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Dolomite: a low cost thermochemical energy storage material†

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The thermal energy storage properties of dolomite, $\text{CaMg}(\text{CO}_3)_2$, from three sources (commercial, mined, and synthetic) are investigated as a potential solid–gas thermochemical energy store for concentrating solar thermal power (CSP) plants. The reversible carbon dioxide release/absorption cycle can be used to store and release large quantities of thermal energy that can be harnessed near 550 °C. To enable carbon dioxide absorption, a novel molten salt eutectic $\text{NaCl} : \text{MgCl}_2$ mixture was added to the dolomite, which proved effective. Curiously, the mined (unrefined) sample proved to have the best calcination/carbonation properties with a stable cyclic stability of ~50 mol% CO_2 over 10 cycles. A morphological investigation by scanning electron microscopy reveals the mined sample contains impurities (e.g. quartz) that prevent detrimental agglomeration of dolomite and its reaction products (CaCO_3 and MgO). High levels of porosity in the mined dolomite also provide short gas–solid reaction pathways during carbonation. A commercial dolomite source contained too many impurities, resulting in the formation of high levels of Ca_2SiO_4 that acted as a Ca-sink, lowering CO_2 capacity. Finally, a high purity synthetic dolomite sample displayed high levels of agglomeration and phase segregation, lowering reversible CO_2 capacity, which was attributed to a lack of impurities that restrict agglomeration.

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Introduction

Renewable energy sources are becoming cheaper and cost-competitive with fossil fuels. However, most renewable energy supplies still lack an energy storage solution, restricting them to intermittent operation based on variable energy fluxes (e.g. wind and solar).¹ Concentrating solar thermal power (CSP) plants utilise solar power in a technologically simple manner to generate and store thermal energy, which is used to heat water and run a steam turbine based electrical generator.² Currently, a number of 1st generation CSP plants utilise molten salt technology to store thermal energy *via* the specific heat of the liquid when heating from 290–565 °C.^{2,3} However, the specific heat capacity (150 kJ kg⁻¹ per 100 °C) of state-of-the-art molten nitrate salts (40NaNO₃ : 60KNO₃) is low compared to other phase change or thermochemical energy storage options (Table 1). Most importantly, the low energy density requires large quantities of salt to store large amounts of energy, resulting in

high overall energy storage costs despite its relatively low cost per kg (Fig. 1a). Thus, 2nd generation energy storage systems with higher energy density and operational temperature will offer lower overall CSP plant costs and higher technology uptake by industrial and commercial sectors.³

There are a wide selection of chemicals available to replace molten salts as thermal energy storage materials, including: latent heat storage materials (phase change materials, PCMs) such as inorganic salts/salt eutectics and metals/metal alloys; solid sensible storage media such as concrete and ceramics, where the heat transfer fluid is circulated around the media; and also thermochemical materials such as metal hydrides and metal carbonates.³ Metal hydrides have been identified as promising thermochemical heat storage options for concentrating solar thermal power because of their impressive energy densities.^{4–7} Although metal hydrides have high material costs, it is offset by the low quantity of material necessary (Table 1, Fig. 1a).⁸ Therefore, metal hydrides offer potential cost savings over current generation molten salt technology.⁹ Despite the benefits of metal hydrides, they remain difficult to handle in air due to their sensitivity to oxygen and moisture. Significant cost savings for 2nd generation CSP plants may also be realised when considering much cheaper materials, including directly minable minerals such as metal carbonates (Table 1 and Fig. 1a). A recent techno-economic assessment determined the cost of various oxide and carbonates as thermal energy storage

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Table 1 Thermochemical, energy density, and conservative cost parameters for energy storage materials in a comparable CSP scenario^{11,12}

	Molten salt (40NaNO ₃ : 60 KNO ₃)	MgH ₂ ⇌ Mg + H ₂	CaH ₂ ⇌ Ca + H ₂	MgCO ₃ ⇌ MgO + CO ₂	CaCO ₃ ⇌ CaO + CO ₂	CaMg(CO ₃) ₂ ⇌ MgO + CaCO ₃ + CO ₂
Enthalpy ΔH (kJ mol ⁻¹ CO ₂ or H ₂)	39.0	74.1	162.4	96.7	165.8	125.8
Molar mass (g mol ⁻¹)	94.60	26.33	42.10	84.31	100.09	184.40
Density (g cm ⁻³)	2.17	1.42	1.7	2.96	2.71	2.85
Capacity (wt% CO ₂ or H ₂)	—	7.66	4.79	52.2	44.0	23.9
Gravimetric energy density (kJ kg ⁻¹)	413	2811	3857	1147	1657	682
Volumetric energy density ^c (MJ m ⁻³)	895	3995	6557	3396	4489	1944
Operating temperature range (°C)	290–565	300–400	>1020	~450	>880	~590
Carnot efficiency ^a (%)	46	47	77	58	74	65
Estimated practical efficiency (%)	27	27	52	35	49	41
Mass required ^b (tonnes)	5250	755	339	1500	819	2260
Volume required ^b (m ³)	2420	532	199	507	302	793
Materials cost (\$ per tonne) ¹³	630	3000	6000	500	60	60
Total materials cost required ^b (\$)	3 307 500	2 260 000	2 030 000	750 000	49 000	136 000

^a Lower temperature. ^b To generate 1 TJ of electrical energy, parameters for pure material only. ^c Based on crystalline density.

materials including dolomite.¹⁰ It noted that despite the low cost of the material, parasitic energy losses, including employing a gas compressor to store the CO₂, would consume 82% of the energy produced, leading to a total capital cost of \$82 per MJ. Cheaper storage of CO₂ without compression must be considered including: operating at higher temperatures to promote gas operating pressures of above 64 bar (promoting liquefaction of CO₂);¹⁴ or the use of low-temperature CO₂ storage in another carbonate material.

A host of research has been conducted to investigate the mineral sequestration of carbon dioxide into metal oxides, forming metal carbonates.¹⁵ The process is reversible, where CO₂ can be cycled in and out of the metal oxide, in highly exothermic and endothermic reactions. This process embodies the thermochemical process that can be harvested for thermal energy storage applications.^{4,9,16} In a CSP application, solar energy would heat a metal carbonate in the daytime, which absorbs energy (endothermic process) and releases CO₂ forming a metal oxide. The CO₂ would be stored in pressure vessels

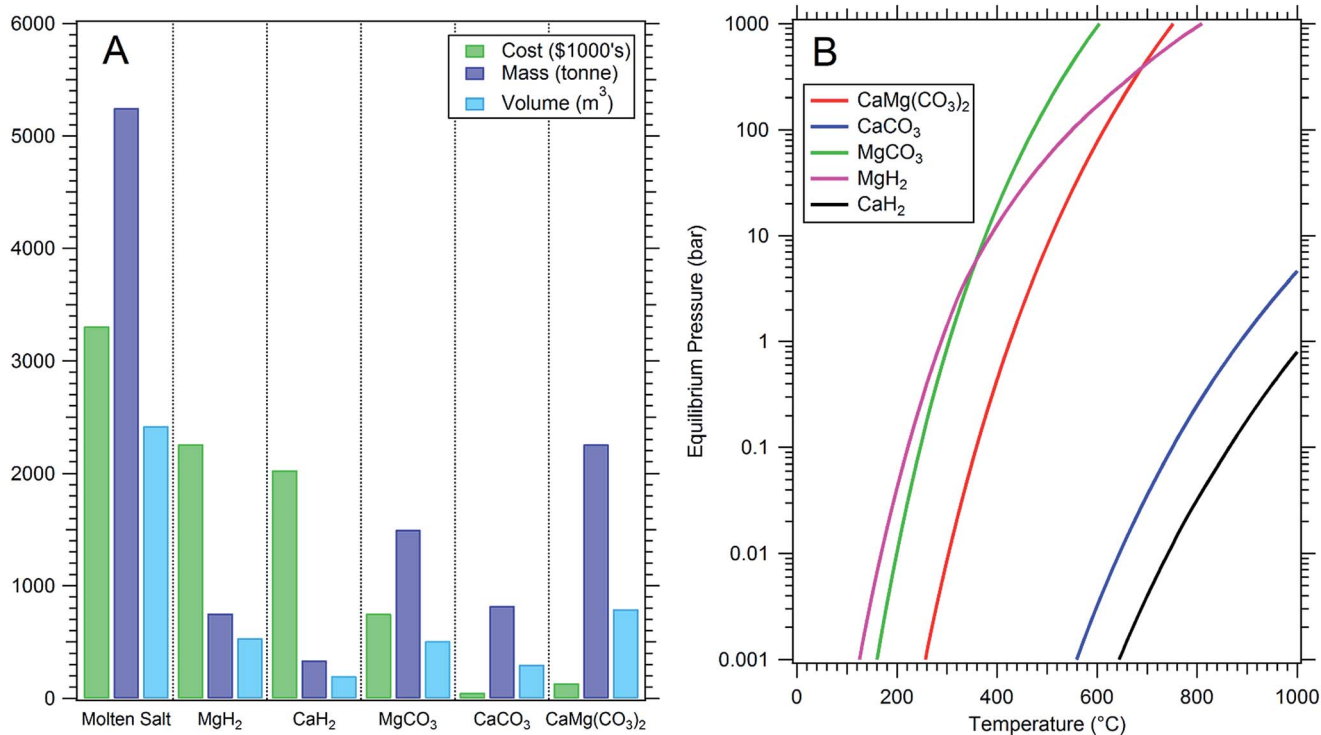
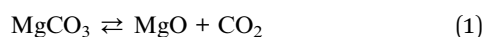


Fig. 1 (A) Comparison of thermal energy storage options to generate 1 TJ of electrical energy (see legend for y axis labels). (B) Equilibrium pressure of thermochemical energy storage materials based on thermodynamics of reaction.¹¹

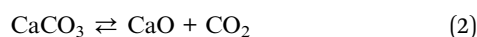


within this closed system. At night, when solar energy fades, CO₂ would reabsorb into the metal oxide in an exothermic process, generating heat. In fact, the entire process is self-limiting, governed by the thermodynamics of the solid–gas reactions, which define a temperature dependent equilibrium pressure (Fig. 1b).

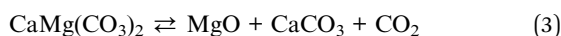
Carbonate rocks such as magnesite (MgCO₃), calcite (CaCO₃) and dolomite (CaMg(CO₃)₂) form approximately 20% of the sedimentary crust on the Earth.¹⁷ The abundance and basic processing requirements result in extremely low materials costs. Crushed limestone (mostly CaCO₃) was sold on the gigatonne scale in the USA in 2001 at \$5.19 per tonne on average, whereas dolomite had a value of \$5.64 per tonne.¹⁸ Calcined products, high-calcium quicklime (CaO) and dolomitic quicklime (CaO, MgO) are produced at moderate purity (>95%) with a value of approximately \$60 per tonne, whereas magnesia (MgO) is more expensive, ranging from \$90–500 per tonne, varying based on purity and source. The thermodynamics of calcination and carbonation are well-known:¹¹



$$\Delta H_{1\text{bar}} = 98.9 \text{ kJ mol}^{-1} \text{ CO}_2, \Delta S_{1\text{bar}} = 171.3 \text{ J mol}^{-1} \text{ CO}_2/\text{K}$$



$$\Delta H_{1\text{bar}} = 165.8 \text{ kJ mol}^{-1} \text{ CO}_2, \Delta S_{1\text{bar}} = 143.0 \text{ J mol}^{-1} \text{ CO}_2/\text{K}$$



$$\Delta H_{1\text{bar}} = 126.2 \text{ kJ mol}^{-1} \text{ CO}_2, \Delta S_{1\text{bar}} = 180.6 \text{ J mol}^{-1} \text{ CO}_2/\text{K}$$

Here, the bimetallic carbonate, dolomite, CaMg(CO₃)₂, only releases one unit of CO₂, before forming more stable CaCO₃, which will not decompose until temperatures >800 °C.

Dolomite offers a thermodynamic stabilisation in comparison to MgCO₃, where the carbon dioxide is released at a higher temperature from the bimetallic material. Conveniently, once carbon dioxide is released, only stable metal oxides or carbonates remain in the solid-phase, reducing problems seen in other thermochemical heat storage materials at high temperatures due to kinetically hindering agglomeration and metallic vaporisation.^{8,9}

During the carbonation of metal oxides an impervious (or diffusion limiting) metal carbonate layer can form that restricts further CO₂ absorption.^{19,20} As such, the reaction kinetics of carbonation are governed by particle size, pore structure and surface area. A major problem is the cyclic stability, or reversibility, of carbonation and calcination reactions, as ~10 000 cycles are required over the 30 year operating lifetime of a CSP plant.² An example is CaO/CaCO₃, which only retains 7–8 wt% CO₂ capacity after 500 cycles.²¹ Natural CaO–MgO sorbents (20–40 μm) derived from dolomite have previously been investigated for CO₂ capture, but displayed no carbonation of MgO and poor cycling capacity in the CaO/CaCO₃ component of the sample

(~60% reversible carbonation after 50 cycles).²² In general, very poor carbonation rates are observed for MgO (<25%),¹⁹ which have been shown to improve in the presence of high pressure steam, but cycling studies have yet to be conducted.

Sodium nitrate (NaNO₃) has been heralded as a molten-salt based catalyst for dolomite and other MgO-containing CO₂ absorbents.^{23,24} Zhang *et al.* studied its effect in detail and provided an illustrative schematic of the process.²³ In general, the mechanism promoting CO₂ absorption relies on the low melting point of NaNO₃ (306.5 °C), where CO₂ absorption correlates with the presence of a liquid phase, even below the melting point in the presence of “pre-melting” or “surface melting”.²³ Carbon dioxide adsorbs onto the surface of an MgO particle and then migrates to the phase boundary between solid MgO and liquid NaNO₃, where the molten salt promotes reactivity and the formation of a metal carbonate. Here, the catalytic effect is assigned to the slight solubility of MgO,^{25,26} CO₂,²⁷ and other carbonate components (such as CaCO₃) in the molten salt surface layer, where they gain enhanced mobility and can precipitate out as a double-salt, *e.g.* dolomite. Note that NaNO₃ decomposes above 570 °C.^{28,29} Therefore, in the present work the performance of a new, higher temperature molten salt system (NaCl/MgCl₂) is studied as an alternative. It is noted that the addition of 20 wt% salt additive will decrease the reported energy density (546 kJ kg⁻¹ instead of 682 kJ kg⁻¹) in Table 1 and will also affect the system cost.

Experimental

Mined dolomite, CaMg(CO₃)₂, was supplied from commercial supplier Richgro and mine site Watheroo Dolomitic Lime, WA, Australia (samples: Rich-A, Wath-A). Subsequently, these dolomite samples were heated from RT to 1000 °C at a rate of 6.5 °C min⁻¹ and held isothermal for 1 h under an Ar flow of 2 L min⁻¹ to form predominantly CaO and MgO (calcined samples: Rich-B, Wath-B). As a reference, a high purity synthetic sample was prepared: metal nitrate precursors were supplied from Sigma-Aldrich, Mg(NO₃)₂·6H₂O (99%) and Ca(NO₃)₂·4H₂O (99%). A 1 M aqueous solution of 1 : 1 Mg(NO₃)₂ and Ca(NO₃)₂ was heated to 115 °C *in vacuo*, while stirring, to uniformly co-crystallise the reagents. The resulting white powder was subsequently heated from RT to 650 °C at a rate of 5 °C min⁻¹ and held isothermal for 3 h under an Ar flow of 2 L min⁻¹, to form the corresponding metal oxides (MgO and CaO; sample: Synth-B). A eutectic mixture of 0.438MgCl₂ : 0.562NaCl (m.p. 442 °C)³⁰ was ground within an argon glovebox (mBraun < 1 ppm H₂O, O₂) and mixed in a mortar and pestle with samples Synth-B and Wath-B in a 20 wt% loading. An overview of samples and preparation/treatment methods is given in Table 2.

X-Ray diffraction (XRD) was performed using a Bruker D8 Advance diffractometer equipped with a CuKα_{1,2} source in flat-plate geometry mode. Salt-containing samples were analysed in an argon-filled poly(methyl methacrylate) dome, which results in a broad hump in the diffraction background around 2θ = 20°. Data were collected using a Lynxeye PSD detector from 5–80° 2θ at 0.04° s⁻¹. *In situ* synchrotron powder X-ray diffraction (SR-



XRD) was performed at the Powder Diffraction (PD) beamline at the Australian Synchrotron in Melbourne, Australia.³¹ The annealed powder was loaded in a quartz capillary (outer diameter 0.7 mm, wall thickness 0.01 mm) while inside a glove box, and mounted to a sample holder constructed from Swagelok tube fittings using a graphite ferrule. The sample holder was then connected to a vacuum manifold and the capillary heated with a hot air blower with a heating rate of 6 °C min⁻¹ under a static argon environment. One-dimensional SR-XRD patterns (monochromatic X-rays with $\lambda = 0.8263076 \text{ \AA}$) were continuously collected using a Mythen microstrip detector³² with an exposure time of 30 s for each of two detector positions. The capillary was continuously oscillated through 100° during exposure to improve the powder averaging and ensure even heating. Diffraction patterns were quantitatively analysed with the Rietveld method³³ using TOPAS (Bruker-AXS).

Scanning Electron Microscopy (SEM) was undertaken on a Tescan Lyra3 field emission microscope coupled with secondary electron (SE) and backscattered electron (BSE) detectors, and an Oxford Instruments Energy Dispersive Spectroscopy (EDS) detector. Samples were sputter coated with 5 nm of Pt and were either dispersed as a powder or embedded in epoxy resin and mechanically polished using an ethanol working fluid before imaging.

Nitrogen (N₂) adsorption/desorption measurements were performed at 77 K using a Micromeritics ASAP 3020 system (Micromeritics, Nor-cross, GA, USA) to determine the specific surface area and the porosity of the samples. N₂ adsorption data, at relative pressures (P/P_0) between 0.05 and 0.30, were used to calculate the specific surface area by employing the Brunauer–Emmett–Teller (BET) multi-point method.³⁴ The Barrett–Joyner–Halenda (BJH) method from the absorption branch of the isotherm³⁵ was used to calculate the pore size

distributions and volumes in the macro- (>50 nm) and meso-ranges (2–50 nm).

Simultaneous Differential Scanning Calorimetry-Thermogravimetric Analysis (DSC-TGA) was performed on a Mettler Toledo DSC 1 instrument using sample masses of ~10 mg. Samples were measured at a heating rate of 10 °C min⁻¹ under an argon flow of 20 mL min⁻¹. The temperature accuracy of this instrument is $\pm 0.2 \text{ °C}$, while the balance has an accuracy of $\pm 20 \text{ }\mu\text{g}$.

Cyclic measurements of CO₂ absorption/desorption were performed on a High Energy PCTpro E&E at 450–550 °C for >10 cycles. Desorption was carried out under an initial back pressure of 2 bar CO₂ into a reservoir volume of 165 cm³. Absorption was carried out under an initial pressure of 30 bar CO₂ in a reservoir volume of 12.3 cm³. The digital pressure transducer (Rosemount 3051S) had a precision and accuracy of 14 mbar. Room temperature measurements were recorded using a 4-wire platinum resistance temperature detector (RTD) while sample temperatures were measured using a K-type thermocouple.

Results and discussion

Mined dolomite samples (Rich-A and Wath-A) were analysed, as calcite (CaCO₃), magnesite (MgCO₃), quartz (SiO₂), and additional silicates. Analysis of the crystalline component of the samples showed high purity dolomite for the Watheroo batch (91.8 wt%), but lower purity for the Richgro batch (70.8 wt%). The problem with impurities, especially silicates, becomes apparent when the dolomite samples are calcined at 1000 °C (Rich-B and Wath-B) where silicates react with the calcium-rich components to form larnite (Ca₂SiO₄) and/or tricalcium aluminate (Ca₃Al₂O₆). Here, calcium is effectively leached from the active carbonate/oxide component of the samples and trapped in a thermodynamic sink (Fig. S2†). Table 3 shows that

Table 2 Sample Overview

Sample	Sample name	Primary content	Temperature	Time	Gas	Catalyst
As supplied						
Richgro dolomite	Rich-A	CaMg(CO ₃) ₂	—	—	—	—
Watheroo dolomite	Wath-A	CaMg(CO ₃) ₂	—	—	—	—
Thermally outgassed						
Synthetic CaO/MgO	Synth-B	MgO/CaO	650 °C	3 h	—	—
Synthetic CaO/MgO (with catalyst)	Synth-B-cat	MgO/CaO	650 °C	3 h	—	NaCl : MgCl ₂
Calcined Richgro (from Rich-A)	Rich-B	MgO/CaO	1000 °C	1 h	—	—
Calcined Watheroo (from Wath-A)	Wath-B	MgO/CaO	1000 °C	1 h	—	—
Calcined Watheroo (from Wath-A, with catalyst)	Wath-B-cat	MgO/CaO	1000 °C	1 h	—	NaCl : MgCl ₂
Carbonated (single-step 50 bar CO₂)						
Carbonated synthetic (from Synth-B)	Synth-C	MgO/CaCO ₃	500 °C	54 h	CO ₂	—
Carbonated synthetic (from Synth-B, with catalyst)	Synth-C-cat	CaMg(CO ₃) ₂	500 °C	8 h	CO ₂	NaCl : MgCl ₂
Re-carbonated Watheroo (from Wath-B)	Wath-C	MgO/CaCO ₃	500 °C	18 h	CO ₂	—
CO₂ cycled						
11-Times cycled Watheroo (from Wath-B-cat in the absorbed state)	Wath-D-cat	CaMg(CO ₃) ₂ /MgO/CaCO ₃	450–550 °C	110 h	CO ₂	NaCl : MgCl ₂
14-Times cycled synthetic (from Synth-B-cat in the absorbed state)	Synth-D-cat	CaMg(CO ₃) ₂ /MgO/CaCO ₃	550 °C	450 h	CO ₂	NaCl : MgCl ₂



sample Rich-B only contains 5.1 wt% CaO and Wath-B contains 27.8 wt% CaO, compared to an ideal 58.2 wt% CaO content. In contrast, the Synth-B sample has a 57.7 wt% CaO, close to the theoretical quantity. Irreversible containment of CaO in silicates obviously makes it impossible to carbonate (with CO₂ gas) the samples back to their original dolomite content.

A single-step carbonation of thermally outgassed samples (Wath-B, Synth-B) was attempted at 500 °C and 50 bar CO₂ (18–54 h). X-Ray diffraction results of the carbonated samples (Wath-C, Synth-C), presented in Table 3 and Fig. S3,† demonstrate that carbonation was successful, but only for the CaO portion of the sample, where MgO proved to be unreactive under these conditions. From a thermodynamic viewpoint, the reaction between CO₂ and pure MgO should not occur at 500 °C and 50 bar CO₂ (173 bar CO₂ is required, Fig. 1b and reaction (1)).⁴¹ This has previously been shown in the literature for MgO, as discussed in the introduction. However, the formation of dolomite from CaO, MgO, and CO₂ only requires 8 bar of CO₂ at 500 °C (Fig. 1b and reaction (3)).⁴¹ Hence, the dolomite formation reaction is a thermodynamically destabilised pathway. Unfortunately, it is apparent that dolomite is not formed at 500 °C and 50 bar CO₂ due to a kinetic barrier, despite favourable thermodynamics.

The addition of a novel molten salt catalyst (NaCl/MgCl₂) to the synthetic batch (Synth-C-cat) allows for near complete conversion to dolomite (Table 3, Fig. S4†). The (0 1 5) Bragg ordering reflection for dolomite is present at ~18.8° 2θ in the synthetic dolomite diffraction data displayed in Fig. 2, which would not be present for magnesium-rich calcite, thus proving the formation of crystallographically ordered dolomite.³⁶ The molten salt acts as a catalyst to dolomite formation, overcoming the kinetic barrier to CO₂ absorption, presumably due to its capability of dissolving reactants. Thus, the metal halide eutectic salt mixture is an effective carbonation catalyst for dolomite.

Fig. 2 illustrates *in situ* powder XRD patterns obtained during thermolysis of the dolomite samples Rich-A, Wath-A, and Synth-C-cat, heated to 850 °C under argon. During thermal treatment the dolomite gradually decomposes to form MgO and CaCO₃. The commercial dolomite samples begin decomposition at higher temperatures (Rich-A: 550 °C, Wath-A: 800 °C) than the synthetic dolomite (Synth-C-cat: 500 °C), which also decomposes more gradually. Interestingly, both larnite (Ca₂SiO₄) and Ca₃Al₂O₆ are not observed at these temperatures, in contrast to their large crystalline fraction after heat treatment at 1000 °C (in Rich-B and Wath-B). This is advantageous as calcium is still present as CaCO₃ and can still react to form dolomite upon re-carbonation. The synthetic sample Synth-C-cat is shown in Fig. 2c, with the RT diffraction pattern illustrating the high dolomite content, with traces of received, by X-ray diffraction (Fig. S1,† Table 3), primarily showing dolomite, CaMg(CO₃)₂, but also a minor content of MgO and the salt-based catalyst. One of the salt phases is hydrated after air exposure, but more interestingly it has the composition CaMg₂Cl₆(H₂O)₁₂. The originally added catalytic salt was a mixture of NaCl and MgCl₂, but CaCl₂ forms during calcination, likely through a known metathesis reaction between MgCl₂ and CaO.³⁷ The

NaCl : MgCl₂ eutectic has a melting point of 442 °C (0.562 : 0.438), but there is also an eutectic in the NaCl : CaCl₂ system at 504 °C (0.479 : 0.521),³⁸ which may also act as an active carbonation catalyst.

Fig. 3 illustrates TGA data collected for each of the dolomite samples from the *in situ* powder XRD study (Rich-A, Wath-A, Synth-C-cat). Each of the samples indicate a lower temperature of decomposition than previously observed for natural dolomite (typically 700–850 °C) under argon flow.^{39,40} This is most likely due to the heating rate, the quantity of impurities, and the particle size variation between various batches of dolomite. This variation has been well documented in previous studies.^{39,40} The mass losses measured during heating are consistent with the expected mass losses calculated from Rietveld analysis (Table 3) assuming full decomposition of dolomite. The synthetic sample (Synth-C-cat) loses 36 wt%, compared to the theoretically expected 38 wt% (considering the 20 wt% catalyst). The Richgro and Watheroo samples also show mass losses close to their expected content (Rich-A: 35 wt% loss when expecting 34 wt%, and Wath-A: 44 wt% loss when expecting 44 wt%). The decomposition kinetics of all three samples differ dramatically. The Watheroo batch (Wath-A) exhibits a rapid single-step decomposition near 700 °C, with a minor mass loss at 550 °C. Whereas, the Richgro batch (Rich-A) exhibits a slow single-step mass loss from 500–700 °C. The synthetic catalysed dolomite (Synth-C-cat) exhibits a two-step decomposition (at 475 and 650 °C) before a slight mass loss at high temperature (>800 °C), which is likely due to vaporisation of the molten salt. It is clear that complete calcination is observed in all samples by 800 °C.

To assess the cyclic CO₂ capacity and thermochemical reversibility of the samples, cyclic pressure measurements were performed (Fig. 4, S5 and Table S1†). Since the reformation of dolomite only occurs with the addition of 20 wt% NaCl : MgCl₂ as a catalyst, cycling tests were performed on catalysed versions of calcined samples, starting from the oxide-rich/CO₂ depleted state (Synth-B-cat and Wath-B-cat). The Richgro sample (Rich-B) was omitted due to its very low content of CaO (5.1 wt%, see Table 3).

Absorption of CO₂ in the Watheroo sample (Wath-B-cat, Fig. 4 and Table S1†) reveals that more CO₂ is absorbed on the first cycle compared to the following 10 cycles. The first cycle involves carbonation of CaO, which is not formed again during CO₂ cycling at these pressures and temperatures, where instead CaCO₃ is the more thermodynamically stable compound. The quantity of CO₂ absorbed and desorbed during subsequent cycles is consistently ~50% of the theoretical capacity (of reaction (3)) over 5 h. For the eighth absorption, the temperature was reduced to 500 °C and again on the tenth cycle to 450 °C, for which a slight increase in cyclic capacity is observed (~60% of theoretical capacity). For each of the 11 cycles, the kinetics of absorption and desorption are both slow, with an absorption plateau not being reached within the five hours of data collection, likely due to a diffusion limited process.

After cycling, XRD of the sample in the absorbed state (Wath-D-cat, Table 3, Fig. S6†) indicates that it contains 31.0 wt% dolomite, 19.1 wt% MgO, 33.0 wt% CaCO₃, along with 8.7 wt%





Table 3 Dolomite samples before and after treatment. Quantitative composition of crystalline component (wt%) provided from X-ray diffraction based Rietveld analysis, with major phases shown. Fitting uncertainty of last decimal point in parentheses

	Carbonates and oxides						Silicates and aluminates						Salts			
	Dolomite CaMg(CO ₃) ₂	Calcite CaCO ₃	Magnesite MgCO ₃	Lime CaO	Magnesia MgO		Montmorillonite ^a	Gehlenite ^e	Andradite ^a	Ca ₃ Al ₂ O ₆	SiO ₂	Quartz	Kaolinite Al ₂ Si ₂ O ₅ (OH) ₄	Larnite Ca ₂ SiO ₄	Halite NaCl	CaMg ₂ Cl ₆ (H ₂ O) ₁₂
As supplied																
Rich-A: Richgro dolomite	70.8(6)	9.8(2)	—	—	—	—	4.5(4)	—	—	5.2(2)	8.3(6)	—	—	—	1.31(5)	—
Wath-A: Watheroo dolomite	91.8(6)	2.37(9)	3.0(1)	—	—	1.1(2)	—	—	—	1.70(7)	—	—	—	—	—	—
Thermally outgassed																
Rich-B: Richgro (Rich-A: 1000 °C, Ar)	—	—	—	5.1(1)	33.6(3)	—	—	5.0(5)	—	3.87(9)	—	—	47.8(4)	0.7(1)	—	—
Wath-B: Watheroo (Wath-A: 1000 °C, Ar)	—	—	—	27.8(3)	34.8(4)	—	—	—	6.7(4)	1.31(4)	—	—	29.4(6)	—	—	—
Synth-B: synthetic (650 °C, Ar)	—	—	—	57.7(2)	42.3(2)	—	—	—	—	—	—	—	—	—	—	—
Carbonated (500 °C, single-step 50 bar CO₂)																
Wath-C: Watheroo (Wath-B, no catalyst)	—	47.0(3)	—	—	28.9(2)	—	—	—	1.3(2)	1.62(6)	—	—	21.2(3)	—	—	—
Synth-C: synthetic (Synth-B, no catalyst)	—	57.6(5)	—	11.4(1)	31.0(3)	—	—	—	—	—	—	—	—	—	—	—
Synth-C-cat: synthetic + catalyst (Synth-B, NaCl : MgCl ₂)	93.0(1)	—	—	—	2.07(6)	—	—	—	—	—	—	—	—	3.09(7)	—	1.9(1)
CO₂ cycled																
Wath-D-cat	31.0(2)	33.0(9)	—	—	19.1(6)	—	—	—	—	0.4(1)	—	—	—	6.4(4)	—	1.5(4)
Synth-D-cat	14.3(2)	49.3(2)	—	—	26.8(2)	—	—	—	—	—	—	—	—	6.2(1)	—	3.4(2)

^a Montmorillonite is (Na,Ca)_{0.33}(Al,Mg)₂(Si₄O₁₀)(OH)₂·*n*H₂O, gehlenite is Ca₂Al₂SiO₇, andradite is Ca₃Fe₃Si₃O₁₂.

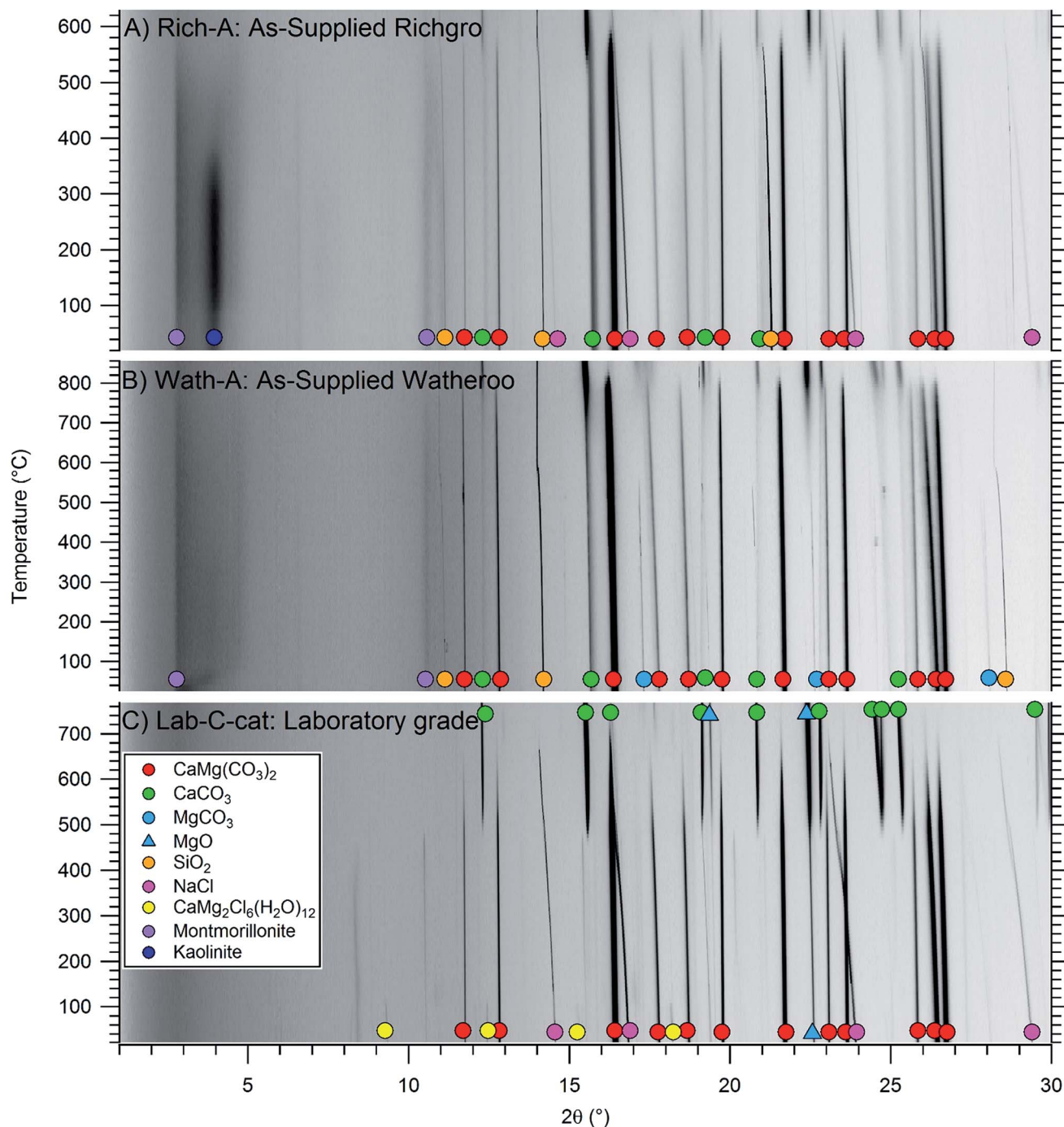


Fig. 2 *In situ* powder X-ray diffraction of (A) Richgro, (B) Watheroo and (C) Laboratory grade dolomite samples heated at 6 °C min^{-1} under 1 bar Ar. ($\lambda = 0.8263076\text{ \AA}$). The same primary end products are formed in all three samples under these conditions of temperature and time.

andradite, 0.4 wt% quartz, 6.4 wt% halite and 1.5 wt% CaCl_2 . These results illustrate that the catalyst enables dolomite formation, similar to the synthetic sample (Synth-C-cat), in contrast to the single-step carbonation at 500 °C (Wath-C, Table 3).

The synthetic sample (Synth-B-cat) was cycled 14 times at 550 °C (Synth-D-cat, Fig. S5 and Table S1†). The initial carbonation under 35 bar CO_2 showed ~ 1.75 mol. equivalent absorption (87.5% theoretical capacity, including the

carbonation of CaO) although subsequent absorptions only allowed for ~ 0.25 mol equivalent of CO_2 uptake (25% theoretical capacity). Complete carbonation was again not observed, even after 12–24 h, showing that the reaction kinetics are very slow. XRD of the sample after the final CO_2 absorption cycle revealed it contained CaCO_3 (49.3 wt%), MgO (26.8 wt%), dolomite (14.3 wt%), NaCl (6.2 wt%) and CaCl_2 (3.4 wt%) (Fig. S6†).



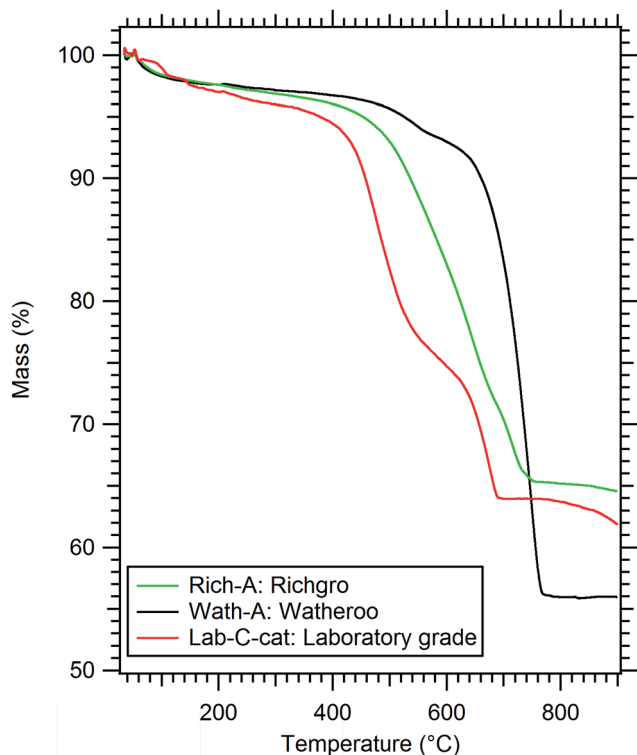


Fig. 3 Thermogravimetric analysis (TGA) of dolomite materials under 1 bar Ar flow, heated at $10\text{ }^{\circ}\text{C min}^{-1}$.

Carbon dioxide cycling of dolomite is therefore possible for both mined and synthetically synthesised materials with the aid of a catalyst, but in this study the commercial material (Watheroo) has a larger cycling capacity of $\sim 50\%$ compared to $\sim 25\%$ of theoretical capacity for the synthetic material. The better cyclability could be due to the additional impurities contained in the

commercial material (quartz, kaolinite and montmorillonite), which also resulted in reasonable carbonation over a 5 h window at $450\text{--}500\text{ }^{\circ}\text{C}$. The geological process of dolomitisation is known to form dolomite when Mg^{2+} replaces Ca^{2+} in calcite.⁴¹ This process has been studied in detail with many researchers attempting to form dolomite in the direct crystallisation from an aqueous solution at temperatures between $60\text{--}220\text{ }^{\circ}\text{C}$.⁴¹ This crystallisation process has been shown to occur on a much longer timescale (hours to days), *via* an Ostwald-ripening mechanism.

To further understand the difference in cyclic capacity between synthetic and mined dolomite samples microscopy investigations were undertaken on CO_2 cycled samples (Synth-D-cat and Wath-D-cat). SEM analysis reveal that the two samples are morphologically different (Fig. 5). Watheroo powders were found to be porous aggregates of fine particles (Fig. 5a and b), displaying higher surface area and better gas migration pathways than synthetic samples (Fig. 5e and f), which were less porous, and more monolithic. However, the drastic differences in phase segregation observed in both samples is most important. The Watheroo sample, as shown by XRD, contains a high SiO_2 content that may be present as quartz, silicates or aluminosilicates, whereas the synthetic sample has greater purity and is Si-free (Table 3). Imaging and chemical analysis of particle cross-sections (Fig. 5c, d and g, h) reveal that the Watheroo samples have intermixed Ca- and Mg-rich components in sub-micron domains, which also include Si-rich compounds. Whereas, the synthetic sample shows larger discrete Ca- or Mg-rich domains indicating a much higher degree of phase segregation ($>5\text{ }\mu\text{m}$). Intermixing of Si-rich compounds with MgO and CaO may act to arrest sintering, agglomeration, and segregation of CaO and MgO. Addition of unreactive phases has previously been shown to act as particle refinement agents in gas-sorption materials, which increases their overall cyclability.⁴² It is feasible that the Si-rich impurities

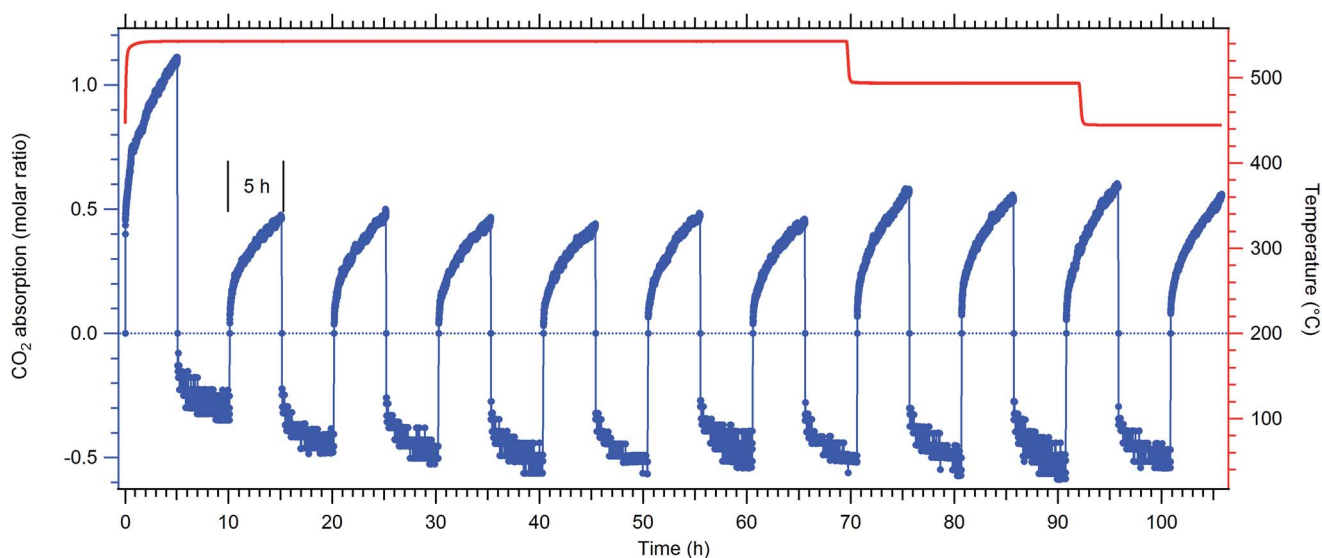
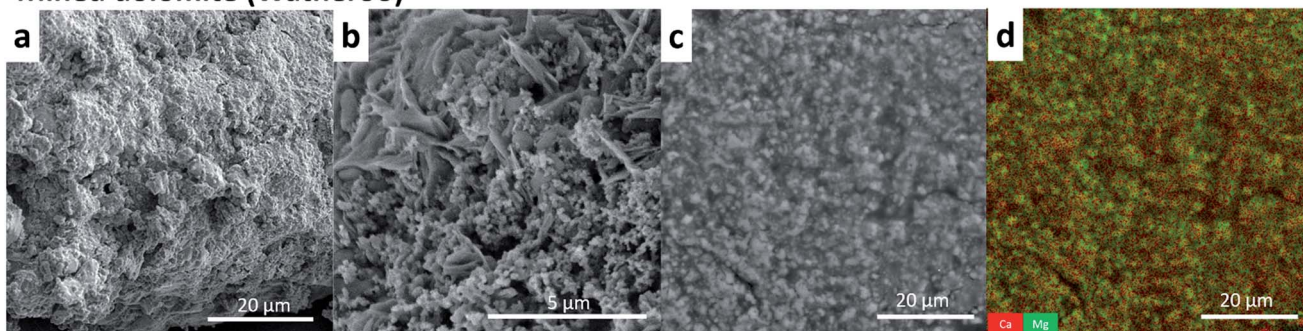


Fig. 4 Carbon dioxide pressure cycling results for catalysed ($\text{NaCl}:\text{MgCl}_2$) Watheroo dolomite (Wath-B-cat) at multiple temperatures (absorption pressure 35 bar, desorption pressure 2 bar). Blue and red curves represent CO_2 absorption/desorption values and temperature, respectively.



Mined dolomite (Watheroo)



Synthetic dolomite

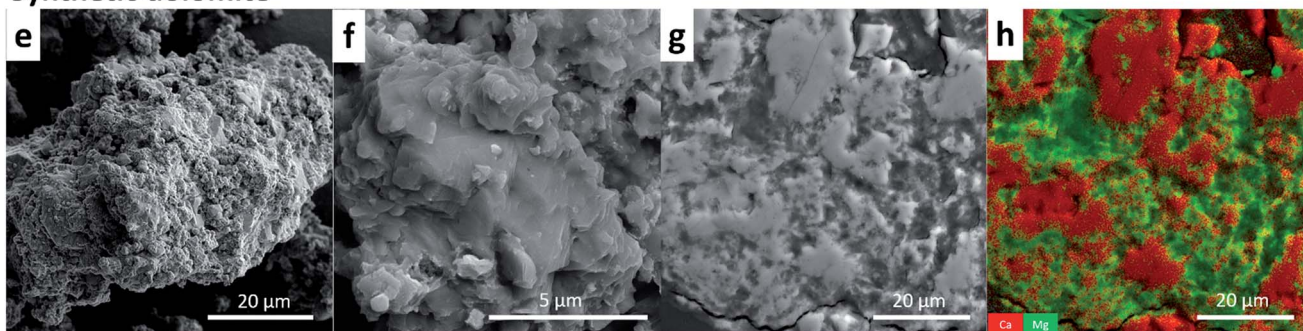


Fig. 5 SEM micrographs of CO₂ cycled mined dolomite (Watheroo, Wath-D-cat, (a–d)) and synthetic dolomite (Synth-D-cat, (e–h)) with secondary electron (SE) or backscattered electron detectors (BSE). (a/e) Typical conglomerated particle (SE), (b/f) higher magnification image of the particle surface (BSE) (c/g) polished cross-section of a powder grain (BSE), (d/h) EDS elemental map of an epoxy embedded cross section (coloured elemental maps overlaid onto BSE image, colour legend in the image).

act as grain size refinement agents and reduce the path length between CaO and MgO reaction sites during dolomite formation, while also retaining a porous morphology, which allows CO₂ migration through the sample.

Nitrogen (N₂) adsorption/desorption measurements (Table S2[†]) are consistent with the SEM observations that the synthetic and mined dolomite samples are morphologically different. The synthetic dolomite (Synth-B) has 0.98 m² g⁻¹ specific surface area and 0.0014 cm³ g⁻¹ cumulative pore volume. Mined dolomite samples, Rich-A and Wath-A, have a specific surface area of 16.90 and 16.78 m² g⁻¹, respectively, which is >16 times higher than the synthetic sample (Synth-B). Their cumulative pore volume is also ~10 times higher than the synthetic dolomite sample, 0.0155 and 0.0146 cm³ g⁻¹.

Conclusions

This study has illustrated the reversible carbonation of dolomite, demonstrating its potential as a 2nd generation concentrating solar thermal power (CSP) thermochemical energy storage material. This was achieved through the use of a novel molten salt catalyst. Three different dolomite sources were assessed for thermochemical energy storage for CSP applications. Catalysed dolomite samples were shown to cycle (~50% capacity) over >10 + cycles from 450–550 °C without additional loss in capacity. This operational temperature range is the same as existing molten nitrate salt energy storage systems but with over an order of magnitude in raw material cost reduction, even

when considering partial CO₂ uptake capacity. These promising results demonstrate that thermochemical energy storage should be targeted for 2nd-generation CSP plants. In fact, metal carbonates offer an incredibly low cost for energy stored per dollar, especially when considering that refining of mined dolomite (and their associated cost) may not be required. Herein, it is shown that impurities within dolomite improve their thermochemical cycling performance, and unrefined mined samples show the most promising results once catalysed by inexpensive salts. Essentially, dolomite could be extracted from the ground and directly deposited into a thermochemical reactor.

Future research should be directed towards improving the cyclic capacity of dolomite, investigating other promising metal carbonate systems, and investigating the design/cost of high temperature reactors to house the powder bed. In addition, low cost volumetric storage options must be identified. This would include liquefying the CO₂ at pressures above 64 bar or the use of low-temperature CO₂ storage in another carbonate material.

Conflicts of interest

There are no conflicts to declare.

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