sub>2</sub> when the WGSR is coupled with *in situ* CO<sub>2</sub> capture and mineralization using (a) Mg(OH)<sub>2</sub> and (b) Mg<sub>2</sub>SiO<sub>4</sub>. The Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub> cases are conducted at 250 °C and 20 atm over 3 g of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst with a sorbent-to-liquid weight ratio of 3:10, leading to mole percentages of 8.48% for Mg(OH)<sub>2</sub> and 3.70% for Mg<sub>2</sub>SiO<sub>4</sub>, respectively.

availability of  $Mg^{2+}$  ions for  $CO_2$  capture is likely decreasing over time. Slower dissolution rates of  $Mg(OH)_2$  over time due to mass transfer arising from the formation of Mgcarbonates<sup>68</sup> or due to the saturation of the solution with  $Mg^{2+}$  ions are attributed to the decrease in  $H_2$  and increase  $CO_2$  compositions at 12 hours.

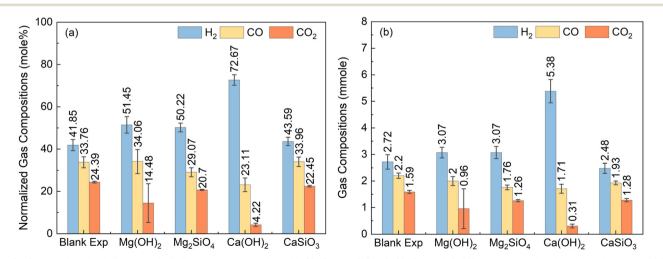
In contrast, Mg<sub>2</sub>SiO<sub>4</sub> for in situ CO<sub>2</sub> capture yielded significantly different results compared to the use of  $Mg(OH)_2$  (Fig. 5(b)).  $H_2$  compositions are 16.9% and 19.2% respectively, at 3 hours and 6 hours of reaction time, while CO compositions are 71.6% and 66.5% in the presence of Mg<sub>2</sub>SiO<sub>4</sub>. These results indicate that the WGSR is the limiting step in the first 6 hours of reaction in the presence of Mg<sub>2</sub>-SiO<sub>4</sub>. This is likely due to the slower dissolution behavior of  $Mg_2SiO_4$  to release  $Mg^{2+}$  ions compared to that of  $Mg(OH)_2$ . As a result, in situ CO<sub>2</sub> capture and subsequent enhancement in H<sub>2</sub> yields is limited in the presence of Mg<sub>2</sub>SiO<sub>4</sub> in the first six hours of the reaction. At 9 hours of reaction time, H<sub>2</sub> composition is 60.3%. CO composition of 66.5% at 6 hours decrease to 15.5% at 9 hours of reaction time (Fig. 5(b)). These results indicate that while the kinetics of WGSR are slower in the first 6 hours of the reaction in the presence of  $Mg_2SiO_4$  compared to in the presence of  $Mg(OH)_2$ , comparable compositions are achieved when the reaction time is 9 hours.

As in the case of  $Mg_2SiO_4$ , a decrease in  $H_2$  compositions and increase in  $CO_2$  compositions is observed when the reaction time is 12 hours.  $H_2$  compositions decrease from 60.3% to 53.0% while  $CO_2$  compositions increase from 24.2% to 35.6% at 12 hours of reaction. The relative decrease in  $H_2$ compositions and increase in  $CO_2$  compositions at 12 hours is attributed to the mass transfer limitations associated with the dissolution of Mg- silicate and Mg-hydroxide. Therefore, these studies indicate that WGSR can be the initial slow step in the presence of  $Mg_2SiO_4$  as opposed to  $Mg(OH)_2$  during the first 6 hours of reaction. Non-monotonic changes in the concentrations of  $H_2$ , CO, and CO<sub>2</sub> observed as the reaction time increases from 9 hours to 12 hours are likely due to limitations from the slow dissolution of Mg(OH)<sub>2</sub> and Mg<sub>2</sub>-SiO<sub>4</sub> to release Mg<sup>2+</sup> ions for CO<sub>2</sub> capture.

#### 3.5 Influence of Mg- and Ca-bearing hydroxides and silicates on enhanced H<sub>2</sub> conversion with *in situ* CO<sub>2</sub> capture

Contrasting the influence of Mg- and Ca-bearing silicates and hydroxides on enhanced H<sub>2</sub> conversion coupled with in situ CO<sub>2</sub> capture can unlock new opportunities for the utilization of a wide range of alkaline feedstocks. Ca-bearing hydroxides and silicates are abundant in fly ash and slags.<sup>69</sup> Mg-bearing hydroxides and silicates are abundant in ultramafic and mafic mine tailings.<sup>70</sup> To this end, experiments are conducted at 300 °C in the presence of 8 bar of CO and 12 bar of N<sub>2</sub> in the gas phase and Mg(OH)<sub>2</sub>, Ca(OH)<sub>2</sub>, Mg<sub>2</sub>SiO<sub>4</sub>, or CaSiO<sub>3</sub> to capture CO<sub>2</sub> in a slurry phase over a reaction time of 12 hours, as noted in section 3.1. Pt/Al<sub>2</sub>O<sub>3</sub> catalyst is used as a low-temperature WGSR catalyst. N2 gas is used as a reference due to its inert nature and constant quantity before and after the reaction which enables the determination of the moles of gas products formed. The influence of  $Mg(OH)_2$ , Ca(OH)<sub>2</sub>, Mg<sub>2</sub>SiO<sub>4</sub>, or CaSiO<sub>3</sub> on compositions of H<sub>2</sub>, CO, and  $CO_2$  are shown in Fig. 6.

The influence of WGSR alone in the absence of Ca- and Mg-bearing hydroxide and silicate determined at the same experimental conditions as the others showed that  $H_2$ , CO, and CO<sub>2</sub> account for 41.85%, 33.76%, and 24.39%, respectively (Fig. 6(a)). Relative to this blank experiment, the highest enhancement in  $H_2$  conversion and CO<sub>2</sub> suppression was observed in the presence of Ca(OH)<sub>2</sub>. In this best case scenario with Ca(OH)<sub>2</sub>,  $H_2$  yield is 72.67% which is 30.82% higher compared to the blank experiment. CO<sub>2</sub> yields are



**Fig. 6** Contrasting the influence of alkaline resources such as  $Mg(OH)_2$ ,  $Mg_2SiO_4$ ,  $Ca(OH)_2$ , and  $CaSiO_3$  on enhancing  $H_2$  yield with *in situ*  $CO_2$  capture and mineralization where (a) represents normalized gas mole percentages and (b) represents exact gas moles. Experiments are conducted at 300 °C, 20 atm for 12 h. The blank experiment is conducted over 3 g of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst without any alkaline source. The test experiments are conducted over 3 g of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst without any alkaline source. The test experiments are conducted over 3 g of Pt/Al<sub>2</sub>O<sub>3</sub> catalyst with a sorbent-to-liquid weight ratio of 3:10, leading to mole percentages of 8.48% for  $Mg(OH)_2$ , 3.70% for  $Mg_2SiO_4$ , 6.80% for  $Ca(OH)_2$ , and 4.45% for  $CaSiO_3$ , respectively.

suppressed by 20.17% to 4.22% relative to the blank experiment indicating that Ca(OH)<sub>2</sub> in the slurry is highly effective in capturing  $CO_2$ emissions. Significant enhancement in WGSR is also evident from the changes in CO compositions which are lowered by 10.65% compared to the blank experiment. In comparison to  $Ca(OH)_2$ , the H<sub>2</sub> yields are  $\sim$ 50–51% in the presence of Mg(OH)<sub>2</sub> and Mg<sub>2</sub>SiO<sub>4</sub> which are lower. Interestingly, CO compositions are 5% lower when  $Mg_2SiO_4$  is used compared to  $Mg(OH)_2$ , indicating WGSR is enhanced in the presence of Mg<sub>2</sub>SiO<sub>4</sub>. However, the higher solubility of Mg(OH)<sub>2</sub> compared to Mg<sub>2</sub>SiO<sub>4</sub> enhances CO<sub>2</sub> capture lowering CO<sub>2</sub> compositions by 6.22% when  $Mg(OH)_2$  is used related to  $Mg_2SiO_4$ .<sup>67</sup>

The reactivity of CaSiO<sub>3</sub> in enhancing H<sub>2</sub> yield and suppressing CO<sub>2</sub> emissions is lower compared to the other alkaline sources. Compared to the blank experiment, the composition of H<sub>2</sub> is 1.74% higher, CO is similar, and CO<sub>2</sub> is  $\sim 2\%$  lower. While these results indicate that CaSiO<sub>3</sub> enhances H<sub>2</sub> conversion and enables CO<sub>2</sub> capture, substantial changes relative to the blank experiment are not observed. The composition data in mmoles are reported in Fig. 6(b) for use in subsequent modeling efforts. The results reported in Fig. 6 indicate that the effectiveness of the alkaline sources in enhancing H<sub>2</sub> yields while capturing CO<sub>2</sub> emissions follows this order:  $CaSiO_3 < Mg_2SiO_4 < Mg(OH)_2 < Ca(OH)_2$ . As hypothesized, the reactivity of Ca- and Mg-bearing hydroxides in enhancing H<sub>2</sub> yield and suppressing CO<sub>2</sub> emissions is higher compared to that of the corresponding silicates. The higher solubility and faster dissolution rates of and Mg-bearing hydroxides compared to the corresponding silicates facilities enhanced CO2 capture and H<sub>2</sub> conversion.<sup>67</sup>

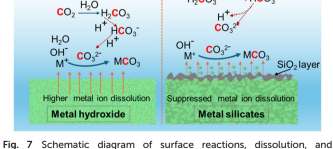
# 3.6 Mechanistic insights underlying enhancement in H<sub>2</sub> conversion coupled with *in situ* CO<sub>2</sub> capture and mineralization

The key reasons for the differences in the reactivity of Caand Mg-bearing silicates vs. hydroxides are the slower kinetics of dissolution of silicates and the formation of a

Carbonation process

(a)Metal hydroxide

(b) Metal Silicates



**Fig.** / Schematic diagram of surface reactions, dissolution, and mineralization process over (a) metal hydroxides and (b) metal silicates sorbents.

silica passivation layer which is known to limit mass transfer.<sup>71,72</sup> Fig. 7 represents the multiphase chemical reaction mechanisms associated with carbon mineralization by harnessing hydroxides versus silicates. While the CO<sub>2</sub> hydration reactions to produce bicarbonate and carbonate species remain unchanged, the leaching of the metal into the solution results in the formation of a silica-rich layer in the case of silicates unlike in hydroxides. It is interesting to note that despite the slower kinetics of Mg-silicate dissolution and the formation of a silica layer that can limit mass transfer, comparable gas phase compositions of H<sub>2</sub>, CO, and CO<sub>2</sub> are noted with  $Mg(OH)_2$  and  $Mg_2SiO_4$  (Fig. 6). However, the kinetics associated with enhancing H2 conversion with in situ CO<sub>2</sub> capture and mineralization can differ significantly as noted in Fig. 5. The slow kinetics of WGSR observed up to 6 hours in the presence of Mg<sub>2</sub>SiO<sub>4</sub> are attributed to limited availability of Mg<sup>2+</sup> ions to capture CO<sub>2</sub> and facilitate the forward reaction. However, beyond 6 hours of reaction time, the observed enhancement in H<sub>2</sub> yields and corresponding decrease in CO2 emissions is attributed to the availability of sufficient Mg<sup>2+</sup> ions to capture CO<sub>2</sub> and aid the forward WGSR to produce H<sub>2</sub>.

Prior studies have shown that temperatures above 100 °C and high partial pressures of CO<sub>2</sub> aid the carbon mineralization of olivine.<sup>39,73</sup> The solubility of CO<sub>2</sub>, and dissolution of Mg<sub>2</sub>SiO<sub>4</sub> to release Mg<sup>2+</sup> ions for capturing CO<sub>2</sub> and producing carbonates at the experimental conditions used in this study are analogous to those reported in prior studies. Therefore, despite the slow initial kinetics of  $Mg_2SiO_4$ , the release of  $Mg^{2+}$  ions over time results in comparable H<sub>2</sub>, CO, and CO<sub>2</sub> compositions as when Mg(OH)<sub>2</sub> is used. It is also interesting to note that Mg<sub>2</sub>SiO<sub>4</sub> is more effective in enhancing WGSR compared to CaSiO<sub>3</sub>. Prior studies reported that CaSiO<sub>3</sub> is reactive at temperatures as low as 110 °C for carbon mineralization, unlike Mg<sub>2</sub>SiO<sub>4</sub>.<sup>74</sup> Possible reasons for the lower-than-expected reactivity of CaSiO<sub>3</sub> are the formation of a silica passivation layer that is thicker and limits mass transfer at the experimental conditions reported in this study. To uncover if CO2 uptake by the alkaline slurry results in the formation of solid carbonates, detailed structural and morphological analyses are investigated and reported in the following section.

# 3.7 Structural and morphological analyses of the alkaline sources

To determine if the captured  $CO_2$  only dissolves in the aqueous phase or reacts to produce solid carbonates, the changes in the structural and morphological features of the alkaline resources are determined. The crystalline features in unreacted and reacted Ca- and Mg-bearing silicates and hydroxides using X-ray diffraction (XRD) measurements (Fig. 8). As shown in Fig. 8(a), the unreacted Mg(OH)<sub>2</sub> exhibits the typical diffractogram of hexagonal Mg(OH)<sub>2</sub> brucite structure with characteristic XRD peaks at 33.2°, 38.1°, 50.6°, 58.7°, 68.3°, and 72.1° (see peaks in black).

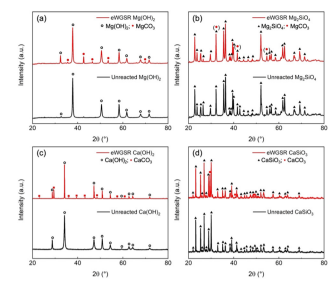


Fig. 8 Determination of the crystalline phases before and after reaction starting with (a)  $Mg(OH)_2$ , (b)  $Mg_2SiO_4$ , (c)  $Ca(OH)_2$ , and (d)  $CaSiO_3$  using X-ray diffraction (XRD) analyses.

These peaks are well in accordance with the reported literature (PDF 44-1482).<sup>75</sup> The diffractogram of the reacted solid (peak in red) revealed the existence of magnesite (MgCO<sub>3</sub>) peaks along with suppression of Mg(OH)<sub>2</sub> intensities. This co-existence indicates the dissolution of Mg(OH)<sub>2</sub> and the capture and conversion of CO<sub>2</sub> to produce MgCO<sub>3</sub>.<sup>73</sup> To investigate if MgCO<sub>3</sub> is formed starting from Mg<sub>2</sub>SiO<sub>4</sub> precursors, XRD data of the unreacted and reacted materials are collected and shown in Fig. 8(b). The X-ray diffraction pattern corresponding to the monoclinic structure of unreacted Mg<sub>2</sub>SiO<sub>4</sub> is shown in Fig. 8(b).

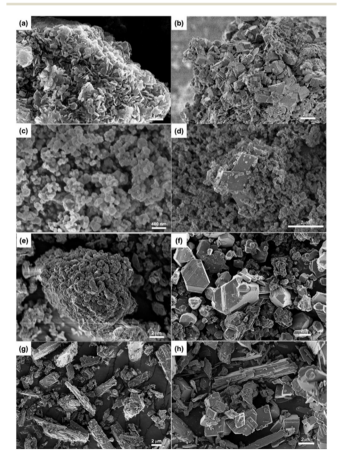
The reacted Mg<sub>2</sub>SiO<sub>4</sub> product exhibits a very similar scattering pattern as the unreacted material, which can results from overlapping between (104) peak of the generated magnesite and (130) peak of the residual Mg<sub>2</sub>SiO<sub>4</sub>.<sup>76,77</sup> To further distinguish the different phases and prove the formation of carbonate, additional characterization is required. Interestingly, the absence of the nesquehonite  $(MgCO_3 \cdot 3H_2O)$  or hydromagnesite  $(Mg_5(CO_3)_4(OH)_2 \cdot 4H_2O)$ phases in our case is due to the higher stability of anhydrous magnesite (MgCO<sub>3</sub>) at elevated temperatures and pressures.<sup>39</sup> Additionally, the formation of Mg-carbonate can be influenced by numerous other factors, including the reaction time, ionic strength or the addition of reagents including salts.<sup>39</sup> Thus, in this study, the formation of anhydrous magnesite (MgCO<sub>3</sub>) can be attributed to the application of high temperatures of 250-300 °C with elevated CO<sub>2</sub> partial pressures in the range of 10-20 atm for reaction durations of 9-12 hours.

The structural arrangement characterization of the Ca cases shows a similar trend with Mg cases, in which the anhydrous calcite (CaCO<sub>3</sub>) is the unique carbonate product due to the favorable reaction conditions. Fig. 8(c) illustrates the XRD peaks of unreacted Ca(OH)<sub>2</sub> at 28.7°, 34.1°, 47.2°,

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50.6°, 54.3°, 62.3°, and 64.3° corresponds to the (100), (101), (102) (110), (111), (201) and (112) planes, respectively. Several studies have reported similar patterns of Ca(OH)2.78,79 After the enhanced WGSR, the reacted product shows the presence of both Ca(OH)<sub>2</sub> and CaCO<sub>3</sub> phases (JCPDS 84-1276 and 85-1108, respectively). The occurrence of peaks at 23.1°, 29.5°, 31.6°, 36.1°, 39.5°, 43.3°, 47.2°, 47.6°, 48.6°, 57.5°, 61.1°, and 64.5° corresponds to (012), (104), (006), (110), (114), (202), (024), (018), (116), (112) (119), and (300) planes, respectively. These XRD patterns corresponded to the rhombohedral crystal structure of CaCO<sub>3</sub>.<sup>80</sup> Furthermore, Fig. 8(d) shows the diffraction peaks of unreacted sorbents can be indexed as the CaSiO<sub>3</sub>.<sup>81</sup> For the reacted CaSiO<sub>3</sub>, the XRD peaks resemble the co-occurrence of CaSiO<sub>3</sub> and CaCO<sub>3</sub> phases, which is a low-temperature polymorph of calcium carbonate (JCPDS 00-005-0586).82

Furthermore, the morphologies of the reacted  $Mg(OH)_2$ ,  $Ca(OH)_2$ ,  $Mg_2SiO_4$ , and  $CaSiO_3$  materials are determined using FE-SEM to identify carbonate-bearing phases. Fig. 9(b) shows the cube-like structure of  $MgCO_3$  with the appearance of residual  $Mg(OH)_2$  after carbon mineralization, confirming the carbonate formation after the enhanced WGSR. Similarly,



**Fig. 9** Morphology of (a) unreacted  $Mg(OH)_2$ ; (b) reacted  $Mg(OH)_2$ ; (c) unreacted  $Mg_2SiO_4$ ; (d) reacted  $Mg_2SiO_4$ ; (e) unreacted  $Ca(OH)_2$ ; (f) reacted  $Ca(OH)_2$ ; (g) unreacted  $CaSiO_3$ ; (h) reacted  $CaSiO_3$  determined by scanning electron microscope measurements.

Table 2 The surface area, pore volume, and mean pore diameter of the unreacted and reacted alkaline sources as determined using BET analyses and
the BJH method

	Unreacted alkaline sources			Reacted alkaline sources				
	Surface area $(m^2 g^{-1})$	Pore volume $(cm^3 g^{-1})$	Mean pore diameter (nm)	Surface area $(m^2 g^{-1})$	Pore volume $(cm^3 g^{-1})$	Mean pore diameter (nm)		
$Mg(OH)_2$	9.78	0.047	4.01	6.48	0.05	3.93		
Mg <sub>2</sub> SiO <sub>4</sub>	17.41	0.050	2.81	14.93	0.08	3.93		
$Ca(OH)_2$	24.29	0.107	3.72	4.81	0.01	1.50		
CaSiO <sub>3</sub>	5.00	0.020	2.66	6.20	0.02	3.94		

the cube-shaped  $MgCO_3$  particles emerged in the case of reacted  $Mg_2SiO_4$  materials (Fig. 9(d)).

Interestingly, the granular particles covered on cubic  $MgCO_3$  particles can be attributed to the residual  $Mg_2SiO_4$  and associated  $SiO_2$  layer resulting from silicate dissolution. Similar phenomena are also observed in the reacted  $Ca(OH)_2$  and  $CaSiO_3$  materials (Fig. 9(f) and (h)). Additionally, the large hexagonal particles indicate the significant dissolution and recrystallization of initial  $Ca(OH)_2$  sorbents (Fig. 9(f)), and the elongated particles correspond to the residual  $CaSiO_3$  materials (Fig. 9(g) and (h)). Interestingly, the XRD analysis does not show the clear existence of crystalline phases rich in silica, indicating the potential presence of an amorphous silica-rich layer or glassy  $SiO_2$  coating that may cause the suppression in metal dissolution in  $CaSiO_3$ ,  $Mg_2SiO_4$ , responsible for the low extents of carbon mineralization.

As a further clarification of the sorbents' morphological evolution, the surface area and the pore distribution determined by BET analysis are listed in Table 2. The decreased surface area in Mg(OH)2, Mg2SiO4, and Ca(OH)2 cases indicates the formation of larger carbonate particles like  $MgCO_3$  and  $CaCO_3$  (Fig. 9(a)-(f)). Notably, the significant surface area reduction in the Ca(OH)<sub>2</sub> case aligns with its more effective CO<sub>2</sub> mineralization and the corresponding greater carbonate formation. In contrast, due to the larger particle size and limited pore volume in the unreacted CaSiO<sub>3</sub> compared with the generated CaCO<sub>3</sub>, the small increase in the surface area and pore volume after reaction are attributed to the dominance of dissolution over carbon mineralization (Fig. 9(g) and (h)). Furthermore, the pore size distributions determined using the BJH model are illustrated in Fig. S4.† The unreacted hydroxide sorbents exhibit distributions, centered relatively narrow pore at approximately 4.01 nm for Mg(OH)2 and 3.72 nm for significant  $Ca(OH)_2$ . Post-reaction, pore structure rearrangements are observed, leading to reductions in both pore volume and diameter, particularly for Ca(OH)<sub>2</sub>. Additionally, the formation of MgCO<sub>3</sub> results in more wellordered pore and particle size distributions compared to the Mg<sub>2</sub>SiO<sub>4</sub> precursor. Conversely, significant changes in the pore size distribution of CaSiO<sub>3</sub> before and after reaction are not observed due to the relatively low extent of carbon mineralization.

Attenuated total reflection Fourier transform infrared (ATR-FTIR) analysis is also performed on unreacted sorbents

and reacted products to confirm carbonate formation after the enhanced WGSR. As shown in Fig. 10(a) and (c), unreacted hydroxide sorbents exhibited only one peak between 3600-3700 cm<sup>-1</sup>, indicating the purity of starting materials (Mg(OH)<sub>2</sub>, Ca(OH)<sub>2</sub>).<sup>83</sup> On the other hand, the occurrence of new peaks around 1400-1500 cm<sup>-1</sup>, 850-890 cm<sup>-1</sup>, and 700-750 cm<sup>-1</sup> corresponds to C-O asymmetric stretching vibrations, the C-O bending vibration, and the inplane bending vibration of O-C-O, respectively, in the carbonated products formed after the enhanced WGSR.84,85 Furthermore, peaks shown in ATR-FTIR of unreacted Mg<sub>2</sub>-SiO<sub>4</sub> sorbents around 600-650 cm<sup>-1</sup> are consistent with SiO<sub>4</sub> bending and stretching modes, respectively, while the peak around 1000 cm<sup>-1</sup> represents asymmetric stretching modes of Si-O-Si.86,87 As shown in Fig. 10(b), despite its very low intensity, the peak at 1100 cm<sup>-1</sup> could be interpreted as a symmetric C-O stretching vibration band indicating the carbon mineralization of Mg<sub>2</sub>SiO<sub>4</sub>.<sup>84</sup> Moreover, in the case of CaSiO<sub>3</sub> materials, the presence of peaks approximately 600- $700 \text{ cm}^{-1}$ ,  $900 \text{ cm}^{-1}$  corresponds to the symmetrical stretching of O-Si-O bonds and Si-O-Ca bonds, while the peaks between 1000-1100 cm<sup>-1</sup> correspond to a symmetrical stretching of Si-O-Si bonds, respectively.88 As shown in Fig. 10(d), the typical peaks from silicate remained constant

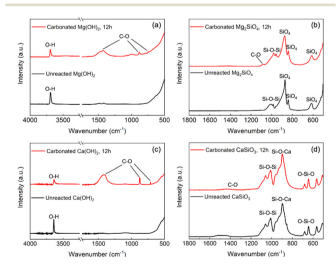


Fig. 10 Identification of different functional groups using attenuated total reflection-Fourier transform infrared (ATR-FTIR) spectroscopy measurements in unreacted and reacted (a) Mg(OH)<sub>2</sub>; (b) Mg<sub>2</sub>SiO<sub>4</sub>; (c) Ca(OH)<sub>2</sub>; (d) CaSiO<sub>3</sub> materials.

after the enhanced WGSR with a little shoulder around 1400 cm<sup>-1</sup>, indicating the asymmetric stretching mode of C–O, proving the carbon mineralization of  $CaSiO_3$  material.<sup>85</sup>

Further, to investigate the effect of eWGSR on the metal hydroxides and metal silicates, XPS analyses are conducted. The XPS spectra were calibrated based on the carbon C-C peak (284.8 eV).<sup>89</sup> Fig. S5(a1) and (a2)† shows the O 1s deconvolution result for unreacted and reacted Mg(OH)<sub>2</sub> materials. The high-resolution XPS spectrum of unreacted Mg  $(OH)_2$  exhibit a peak at binding energy of 531.7 eV which suggests the existence of hydroxyl (OH) species. Furthermore, the reacted Mg(OH)<sub>2</sub> materials showed slight changes in O 1s spectrum (Fig. S5(a2)<sup>†</sup>). Binding energy (BE) peaks are observed at 531.8 eV and 533.34 eV which correspond to hydroxyl (OH) and carbonate (CO<sub>3</sub><sup>2-</sup>) species, respectively.<sup>90,91</sup> Similar trends are observed within the unreacted and reacted Ca(OH)<sub>2</sub> materials. The hydroxyl (OH) peak at 531.9 eV exclusively appears in the unreacted Ca(OH)2 while the hydroxyl peak at 531.8 eV and carbonate peak at 532.9 coexists in the reacted Ca(OH)<sub>2</sub> material (Fig. S5(b1) and (b2)<sup>†</sup>). To further determine the influence of enhanced WGSR on metal silicates (CaSiO<sub>3</sub> and Mg<sub>2</sub>SiO<sub>4</sub>), the valance state of the oxygen in the unreacted and reacted silicates are studied using XPS analyses. Fig. S6<sup>†</sup> shows the carbon (C1s) spectra from unreacted and reacted Mg<sub>2</sub>SiO<sub>4</sub>, CaSiO<sub>3</sub> materials consisting of two main components arising from C-C (~284.4 eV) and O=C (carboxyl, ~288.6 eV).92,93 Compared to unreacted counterparts, the reacted materials exhibit higher intensity signal of C 1s indicating the presence of carbonate (CO<sub>3</sub><sup>2-</sup>) species. The C 1s peak values observed between 289.5 eV and 289.7 eV are well-matched with the carbonate  $(CO_3^{2-})$  species in the reported literature.<sup>94,95</sup>

Insights into the changes in the structure of silicon are obtained from XPS analyses. Fig. S7(a1)<sup>†</sup> shows that O 1s peak at 531.35 eV corresponds to the SiO<sub>4</sub> tetrahedra in the unreacted  $Mg_2SiO_4$ .<sup>96</sup> Additionally, the O1s peak tail at a higher binding energy region around 533.6 eV could result from minor amorphous SiO<sub>2</sub> content.<sup>97</sup> On the other hand, the reacted  $Mg_2SiO_4$  materials ((Fig. S7(a2))) show deconvoluted peaks at 531.1, 532.3, and 533.5 eV which corresponds to the residual Mg<sub>2</sub>SiO<sub>4</sub>, the generated carbonate and the amorphous SiO<sub>2</sub> resulting from silicate dissolution.98-101 On the contrary, Fig. S7(b1)† shows that unreacted CaSiO<sub>3</sub> exhibits a SiO<sub>2</sub> peak at 533.5 eV other than the typical  $SiO_3$  peak at 532 eV, indicating the presence of an amorphous phase in the unreacted sorbent.97,102 After the enhanced WGSR, the O1s peaks of the reacted CaSiO<sub>3</sub> shift towards a high binding energy range. More specifically, the existence of deconvoluted O1s peaks at 531.4 eV, 532.9 eV, and 533.3 eV correspond to the residual SiO<sub>3</sub>, generated carbonate and SiO<sub>2</sub>, which confirms carbonate formation and the dissolution of calcium silicate material.

Further, Fig. S7(c1) and (d1)<sup>†</sup> depicts the existence of Si2p broad peak at the binding energy of 102.7 eV and 103.2 eV for unreacted  $Mg_2SiO_4$  and  $CaSiO_3$ , respectively.<sup>103</sup> It's worth noting that the SiO<sub>2</sub> peaks around 104 eV appear in both the

silicates, corresponding to the results from O1s peak.<sup>104</sup> However, after carbon mineralization, Si 2p deconvoluted spectra shows the diminished  $SiO_4$  or  $SiO_3$  peaks with intensified  $SiO_2$  peaks for both silicates, indicating the dissolution of silicate and the generation of Si-rich passivation layer.<sup>105,106</sup> Additionally, the relatively high amorphous  $SiO_2$  component in the unreacted CaSiO<sub>3</sub> compared to Mg<sub>2</sub>SiO<sub>4</sub> likely suppresses the dissolution of CaSiO<sub>3</sub>, thus resulting in lowering H<sub>2</sub> yields compared to Mg<sub>2</sub>SiO<sub>4</sub> materials.

To quantitatively evaluate the carbonate phases, the carbon/metal ratios are calculated before and after the enhanced WGSR based on the atom% from XPS (Fig. S8<sup>+</sup>). The detected carbon phase in the unreacted carbonate-free sorbents can be attributed to the adventitious carbon on samples' surface due to the carbon contamination during air exposure.<sup>107</sup> However, neglecting the impact from adventitious carbon, the carbon/metal atomic ratio after the enhanced WGSR increased to 0.58 with Mg(OH)2, 0.78 with Ca(OH)<sub>2</sub>, 0.06 with Mg<sub>2</sub>SiO<sub>4</sub>, and 0.1 with CaSiO<sub>3</sub>, respectively. In terms of the carbonate fraction, the increased carbon/metal ratio of Mg<sub>2</sub>SiO<sub>4</sub> requires doubling to compare with other sources due to its doubled stoichiometric coefficient of metal ion, which results in a 0.12 increase in Mg<sub>2</sub>SiO<sub>4</sub> carbon/metal atomic ratio after the enhanced WGSR. In this case, the varying trend in carbon/metal atomic ratio increases perfectly aligns with the enhanced WGSR:  $Ca(OH)_2 > Mg(OH)_2 > Mg_2SiO_4 > CaSiO_3$ , indicating the strong positive correlation between the enhanced H<sub>2</sub> yields and carbonate formation.

# 3.8 Insights on the mechanisms of the enhanced WGSR and *in situ* CO<sub>2</sub> capture and carbon mineralization

Given the multiphase reaction described in Fig. 1, the key reaction mechanisms involved in the enhanced WGSR can be delineated as CO oxidation and CO<sub>2</sub> capture. To be more specific, the CO oxidation proceeds *via* two potential approaches simultaneously: (i) the gaseous-phase WGSR with the assistance of noble-metal (Pt/Al<sub>2</sub>O<sub>3</sub>) catalyst (eqn (1)) and (ii) the aqueous-phase CO reaction to form formate followed by subsequent formate decomposition (eqn (14) and (15)).<sup>108</sup>

$$CO_{(aq)} + OH_{(aq)} \rightarrow HCOO_{(aq)}$$
 (14)

$$HCOO_{(aq)}^{-} + H_2O_{(aq)} \rightarrow CO_{2(g)} + H_{2(g)} + OH_{(aq)}^{-}$$
(15)

The first mechanism results in maximum  $H_2$  production, while the second mechanism consumes part of the CO supply yet may not produce  $H_2$  eventually due to the limitation of formate decomposition. It is well-reported that the reaction between CO and hydroxide species leads to a competition between CO and CO<sub>2</sub> dissolution in the second mechanism. According to previous studies, higher CO partial pressure in the system may result in a larger amount of expelled CO<sub>2</sub> from the solution and restrict the subsequent carbon

mineralization step.<sup>109</sup> In this situation, the alkalinity of the slurry becomes an essential factor of the aqueous-phase CO oxidation: more sufficient hydroxide ion supply alleviates the competition between CO and CO<sub>2</sub> and promotes both H<sub>2</sub> production and CO2 capture, which explains the enhancement in performance of hydroxide in this study when compared to their silicate counterparts (Fig. 6). Additionally, to evaluate the impact of the non-decomposed formate, the separated aqueous phase after reactions are characterized by liquid-state nuclear magnetic resonance (NMR), as shown in Table S3.† Although the blank experiment without alkaline sorbent exhibits a relatively high residual formate level in the aqueous phase, only trace amounts of formate (<200 ppm) were detected in the aqueous phase when alkaline resources were applied. The observation indicates the presence of alkaline sorbents effectively catalyzes the decomposition of the generated formate, leading to the near-complete oxidization of dissolved CO into CO<sub>2</sub>.

On the other hand, the generated  $CO_2$  from both CO oxidation mechanisms further hydrated to form bicarbonate and carbonate species and released protons into the aqueous phase (eqn (16)).<sup>110</sup> Simultaneously, the dissolution of the alkaline resources released the metal cations ( $M^{2+}$ ), which could be enhanced by the increasing proton level. Consequently, the metal cations combine with carbonate species ( $CO_3^{2-}$ ,  $HCO_3^{-}$ ) precipitated out when the solubility limit is reached (eqn (17)). In our case, the carbonate products tend to be in anhydrous form due to their favored chemical stability at the operating temperature (250–300 °C) in this multiphase reaction environment,<sup>111</sup> which is also confirmed by the characterization of the solid products.

$$CO_{2(aq)} + H_2O_{(l)} \rightarrow HCO_{3(aq)}^- + H_{(aq)}^+ \rightarrow CO_{3(aq)}^{2-} + 2H_{(aq)}^+$$
 (16)

$$M_{(aq)}^{2+} + CO_{3(aq)}^{2-} \to MCO_{3(s)}$$
 (17)

## 4. Conclusions

In this study, the feasibility of the enhanced WGSR integrated with in situ thermodynamically downhill CO<sub>2</sub> capture and mineralization is investigated in the presence of Mg- and Ca-hydroxides and silicates to co-produce H<sub>2</sub> and carbonate-bearing materials. Calibrated tuning of the multiphase chemical interactions is unlocked by deriving the thermodynamical models of the multiphase reactions, where the theoretical prediction of the CO conversion is mutually corroborated with the experimental results. For example, 30.82% more H<sub>2</sub> yields with 20.17% less CO<sub>2</sub> emission achieved with Ca(OH)<sub>2</sub> compared with independent WGS reaction without in situ CO2 capture and mineralization. Thus, these studies proved the hypothesis that the in situ capture and mineralization of gaseous CO<sub>2</sub> depletes CO<sub>2</sub> gas compositions and favors the forward reaction to produce H<sub>2</sub>. The higher solubility and dissolution rates of Ca- and Mghydroxides favors enhanced  $H_2$  yields. Comparable compositions of  $H_2$  are achieved with  $Mg(OH)_2$  and  $Mg_2SiO_4$ when reacted for 12 hours at 300C despite initial slower kinetics of CO conversion with  $Mg_2SiO_4$  in the first 6 hours of the reaction.

Among the studied alkaline resources for enhanced WGSR,  $Ca(OH)_2$  exhibited unbeatable  $CO_2$  capture over the other sorbents (CO<sub>2</sub> compositions:  $Ca(OH)_2$  (4.22%) <  $Mg(OH)_2$  (14.48%) <  $Mg_2SiO_4$  (20.7%) <  $CaSiO_3$  (22.45%) < blank experiment (24.39%) respectively), when feeding a mixture of CO and N2 at 300 °C and 20 bar. Additionally, the temporal evolution in H<sub>2</sub> yields and CO<sub>2</sub> capture revealed that the kinetic limitation of the overall reaction was dependent on the sorbent dissolution behavior, especially for the silicate sorbents. The amorphous Si-rich layer generated from silicate dissolution and mineralization passivated the alkaline resources and restricted the rate of enhanced WGSR, particularly in the case of CaSiO<sub>3</sub>. The dissolution of Mg- and Ca-bearing hydroxides and silicates, formation of the silica passivation layer, and anhydrous metal carbonate products in the reacted alkaline resources are confirmed from structural and morphological analyses. Detailed characterization and analyses confirmed that anhydrous carbonates (MgCO<sub>3</sub>, CaCO<sub>3</sub>) are the dominant carbonate bearing products without any associated hydrated carbonate formations. Quantitative analyses of the reacted products showed that the carbonate content corresponded directly to the trend in H<sub>2</sub> yields. Evidence of comparable H<sub>2</sub> yields with  $Mg_2SiO_4$  and  $Mg(OH)_2$  indicates the feasibility of directly using silicates for enhanced WGSR as opposed to extracting Mg(OH)<sub>2</sub> from Mg<sub>2</sub>SiO<sub>4</sub> resources, thus enabling more material and energy efficient pathways for producing H<sub>2</sub> with inherent CO<sub>2</sub> capture and storage in the form on durable Mgand Ca-bearing carbonates. This approach can also unlock new opportunities in harnessing renewable biomethane for co-producing H<sub>2</sub> with inherent CO<sub>2</sub> capture and mineralization to produce carbonates for use in construction materials by harnessing energy from renewable sources, thus enabling carbon-negative pathways for a sustainable energy and resource future.

## Data availability

The data supporting this article have been included in the main manuscript and as part of the ESI.<sup>†</sup>

## Author contributions

Greeshma Gadikota – supervision, conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, resources, software, validation, visualization, writing – original draft, writing – review & editing. Xun Gao – conceptualization, data curation, formal analysis, investigation, methodology, resources, software, validation, visualization, writing – original draft, writing – review & editing; Divya Prasad – investigation, validation, visualization, writing – review & editing; Mahadeo A. Mahadik – investigation, validation, visualization, writing – review & editing.

## Conflicts of interest

Greeshma Gadikota is the co-founder of Carbon To Stone which is commercializing technologies for industrial decarbonization and carbon management. The other authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

## Acknowledgements

All the analytical lab work has been performed at the School of Civil and Environmental Engineering, Cornell University. The authors acknowledge the use of shared facilities at the Cornell Center for Materials Research (CCMR). The DOE CAREER Award supports G. G. and X. G.'s contributions through the Office of Science: DE-SC0020263. The authors also gratefully acknowledge the support of Ivan Kuzmenko at APS and Peilong Lu at Cornell University for assisting in this effort.

## References

- 1 R. E. H. Sims, H. H. Rogner and K. Gregory, *Energy Policy*, 2003, **31**, 1315–1326.
- 2 International Energy Agency (IEA), *Global Energy Review:* CO<sub>2</sub> Emissions in 2021, IEA, 2022.
- 3 D. Prasad, K. N. Patil, N. K. Chaudhari, H. Kim, B. M. Nagaraja and A. H. Jadhav, *Catal. Rev.: Sci. Eng.*, 2022, 64, 356–443.
- 4 M. A. Rosen and S. Koohi-Fayegh, *Energy, Ecol. Environ.*, 2016, 1, 10–29.
- 5 M. A. Pellow, C. J. M. Emmott, C. J. Barnhart and S. M. Benson, *Energy Environ. Sci.*, 2015, 8, 1938–1952.
- 6 I. K. Kapdan and F. Kargi, *Enzyme Microb. Technol.*, 2006, **38**, 569–582.
- 7 P. Nikolaidis and A. Poullikkas, *Renewable Sustainable Energy Rev.*, 2017, **67**, 597–611.
- 8 J. D. Holladay, J. Hu, D. L. King and Y. Wang, *Catal. Today*, 2009, **139**, 244–260.
- 9 M. I. Temkin, Adv. Catal., 1979, 28, 173-287.
- 10 C. Rhodes, G. J. Hutchings and A. M. Ward, *Catal. Today*, 1995, 23, 43–58.
- 11 C. Ratnasamy and J. P. Wagner, *Catal. Rev.:Sci. Eng.*, 2009, **51**, 325–440.
- 12 H. Bohlbro, J. Catal., 1964, 3, 207–215.
- 13 W. Xu, R. Si, S. D. Senanayake, J. Llorca, H. Idriss, D. Stacchiola, J. C. Hanson and J. A. Rodriguez, *J. Catal.*, 2012, 291, 117–126.
- 14 P. Mierczynski, W. Maniukiewicz and T. P. Maniecki, *Cent. Eur. J. Chem.*, 2013, **11**, 912–919.
- 15 D. W. Jeong, W. J. Jang, J. O. Shim, W. B. Han, H. S. Roh, U. H. Jung and W. L. Yoon, *Renewable Energy*, 2014, 65, 102–107.

- 16 X. Zhang, M. Zhang, Y. Deng, M. Xu, L. Artiglia, W. Wen, R. Gao, B. Chen, S. Yao, X. Zhang, M. Peng, J. Yan, A. Li, Z. Jiang, X. Gao, S. Cao, C. Yang, A. J. Kropf, J. Shi, J. Xie, M. Bi, J. A. van Bokhoven, Y. W. Li, X. Wen, M. Flytzani-Stephanopoulos, C. Shi, W. Zhou and D. Ma, *Nature*, 2021, **589**, 396–401.
- 17 L. Chen, Z. Qi, S. Zhang, J. Su and G. A. Somorjai, *Catalysts*, 2020, **10**, 858.
- 18 E. R. Van Selow, P. D. Cobden, P. A. Verbraeken, J. R. Hufton and R. W. Van Den Brink, *Ind. Eng. Chem. Res.*, 2009, 48, 4184–4193.
- 19 J. Gibbins and H. Chalmers, *Energy Policy*, 2008, 36, 4317-4322.
- 20 I. Sreedhar, T. Nahar, A. Venugopal and B. Srinivas, Renewable Sustainable Energy Rev., 2017, 76, 1080-1107.
- 21 R. Ben-Mansour, M. A. Habib, O. E. Bamidele, M. Basha, N. A. A. Qasem, A. Peedikakkal, T. Laoui and M. Ali, *Appl. Energy*, 2016, **161**, 225–255.
- 22 R. Khalilpour, K. Mumford, H. Zhai, A. Abbas, G. Stevens and E. S. Rubin, *J. Cleaner Prod.*, 2015, **103**, 286–300.
- 23 C. A. Scholes, M. T. Ho, D. E. Wiley, G. W. Stevens and S. E. Kentish, *Int. J. Greenhouse Gas Control*, 2013, 17, 341–348.
- 24 A. Sanna, M. Uibu, G. Caramanna, R. Kuusik and M. M. Maroto-Valer, *Chem. Soc. Rev.*, 2014, **43**, 8049–8080.
- 25 H. M. Jang, K. B. Lee, H. S. Caram and S. Sircar, *Chem. Eng. Sci.*, 2012, **73**, 431–438.
- 26 S. M. Kim, A. Armutlulu, A. M. Kierzkowska, D. Hosseini, F. Donat and C. Müller, *Sustainable Energy Fuels*, 2020, 4, 713–729.
- 27 M. S. Duyar, R. J. Farrauto, M. J. Castaldi and T. M. Yegulalp, *Ind. Eng. Chem. Res.*, 2014, 53, 1064–1072.
- 28 C. Zhang, Y. Li, Z. He, J. Zhao and D. Wang, *Appl. Catal.*, *B*, 2022, **314**, 121474.
- 29 T. Noor, M. V. Gil and D. Chen, Appl. Catal., B, 2014, 150– 151, 585–595.
- 30 R. W. Stevens, A. Shamsi, S. Carpenter and R. Siriwardane, *Fuel*, 2010, **89**, 1280–1286.
- 31 Y. Hu, H. Cui, Z. Cheng and Z. Zhou, Chem. Eng. J., 2019, 377, 119823.
- 32 C. H. Lee and K. B. Lee, Appl. Energy, 2017, 205, 316-322.
- 33 S. Sun, C. Zhang, S. Chen, X. Zhao, Y. Wang, S. Xu and C. Wu, R. Soc. Open Sci., 2023, 10, 230067.
- 34 K. S. Lackner, C. H. Wendt, D. P. Butt, E. L. Joyce and D. H. Sharp, *Energy*, 1995, 20, 1153–1170.
- 35 D. E. Giammar, R. G. Bruant and C. A. Peters, *Chem. Geol.*, 2005, **217**, 257–276.
- 36 P. B. Kelemen, R. Aines, E. Bennett, S. M. Benson, E. Carter, J. A. Coggon, J. C. De Obeso, O. Evans, G. Gadikota, G. M. Dipple, M. Godard, M. Harris, J. A. Higgins, K. T. M. Johnson, F. Kourim, R. Lafay, S. Lambart, C. E. Manning, J. M. Matter, K. Michibayashi, T. Morishita, J. Noël, K. Okazaki, P. Renforth, B. Robinson, H. Savage, R. Skarbek, M. W. Spiegelman, E. Takazawa, D. Teagle, J. L. Urai and J. Wilcox, in *Energy Procedia*, Elsevier Ltd, 2018, vol. 146, pp. 92–102.

- 37 B. P. McGrail, H. T. Schaef, A. M. Ho, Y. J. Chien, J. J. Dooley and C. L. Davidson, *J. Geophys. Res.:Solid Earth*, 2006, **111**, 1–13.
- 38 K. S. Lackner, Annual Review of Energy and the Environment, 2002, 27, 193–232.
- 39 G. Gadikota, J. Matter, P. Kelemen and A. H. A. Park, *Phys. Chem. Chem. Phys.*, 2014, **16**, 4679–4693.
- 40 G. Gadikota, J. Matter, P. Kelemen, P. V. Brady and A. H. A. Park, *Fuel*, 2020, 277, 117900.
- 41 G. Gadikota, E. J. Swanson, H. Zhao and A. H. A. Park, *Ind. Eng. Chem. Res.*, 2014, 53, 6664–6676.
- 42 R. Baciocchi, G. Costa, A. Polettini, R. Pomi and V. Prigiobbe, *Energy Procedia*, 2009, **1**, 4851–4858.
- 43 R. Zevenhoven, S. Teir and S. Eloneva, *Energy*, 2008, 33, 362–370.
- 44 C. R. Müller, R. Pacciani, C. D. Bohn, S. A. Scott and J. S. Dennis, *Ind. Eng. Chem. Res.*, 2009, **48**, 10284–10291.
- 45 E. R. van Selow, P. D. Cobden, A. D. Wright, R. W. van den Brink and D. Jansen, *Energy Procedia*, 2011, 4, 1090–1095.
- 46 Y. Liu, Z. Li, L. Xu and N. Cai, *Ind. Eng. Chem. Res.*, 2012, 51, 11989–11997.
- 47 J. Boon, K. Coenen, E. van Dijk, P. Cobden, F. Gallucci and M. van Sint Annaland, *Adv. Chem. Eng.*, 2017, **51**, 3–96.
- 48 D. Iruretagoyena, K. Hellgardt and D. Chadwick, *Int. J. Hydrogen Energy*, 2018, **43**, 4211–4222.
- 49 T. J. Stadler, P. Barbig, J. Kiehl, R. Schulz, T. Klövekorn and P. Pfeifer, *Energies*, 2021, **14**, 355.
- 50 J. Fagerlund, J. Highfield and R. Zevenhoven, *RSC Adv.*, 2012, 2, 10380–10393.
- 51 L. Zhao, L. Sang, C. Jun, J. Ji and H. H. Teng, *Environ. Sci. Technol.*, 2010, 44, 406–411.
- 52 A. I. Fernández, J. M. Chimenos, M. Segarra, M. A. Fernández and F. Espiell, *Hydrometallurgy*, 1999, 53, 155–167.
- 53 W. K. O'Connor, D. C. Dahlin, G. E. Rush, S. J. Gerdemann, L. R. Penner and R. P. Nilsen, Aqueous Mineral Carbonation: Mineral Availability, Pretreatment, Reaction Parametrics, and Process Studies, DOE/ARC-TR-04-002, Albany Research Center, Albany, 2005.
- 54 A. I. Fernández, J. M. Chimenos, M. Segarra, M. A. Fernández and F. Espiell, *Hydrometallurgy*, 1999, 53, 155–167.
- 55 J. S. Loring, C. J. Thompson, C. Zhang, Z. Wang, H. T. Schaef and K. M. Rosso, *J. Phys. Chem. A*, 2012, **116**, 4768–4777.
- 56 W. J. J. Huijgen, G. J. Witkamp and R. N. J. Comans, *Environ. Sci. Technol.*, 2005, **39**, 9676–9682.
- 57 G. Gadikota, Nat. Rev. Chem., 2020, 4, 78-89.
- 58 M. Bilal and S. D. Jackson, *Appl. Catal.*, *A*, 2017, 529, 98–107.
- 59 R. Y. Chein, Y. H. Lin, Y. C. Chen, Y. P. Chyou and J. N. Chung, *Int. J. Hydrogen Energy*, 2014, **39**, 18854–18862.
- 60 I. Kocemba, I. Śmiechowicz, M. Jędrzejczyk, J. Rogowski and J. M. Rynkowski, *Catalysts*, 2021, **11**, 1475.
- 61 E. Baraj, K. Ciahotný and T. Hlinčík, *Fuel*, 2021, **288**, 119817.

- 62 P. Ochonma, C. Noe, S. Mohammed, A. Mamidala and G. Gadikota, *React. Chem. Eng.*, 2023, 8, 1943–1959.
- 63 B. R. Smith, J. M. Loganathan and M. Shekhar Shantha, *Int. J. Chem. React. Eng.*, 2010, 8, 1–32.
- 64 K. J. Fricker, Magnesium Hydroxide Sorbents for Combined Carbon Dioxide Capture and Storage In Energy Conversion Systems, *PhD Thesis*, Columbia University, 2014.
- 65 D. P. Harrison, Ind. Eng. Chem. Res., 2008, 47, 6486-6501.
- 66 E. Abbasi, A. Hassanzadeh, S. Zarghami, H. Arastoopour and J. Abbasian, *Fuel*, 2014, **137**, 260–268.
- 67 J. L. Palandri and Y. K. Kharaka, Open-File Rep. U. S. Geol. Surv., 2004, 1068, 1–64.
- 68 B. Li, L. Wei, H. Yang, X. Wang and H. Chen, *Energy*, 2014, 68, 248–254.
- 69 Z. Giergiczny, Cem. Concr. Res., 2019, 124, 105826.
- 70 A. Polat, P. W. U. Appel and B. J. Fryer, Gondwana Res., 2011, 20, 255–283.
- 71 S. V. Golubev, O. S. Pokrovsky and J. Schott, *Chem. Geol.*, 2005, 217, 227–238.
- 72 G. Rim, A. K. Marchese, P. Stallworth, S. G. Greenbaum and A.-H. A. Park, *Chem. Eng. J.*, 2020, **396**, 125204.
- 73 S. Katre, P. Ochonma, H. Asgar, A. M. Nair, K. Ravi and G. Gadikota, *Phys. Chem. Chem. Phys.*, 2024, 26, 9264–9283.
- 74 S. J. Gerdemann, W. K. O'Connor, D. C. Dahlin, L. R. Penner and H. Rush, *Environ. Sci. Technol.*, 2007, 41, 2587–2593.
- 75 R. Kurosawa, M. Takeuchi and J. Ryu, *RSC Adv.*, 2021, **11**, 24292–24311.
- 76 W. Liang, Y. Yin, L. Wang, L. Chen and H. Li, J. Alloys Compd., 2017, 702, 346–351.
- 77 Z. F. Cheng, X. Hu, Y. Li and Z. Y. Ling, J. Am. Ceram. Soc., 2016, 99, 2688–2692.
- 78 Y. H. Lee, H. Eom, S. M. Lee and S. S. Kim, RSC Adv., 2021, 11, 8306–8313.
- 79 H. Zheng, Y. He, Y. Zhu, L. Liu and X. Cui, RSC Adv., 2021, 11, 12476–12483.
- 80 K. S. P. Karunadasa, C. H. Manoratne, H. M. T. G. A. Pitawala and R. M. G. Rajapakse, *J. Phys. Chem. Solids*, 2019, 134, 21–28.
- L. Ernawati, R. A. Wahyuono, A. D. Laksono, A. Ningrum, K. Handayani and A. Sabrina, *IOP Conf. Ser.:Mater. Sci. Eng.*, 2021, **1053**, 012001.
- 82 R. Lakshmi, V. Velmurugan and S. Sasikumar, *Combust. Sci. Technol.*, 2013, **185**, 1777–1785.
- 83 T. N. Ramesh and A. Taj, Int. J. Sci. Res., 2013, 1, 58.
- 84 C. H. Park, J. H. Lee, E. Jang, K. B. Lee and J. H. Kim, *Chem. Eng. J.*, 2017, **30**7, 503–512.
- 85 A. Z. Noah, M. A. El Semary, A. M. Youssef and M. A. El-Safty, *Egypt. J. Pet.*, 2017, 26, 33–40.
- 86 S. H. Tamin, S. B. R. S. Adnan, M. H. Jaafar and N. S. Mohamed, *Ionics*, 2018, 24, 2665–2671.
- 87 A. Saberi, B. Alinejad, Z. Negahdari, F. Kazemi and A. Almasi, *Mater. Res. Bull.*, 2007, **42**, 666–673.
- 88 M. V. Reddy and M. Pathak, J. Inorg. Organomet. Polym. Mater., 2018, 28, 2187–2195.

- 89 K. H. Tan, A. Iqbal, F. Adam, N. H. H. Abu Bakar, M. N. Ahmad, R. M. Yusop and H. Pauzi, *RSC Adv.*, 2019, 9, 38760–38771.
- 90 J. P. Holgado, G. Munuera, J. P. Espinos and A. R. Gonzalez-Elipe, *Appl. Surf. Sci.*, 2000, **158**, 164–171.
- 91 P.-C. Huang, A. C. Vilando, T.-H. Ha and M.-C. Lu, Sustainable Environ. Res., 2024, 34, 11.
- 92 X. Y. Peng, X. X. Liu, D. Diamond and K. T. Lau, *Carbon*, 2011, 49, 3488–3496.
- 93 Y. Xu, H. Bai, G. Lu, C. Li and G. Shi, J. Am. Chem. Soc., 2008, 130, 5856–5857.
- 94 C. M. Chen, J. Q. Huang, Q. Zhang, W. Z. Gong, Q. H. Yang, M. Z. Wang and Y. G. Yang, *Carbon*, 2012, 50, 659–667.
- 95 A. Dufourny, C. Julcour, J. Esvan, L. Cassayre, P. Laniesse and F. Bourgeois, *Front. Clim.*, 2022, 4, 946735.
- 96 B. Han, Y. Yang, J. Li, H. Deng and C. Yang, Int. J. Electrochem. Sci., 2018, 13, 9166–9182.
- 97 M. T. Nichols, W. Li, D. Pei, G. A. Antonelli, Q. Lin, S. Banna,
  Y. Nishi and J. L. Shohet, *J. Appl. Phys.*, 2014, 115, 094105.
- 98 M. J. Guittet, J. P. Crocombette and M. Gautier-Soyer, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2001, 63, 125117.
- 99 A. A. Khassin, T. M. Yurieva, M. P. Demeshkina, G. N. Kustova, I. Sh. Itenberg, V. V. Kaichev, L. M. Plyasova, V. F. Anufrienko, I. Yu. Molina, T. V. Larina, N. A. Baronskaya and V. N. Parmon, *Phys. Chem. Chem. Phys.*, 2003, 5, 4025–4031.

- 100 H. Xu, Q. Wang, H. Xiao, X. Li, X. Su, M. Tang, L. Chen and S. Li, *RSC Adv.*, 2019, 9, 1319–1326.
- 101 J. Esvan, G. Berger, S. Fabre, E. Bêche, Y. Thébault, A. Pages and C. Charvillat, *Geochim. Cosmochim. Acta*, 2022, 335, 124–136.
- 102 L. Liu, J. Liu, L. Zhao, Z. Yang, C. Lv, J. Xue and A. Tang, *Environ. Sci. Pollut. Res.*, 2019, 26, 8721–8736.
- 103 X. Gao, H. Asgar, I. Kuzmenko and G. Gadikota, *Microporous Mesoporous Mater.*, 2021, **327**, 111381.
- 104 A. Kaur, P. Chahal and T. Hogan, *IEEE Electron Device Lett.*, 2016, 37, 142–145.
- 105 D. A. Shirley, High-Resolution X-Ray Photoemission Spectrum of the Valence Bands of Gold, *Phys. Rev. B: Solid State*, 1972, 5, 4709.
- 106 T. V. Larina, L. S. Dovlitova, V. V. Kaichev, V. V. Malakhov, T. S. Glazneva, E. A. Paukshtis and B. S. Bal'Zhinimaev, *RSC Adv.*, 2015, 5, 79898–79905.
- 107 G. Greczynski and L. Hultman, *ChemPhysChem*, 2017, 18, 1507–1512.
- 108 D. C. Elliott, R. T. Hallen and L. J. Sealock, *Ind. Eng. Chem. Process Des. Dev.*, 1983, 22, 431–435.
- 109 W. A. R. Slegeir, R. S. Sapienza, R. Rayford and L. Lam, Organometallics, 1982, 1, 1728–1730.
- 110 A. H. G. Cents, D. W. F. Brilman and G. F. Versteeg, *Chem. Eng. Sci.*, 2005, **60**, 5830–5835.
- 111 M. Hänchen, V. Prigiobbe, R. Baciocchi and M. Mazzotti, *Chem. Eng. Sci.*, 2008, **63**, 1012–1028.