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Article

Experimental Investigation and Simplistic Geochemical Modeling of CO₂ Mineral Carbonation Using the Mount Tawai Peridotite

Omeid Rahmani ^{1,*}, James Highfield ², Radzuan Junin ^{3,4}, Mark Tyrer ⁵ and Amin Beiranvand Pour ⁶

- Department of Petroleum Engineering, Mahabad Branch, Islamic Azad University, Mahabad 59135-433, Iran
- ² 560 Yishun Avenue 6 #08-25 Lilydale, Singapore 768966, Singapore; James03@singnet.com.sg
- Department of Petroleum Engineering, FCEE, Universiti Teknologi Malaysia (UTM), Skudai 81310, Johor, Malaysia; radzuan@petroleum.utm.my or r-razuan@utm.my
- ⁴ Universiti Teknologi Malaysia-Malaysia Petroleum Resources Corporation (UTM-MPRC), Institute for Oil and Gas, Universiti Teknologi Malaysia (UTM), Skudai 81310, Johor, Malaysia
- Mineral Industry Research Organisation, Wellington House, Starley Way, Birmingham International Park, Solihull, Birmingham B37 7HB, UK; m.tyrer@mtyrer.net
- Geoscience and Digital Earth Centre (Geo-DEC), Research Institute for Sustainability and Environment (RISE), Universiti Teknologi Malaysia (UTM), Skudai 81310, Johor, Malaysia; a.beiranvand@utm.my or beiranvand.amin80@gmail.com
- * Correspondence: omeid.rahmani@iau-mahabad.ac.ir or omeidrahmani@gmail.com; Tel.: +98-914-442-2009

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Abstract: In this work, the potential of CO_2 mineral carbonation of brucite $(Mg(OH)_2)$ derived from the Mount Tawai peridotite (forsterite based $(Mg)_2SiO_4$) to produce thermodynamically stable magnesium carbonate $(MgCO_3)$ was evaluated. The effect of three main factors (reaction temperature, particle size, and water vapor) were investigated in a sequence of experiments consisting of aqueous acid leaching, evaporation to dryness of the slurry mass, and then gas-solid carbonation under pressurized CO_2 . The maximum amount of Mg converted to $MgCO_3$ is ~99%, which occurred at temperatures between 150 and 175 °C. It was also found that the reduction of particle size range from >200 to <75 μ m enhanced the leaching rate significantly. In addition, the results showed the essential role of water vapor in promoting effective carbonation. By increasing water vapor concentration from 5 to 10 vol %, the mineral carbonation rate increased by 30%. This work has also numerically modeled the process by which CO_2 gas may be sequestered, by reaction with forsterite in the presence of moisture. In both experimental analysis and geochemical modeling, the results showed that the reaction is favored and of high yield; going almost to completion (within about one year) with the bulk of the carbon partitioning into magnesite and that very little remains in solution.

Keywords: CO₂ sequestration; forsterite; *ex-situ*; *in situ*; mineral carbonation

1. Introduction

Carbon dioxide (CO₂) is the principal greenhouse gas released into the atmosphere during fuel combustion, particularly due to the extensive use of fossil fuels for energy production from coal, oil, and natural gas since the industrial revolution [1]. Moreover, atmospheric CO₂ has recently surpassed 400 ppm, and is predicted to increase to nearly 1000 ppm by the end of the 21st Century [2,3]. The related global temperature rise (exceeding 2 $^{\circ}$ C) [3] will almost certainly result in irreversible climate change with potentially disastrous consequences.

Researchers have studied ways to mitigate the amount of greenhouse gases released into the atmosphere by sequestration of CO₂ through different approaches, including aquifer storage, deep

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sea storage, and mineral carbonation in particular [1,4–14]. Minerals and rocks rich in magnesium (Mg^{2+}) /calcium (Ca^{2+}) are commonly considered as candidates due to their wide availability, low cost, and environmentally benign nature. [8,12,15] During mineral carbonation, CO_2 reacts with Mg^{2+} or Ca^{2+} -rich minerals (e.g., olivine and gypsum) to form solid carbonates, which are expected to be stable over geologic time periods. The minerals olivine $[(Mg,Fe)_2SiO_4]$ and forsterite $[Mg_2SiO_4]$, containing up to 33.6 wt. % Mg, have the highest capacity to trap CO_2 as magnesium carbonates, and a high rate of dissolution among rock-forming silicate minerals [16]. Formation of magnesite from forsterite, the Mg-end member of the olivine solid solution series, is thermodynamically favorable based on the negative Gibbs free energy of Reaction (1).

$$1/2Mg_2SiO_4 + CO_2 \rightarrow MgCO_3 + 1/2SiO_2 + 95 kJ/mol$$
 (Rx. 1)

According to Lackner *et al.* [17] the chemical reactions in mineral carbonation process can be very slow under ambient conditions and, therefore, activation processes such as exposure to acid, heat [8], and water [18–21] have been used to accelerate the carbonation rate. These processes are tedious and require additional energy input, which has made acid/heating approach less attractive than some others. The rates of carbonation can be raised by increasing surface areas of the mineral or its intermediates and by elevating the temperatures, resulting in lower kinetic constraints [15,16,22–29]. Moreover, an *in situ* CO₂ sequestration system with mineral carbonation can be treated as a fully-coupled problem between rock deformation, pore-fluid flow, heat transfer, mass transport, and chemical reaction processes. Three types of models are commonly employed in computational petroleum geoscience and engineering research methodologies. These approaches are geological/geochemical (conceptual), mathematical, and/or numerical simulation.

The purpose of this study was to do *ex situ* (laboratory) studies of the factors affecting the rates of (forsterite) mineral carbonation (*i.e.*, particle size, water vapor, and reaction temperature) in support of an *in situ* geochemical conceptual model. A transport reaction modeling software PHREEQC (version 2.18, US Geological Survey (USGS), Reston, WV, USA), was applied for simulating the chemical reactions and transport processes in the forsterite mineral carbonation process. Pre-treatment of forsterite in a quantitative equivalent of mineral acid (HCl), *i.e.*, dissolution to neutrality, was taken as a suitably fast process (in extracting most of the Mg ion) for practical experimentation. One advantage of this process is that the formation of magnesium carbonate (MgCO₃) releases heat, which can in principle be cycled back to other endothermic steps (see Reactions (2)–(4) in Section 3.3) through heat integration in a commercial process.

2. Results and Discussion

2.1. Mineral Characterization

The elemental composition of the peridotite mineral was determined by XRF analysis as MgO (51.9%), SiO₂ (41.1%) as major components with minor levels of FeO, Al₂O₃, Na₂O, K₂O, and CaO (see details in Table 1).

Al ₂ O ₃	CaO	FeO	MgO	K ₂ O	SiO ₂	Na ₂ O	Cr ₂ O ₃	Volatiles	
								C + CO ₂	H ₂ O

0.204 0.061 5.969 51.921 0.005 41.072 0.083 0.034 < 0.352 0.291 Number of ions on the basis of O Al Ca Fe K Si Cr Na Na Mg 0.004 0.001 0.001 0.161 1.812 0.004 0.995 0.008

Table 1. Chemical composition of fresh peridotite mineral (wt. %) as determined by XRF analysis.

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XRD analysis of the HCl-cleaned starting material (Figure 1) shows a characteristic pattern of olivine ($2\theta = 11.8^{\circ}$, 23.6° , 29.3° , 31.1° , 33.6° , 40.8° , and 43.6°), with some contamination by quartz or free silica ($2\theta = 20.9^{\circ}$, 26.5° , 50.5° and 68.7°). In view of the fairly low level of Fe and its own capacity for carbonation (as FeCO₃), modeling studies (*vide infra*) were based for simplicity on the pure forsterite composition (Mg₂SiO₄), the Mg-end member of the olivine solid solution series.

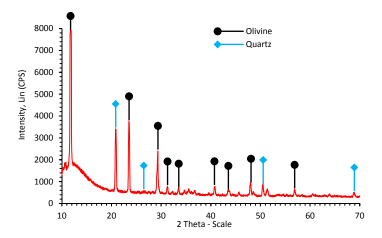


Figure 1. XRD pattern of the starting peridotite mineral.

Morphological and structural changes in the olivine (nominal composition $Mg_{1.84}Fe_{0.16}SiO_4$) at various stages during chemical pretreatment and carbonation, viz., after leaching in HCl, neutralization and precipitation of Mg^{2+} [as $Mg(OH)_2$], and exposure to humid CO_2 at 4.8 bar and 150 °C, are best seen by SEM micrographs and powder XRD in Figures 2 and 3 respectively.

By SEM, the fresh olivine sample (Figure 2a) consisted typically of polycrystalline grains in the millimeter size range. After acid leaching and neutralization of the sieved fraction <75 μ m (Figure 2b), the grains became finer (<5 μ m) and the structure more amorphous. In agreement with the findings of Bearat *et al.* [30] and Kwon *et al.* [18], this is likely due to an amorphous SiO₂ residue after leaching, any (hexagonal) brucite likely being nanocrystalline. Under humid CO₂, the development of polyhedral and more sheet-like crystallites (Figure 2c) was evident, probably representing (hydrated) magnesium carbonate, and finally (Figure 2d), the rhombohedral habit typical of magnesite (MgCO₃) was evident.

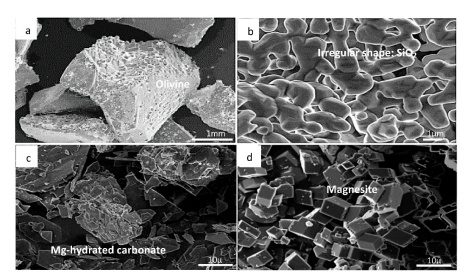


Figure 2. SEM images showing (**a**) the fresh olivine mineral, and morphological changes during chemical pretreatment and carbonation: (**b**) the leached/neutralized sample in the presence of humid CO₂ at 150 °C during 15 min; (**c**) 90 min; and (**d**) 120 min.

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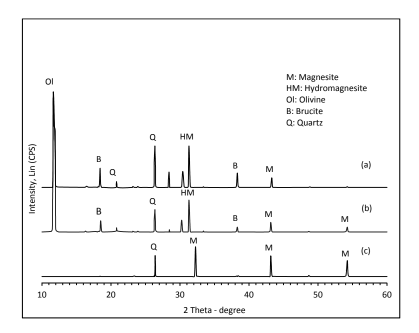


Figure 3. XRD patterns showing structural changes in the olivine mineral during progressive carbonation: (a) after acid leaching, neutralization, and initial exposure of the damp residue to humid CO_2 ($P_{CO2} = 4.8$ bar, T = 150 °C) during 15 min; (b) after 90 min; and (c) after 120 min.

Figure 3 shows XRD data obtained after the same stages of treatment in parallel with the SEM analyses. Diffractogram 3a reveals that although unreacted olivine ($Mg_{1.84}Fe_{0.16}SiO_4$ main reflection at $2\theta = 11.8^{\circ}$) is predominant after only 15 min exposure to humid CO_2 , hydromagnesite ($HM = Mg_5(CO_3)_4(OH)_2 \bullet 4H_2O$ main reflection at $2\theta = 31.3^{\circ}$) and a little magnesite ($M = MgCO_3$ main reflection at $2\theta = 43.15^{\circ}$), had already formed from brucite [$Mg(OH)_2$] precursor, itself created in the leaching/neutralization stage. At longer exposure times (Figure 3b,c), reflections due to brucite and hydromagnesite were progressively replaced by those of magnesite ($2\theta = 32.25^{\circ}$, 43.15° , 54.25°). The ultimate disappearance of olivine is intriguing since it implies direct steam-activated carbonation of the (chemically un-pretreated) mineral, for which evidence has been reported elsewhere [31]. The abundance of quartz (as co-product) cannot be taken as a reliable indicator of the progress of carbonation because it is also a phase contaminant in the original mineral (see Figure 1). Furthermore, similar chemical treatment (flux extraction of Mg^{2+}) from serpentinites did not produce quartz but instead a silica residue of unusual structure [27]. The unsystematic peak intensity of quartz seen here by XRD may be due merely to local inhomogeneity in the samples.

2.2. Effect of Particle Size on the Carbonation of Olivine

The leaching rate was found to be much slower than the optimal carbonation rate (achievable at $175\,^{\circ}\text{C}$ *vide infra*), such that the production of magnesite in these experiments reflects mainly variations in the rate and efficiency of extraction of soluble Mg^{2+} ion by acid treatment. Figure 4 shows that decreasing the grain size from >200 μ m to <75 μ m caused the limiting degree of Mg^{2+} leaching to increase from 35% to 99%, respectively. It is well-known and intuitively obvious that the reaction rate and carbonation degree can be raised by increasing the surface area, e.g., by grinding/sieving, as shown for example by Garcia *et al.* [32]. In this work, the smallest particle size (d < 75 μ m) is the only one offering the prospect of full leaching (99%) in a practical time. As a rule of thumb, 1–2 h for a terrestrial (*ex-situ*) process to achieve >90% conversion is taken as a realistic practical target to limit the scale (and associated costs) of any future installation for CO_2 sequestration. This particle cut and leaching procedure (2 h at 60 $^{\circ}$ C) were therefore applied as standard for all samples in subsequent tests described below.

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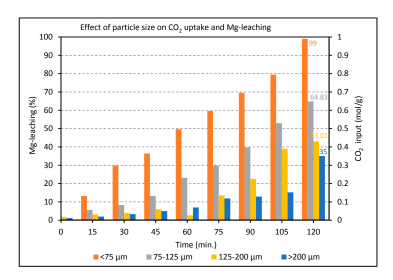


Figure 4. Effect of particle size on Mg^{2+} leaching over a range of time intervals as measured by the volume of CO_2 uptake in the subsequent mineral carbonation process (at 175 °C and 2 h).

2.3. Effect of Temperature on the Carbonation of Olivine

The temperature dependence of the carbonation process was studied over the interval from ambient to 175 °C. The effect of temperature on the amount of Mg conversion after standard leaching treatment (99% extraction) is illustrated in Figure 5. As expected, the temperature had an important effect on the mineral carbonation process consisting of Mg extraction and subsequent MgCO₃ precipitation. The amount of Mg converted at the desired temperatures was measured continuously several times. The maximum extent of carbonation of (99%) was attained at temperatures in excess of 150 °C. The quantification of the MgCO₃ formed in the mineral carbonation process was carried out by titration against HCl at room temperature. Pokrovsky *et al.* [33] demonstrated that the dissolution rate of magnesite at 150 °C is lower than at 25 °C whereas the rates at 100 and 150 °C in acidic solutions are almost the same [34], suggesting a strong decrease of the apparent activation energy above 100 °C. According to Saldi *et al.* [34] the tendency for dissolution rates of MgCO₃ to decrease with increasing temperature could benefit CO₂ sequestration efforts by making magnesite more resistant to dissolution in deeper (hotter) strata, thus preserving the petrophysical integrity of deep carbonate-rich confining reservoirs.

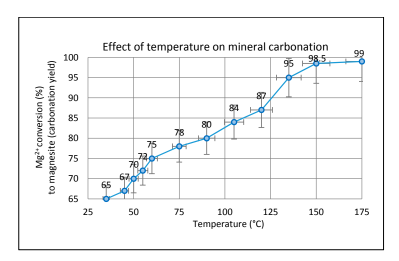


Figure 5. The effect of reaction temperature in gas-solid carbonation of fully extracted Mg^{2+} ion (as $Mg(OH)_2$) to yield $MgCO_3$ ($P_{CO2} = 4.8$ bar, humid CO_2 , t = 2 h).

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It is clear from Figure 5 that several competing factors in the carbonation process reach equilibrium in the temperature range between 150 and 175 $^{\circ}$ C. Probably the key factor is the relative humidity (RH). Previous work has shown that carbonation accelerates on cooling below 200 °C in a fixed partial pressure of steam or RH ≥ 20% [35]. Since the presence of liquid water has been shown to be important, it can also be argued that the effect of temperature on the solubility of CO₂ may influence the carbonation rate. Chen et al. [36] indicated that increasing temperature from 25 °C to 150 °C generally helps the precipitation of magnesite. The carbonate solubility product (K_{sp}), Henry's constant (K_H), and the first- and second-order dissociation constants of carbonic acid (Ka₁, Ka₂) are all functions of temperature. Increasing the reaction temperature decreases the value of K_{sp} and increases the K_H value, which lowers the amount of CO₂ gas in solution at a given pressure. The dissociation constants of carbonic acid (Ka₁ and Ka₂) are also increased by increasing temperature and promoting solution speciation and carbonation. Thus, increasing temperature has conflicting effects, lowering the level of dissolved CO₂ gas via its effect on K_H but promoting magnesite precipitation via effects on Ka₁, Ka₂, and K_{sp}. Furthermore, temperature impacts on the kinetics of magnesite precipitation [37] and also affects the type of Mg-carbonate formed. It is well-known that magnesite precipitation kinetics at ambient temperatures are exceedingly slow, and that metastable hydrated carbonates such as hydromagnesite, dypingite, and nesquehonite almost invariably form instead.

2.4. Effect of Water Concentration

Formal carbonation of forsterite [Reaction (1)] does not involve water molecules explicitly, so any beneficial effect of added water is clearly a kinetic effect. As shown in Figure 6, an indication of the importance of water is seen in the modest level of CO₂ removal in its absence. By analogy with recent works on brucite and serpentinites, almost regardless of the CO₂ pressure utilized, the presence of water vapor in high relative humidity appears crucial to obtaining practical carbonation rates [20,31,38], evidently by establishing a highly polar thin-film aqueous overlayer that facilitates CO₂ ingress into the bulk particle. This is supported by independent studies simulating in situ or geochemical carbon sequestration where CO₂ was assumed to be in the supercritical state. Felmy et al. [39] studied high-surface-area forsterite in the presence of water-saturated scCO₂. They concluded that the nature of the water in contact with the reacting surface is a key factor in the enhanced magnesite formation. When excess water was added to the forsterite particles, a thin water film was formed on the forsterite surface promoting magnesite formation. Loring et al. [40] declared that this water film provides a distinctive situation for the magnesite formation by decreasing the effective Mg²⁺ dehydration energy and simplifying the transformation of nesquehonite to magnesite. Otherwise, the presence of liquid water can allow the formation of magnesium bicarbonate in solution that decomposes upon drying to magnesium carbonate. Moreover, Schaef et al. [41,42] revealed that the addition of water to the saturated system noticeably increases the rate of mineral carbonation, facilitating the overall conversion of nesquehonite to magnesite. No evidence of further carbonation was observed under unsaturated conditions below 50 °C. A similar promoting effect of water on brucite carbonation under scCO₂ was reported by Loring et al. [43] using in situ Fourier-Transform Infrared (FTIR) spectroscopic experiments.

To investigate the effect of humidity, the concentration of water vapor was set at various levels, 5, 10, and 20 vol % prior to carbonation of the brucite extract at 175 °C. As illustrated in Figure 6, above 5 vol % steam, the degree of carbonation increases by almost double in the presence of water vapor such that within 5 min, complete removal of CO_2 (15 vol %) was achieved. However, levels of water vapor exceeding 20 vol % had no additional effect on the rate of removal of CO_2 . It can be concluded that water vapor is able to solvate CO_2 , generate carbonate ions and protons [44], and increase the carbonation degree of $Mg(OH)_2$ as derived from olivine. Moreover, the aforementioned modest uptake of CO_2 (~7 vol %) under "dry" conditions may be due to adventitious water not fully removed from the $Mg(OH)_2$ -containing residue. Vitillo [45] declared that in the presence of water vapor, $MgCO_3$ crystalline phase reappeared increasingly, while the magnesium oxide periclase (MgO) phase gradually disappeared. These observations are well in agreement with the thermodynamic

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data on MgO, Mg(OH)₂, and MgCO₃ systems. The promoting effect of water may be attributed to faster reaction kinetics by offering alternative routes to magnesite via hydrocarbonate intermediates such as dypingite.

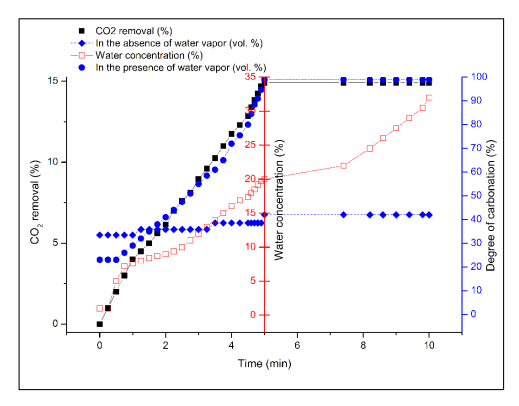


Figure 6. Effect of the presence or absence of water vapor on the rate of CO_2 absorption at 175 °C on 10 g Mg_2SiO_4 (d < 75 μ m) activated by chemical pre-treatment and exposed to 0.5 L/min CO_2 gas ($P_{CO2} = 4.8$ bar, $P_{H2O} = 1.6$ –6.4 bar or 18%–72% RH).

As a comparison to related work on Mg(OH)₂ in the literature, Siriwardane and Stevens [46] reported good absorption kinetics and reasonable capacity for CO₂ in plug-flow reactor experiments over a promoted brucite (~3 mol CO₂/kg or 20 mol %) in "moist" helium at 200 °C. However, the promotional effect of water *per se* was not explored and the sorbent surface area was low (~2.5 m²· g⁻¹). This compares with our thermogravimetric work [38] showing that Mg(OH)₂ extracted from the mineral is typically obtained in high-surface-area form (~25 m²· g⁻¹). These are evidently important factors in attainment of almost quantitative (~100%) carbonation at lower temperature (150–175 °C) in this work, specifically the higher water levels utilized and the better dispersion of brucite derived from the mineral. Based on the thermodynamic equilibrium of Mg(OH)₂ formation from MgO, Siriwardane and Stevens [46] showed that is likely to form Mg(OH)₂ under the high steam environment, which accounts for the subsequent CO₂ uptake. It is important to note that the Mg(OH)₂ system has the far lower heat of sorption. This confirms that the regeneration heat (input) needed to displace CO₂ from MgCO₃ by water (to form Mg(OH)₂) is significantly lower than that required for the decomposition of MgCO₃ (to MgO + CO₂).

2.5. Kinetic Analysis of Mg Extraction by HCl

Different kinetic analyses including expressions for product layer diffusion, film diffusion, chemical reaction control, and a combination of chemical reaction control (Equations (4)–(7), respectively, in Section 3.5), were used to evaluate the integral rate data. The extent of forsterite dissolution, X_E , is taken as fitting parameter but this is actually measured from the amount of magnesite, *i.e.*, the extent of carbonation (= R_{CO2} in Equation (2)), because dissolution is much slower

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than carbonation (of the Mg(OH)₂ extract). Direct carbonation of unreacted forsterite is probably even slower. Thus, dissolution is rate-determining in the overall process. The two best-fit results are illustrated in Figure 7a,b, but the first, a combination of *chemical reaction control* and *product layer diffusion* (Equation (7)) provided the highest correspondence with the measured data. Thus, it can be concluded that a combination of chemical reaction control and product layer diffusion is rate-limiting for Mg extraction.

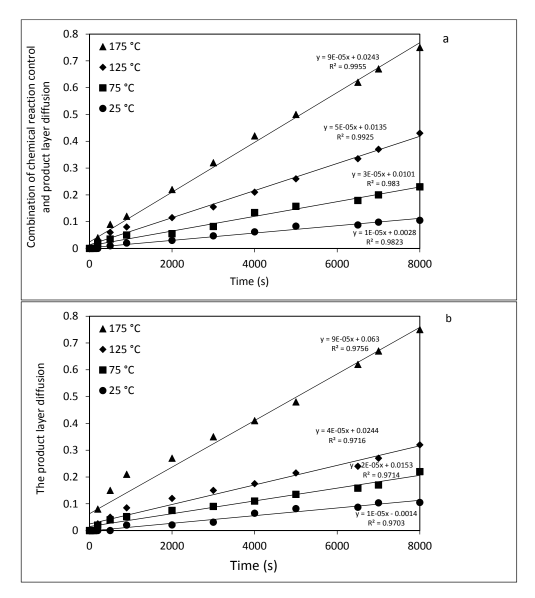


Figure 7. Kinetic analysis of olivine dissolution rate (in HCl) by plotting the combination of chemical reaction control and product layer diffusion (**a**) and the product layer diffusion (**b**) *vs.* time at various reaction temperatures.

Activation energies (E_a) for mineral dissolution were determined from simple log/log plots of the time-independent rate k at various temperatures. These values are presented in Table 2, from which the Arrhenius plots shown in Figure 8 were obtained. Once again, the quality of fit was best for the combination of product layer diffusion and chemical reaction control ($R^2 = 0.9917$) as compared to product layer diffusion only (R = 0.9764). Considering these models as the controlling mechanisms during the dissolution of forsterite, the E_a value was 15.5 kJ/mol for product layer diffusion control, and 16.0 kJ/mol for the combination of product layer diffusion and chemical reaction control. It is likely

that the chemical reaction is initially rate-limiting but product layer diffusion gradually becomes rate limiting as the product layer of silica builds up and the unreacted surface area decreases. According to Gharabaghi *et al.* [47] a low value of E_a indicates that product layer diffusion is rate-controlling, Therefore, considering the values of multiple regression coefficients for different models and calculated E_a for two selected models, it could be concluded that the dissolution rates of forsterite are kinetically regulated by the combination of chemical reaction control and product layer diffusion.

Kinetic Analysis	k	ln k	T (°C)	T (K)	1/T
	9.0243×10^{-5}	-9.313	175	448.15	0.00223
Combination of chemical reaction control and	5.0135×10^{-5}	-9.9007	125	398.15	0.00251
product layer diffusion	3.0101×10^{-5}	-10.4109	75	348.15	0.00287
	1.0028×10^{-5}	-11.5101	25	298.15	0.00335
	9.0635×10^{-5}	-9.3086	175	448.15	0.00223
Product layer diffusion	4.0244×10^{-5}	-10.1205	125	398.15	0.00251
i loduct layer diffusion	2.0153×10^{-5}	-10.8121	75	348.15	0.00287
	1.0014×10^{-5}	-11.5115	25	298.15	0.00335

Table 2. The rate constant calculation for every experiment at different temperatures.

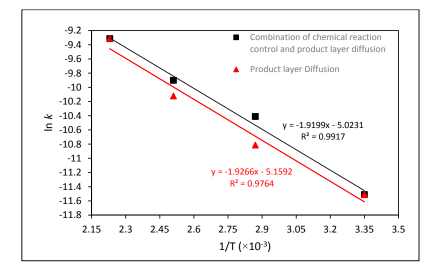


Figure 8. The Arrhenius plots for the extraction of Mg from forsterite using two selected models of the combination of product layer diffusion and chemical reaction control and the product layer diffusion.

The rate constants generated in this study on forsterite are faster than corresponding rate constants reported by Pokrovsky and Schott [48], who only worked at the temperature of 25 °C. Comparison of photomicrographs showing the surface of forsterite presented by Pokrovsky and Schott [48] and the samples from this study (see Figure 2) demonstrates they consist of euhedral and larger crystals due to small adhering particles and their agglomeration in acid solution at the temperature of 25 °C. In the present study, the rate constants are increased because of the far higher density of activated sites per unit surface area at higher temperatures.

2.6. Thermodynamic Considerations on the Mineral Carbonation

As regards the effect of temperature on the system, two physical mechanisms interact; the increase of olivine solubility with temperature and, conversely, the reduction of magnesite solubility with increasing temperature. Heat treatment was reported by O'Connor *et al.* [49] to remove sorbed water and activate the mineral surface. They suggested that the carbonation phase occurs quickly in the olivine powders at high temperatures, from which we expect an increase in the available reactivity of

mineral surface with removal of sorbed water and CO₂ uptake by the mineral surface and, therefore, more rapid reaction.

Additionally, elevated temperature causes an increase of the olivine dissolution rate, which enhances the overall reaction kinetics. Our results concur with this (see Figure 5) up to the maximum temperature studied (175 °C). The lowest CO₂ concentration considered was 0.5% (volume fraction of CO₂ unconsumed) indicating olivine samples have efficiently precipitated CO₂ as carbonate phase, which is in agreement with the findings of Kwon et al. [18]. Interestingly, the CO₂ sequestration capacity of olivine mineral seen here over the temperature range of 150–175 °C was higher than in previous studies [50–55]. We tentatively suggest that this effect is due to either an increase of the available surface area or available moisture (or a combination of both factors). At the highest temperatures, (e.g., the range of 150–175 °C), Hänchen et al. [56] suggest the olivine samples released more Mg into the solution, which seems a reasonable assumption to adopt here and is confirmed by the calculation shown below. At lower temperatures (less than 100 °C), King et al. [12] found that MgCO₃ precipitation was kinetically hindered due to the high activation energy for the de-solvation of the strongly hydrated Mg²⁺ ions (Reaction (2)). According to Hänchen et al. [57] during low temperature experiments, hydromagnesite [Mg₅(CO₃)₄(OH)₂•4H₂O] was precipitated from the solution (Reaction (3)) instead of magnesite (MgCO₃). When the temperature was increased (up to 125 °C), hydromagnesite was transformed to MgCO₃ albeit slowly (more than 15 h).

$$Mg_2SiO_4$$
 (sol) + $2CO_2$ (gas) + $2H_2O$ (liq.) $\rightarrow 2MgCO_3$ (sol) + H_4SiO_4 (aq.) (Rx. 2)

$$5 Mg_2 SiO_4~_{(sol)} ~+~ 8 CO_2~_{(gas)} ~+~ 20 H_2 O~_{(liq.)} \rightarrow 2 [4 Mg CO_3 \bullet Mg (OH)_2 \bullet 4 H_2 O]_{(sol)} ~+~ 5 H_4 SiO_4~_{(aq.)}~~(Rx.~3)$$

The work of Hänchen *et al.* [57] is intriguing, as we found the evidence of hydromagnesite formation in our samples, suggesting that this phase may be as a metastable "intermediate" phase [38] with respect to magnesite, but kinetically favored as a first reaction product under certain conditions (e.g., at low temperature conditions).

2.7. Geochemical Modeling

Equilibrium simulations were performed using PHREEQC (version 2.18—Parkhurst and Appello [58] with data from the Lawrence Livermore National Laboratory (LLNL, Livermore, CA, USA, supplied with the code). The first, very simple calculations examined the saturation state of a solution (initially pure water) when equilibrated with either magnesite, or both magnesite and carbon dioxide gas. Figure 9 shows the saturation index of both solutions with respect to magnesium phases which could be formed from the elements in the simulated system. In each case, the solutions suggest saturation with brucite, Mg(OH)₂, would be reached above 25 °C, so the simulations were repeated, but allowing brucite to precipitate when it would otherwise be oversaturated at ambient pressure and it is these conditions which are shown in Figure 9. It is interesting to note that the solution remains undersaturated with respect to hydromagnesite at all temperatures and the influence of CO₂ saturation is negligible. This reinforces the view that hydromagnesite is an artifact of rapid precipitation, *i.e.*, a process under kinetic control, in preference to magnesite (in line with experiments by Konigsberger *et al.* [59] and Power *et al.* [60]), which, under equilibrium conditions, is the dominant magnesium-bearing phase.

Secondly, attempts were made to simulate the reaction of CO_2 gas with forsterite in the presence of water, with and without varying amounts of HCl. These results also suggest considerable disequilibrium, in that magnesite would *not* be the dominant "sink" for magnesium in the presence of the silicate ions liberated from forsterite dissolution. Examination of the saturation state of solutions in which CO_2 was sequentially added to forsterite-water mixtures, always suggested a greater tendency for sepiolite precipitation, in preference to magnesite. Over the temperature interval 1 to $175\,^{\circ}C$, CO_2 gas was sequentially "reacted" (numerically) with an excess of forsterite in the presence of water and these simulations were repeated in the presence of HCl to represent any carry-over of the acid from

the previous processing step. Although we recognize that the presence of hydrochloric acid has little influence on the thermodynamic equilibrium position, it has a major role in promoting a rapid increase in the forsterite dissolution kinetics, so it was considered here. In all cases, thermodynamic equilibrium was only reached when sepiolite was allowed to precipitate, which effectively prevented magnesite formation. No evidence of other phases being important in these reactions was seen and the influence of residual HCl was negligible.

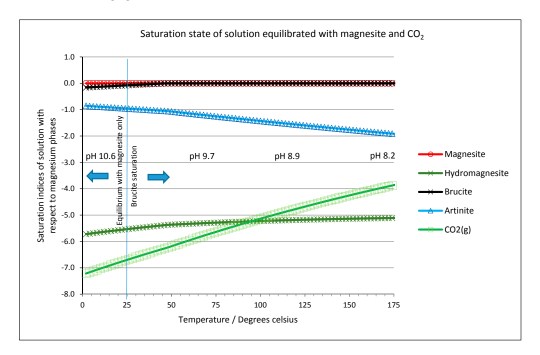


Figure 9. Saturation state of a solution equilibrated with magnesite and carbon dioxide gas $(P_{CO2} = 4.8 \text{ bar})$, allowing brucite to precipitate above 25 °C.

Experimentally, no evidence of sepiolite formation was seen and magnesite certainly dominates the solid phase assemblage after carbonation in this work and in the previous studies cited above. Our observations are that forsterite dissolves and magnesite and amorphous silica precipitate as reaction products (see Figures 2 and 3). Experimental studies on the stability of sepiolite have demonstrated that if the silica activity value is high in alkaline solutions, sepiolite precipitates [61–63]. From this, we must assume that the kinetics of sepiolite formation are very slow indeed and consequently it was excluded from the mineral assemblage in subsequent calculations. Figure 10 shows the solution chemistry predicted in a system where forsterite is carbonated by gaseous CO_2 in the presence of water. One mole of forsterite (in excess) was sequentially reacted with 0.2 mol of CO_2 gas, allowing equilibrium to be reached with both amorphous silica and all possible magnesium phases except sepiolite (which was prevented from forming in the simulations). At all temperatures between 25 °C and 175 °C carbonation is predicted to cause dissolution of forsterite and precipitation of magnesite and amorphous silica, with no other magnesium-bearing phases reaching equilibrium, except sepiolite which was allowed to remain oversaturated in the pore solution.

The solid phase chemistry is shown in Figure 11, which plots the ratio of forsterite dissolved against magnesite precipitated (mole ratio) against the number of moles of carbon dioxide consumed by the reaction. This is in effect the "yield" of the reaction, showing the moles of product formed as a function of the moles of reactant consumed. The theoretical stoichiometry (see Equation (1), above) is that two moles of forsterite react with one mole of CO₂ to produce one mole of magnesite and one mole of amorphous silica. These simulations suggest that this condition is reached quite rapidly. After reaction of only 0.9 mol of CO₂, the reaction is 97% complete in stoichiometric terms; that is to say the bulk of the CO₂ reacts to form magnesite and very little partitions into the liquid

phase (cf. Figure 10). Figure 11 shows that the temperature does have an effect on the extent of reaction and efficiency of carbonation in terms of the overall yield. The pseudo steady-state conditions established after reaction of 0.2 mol of CO₂ are maintained if further reaction is simulated, such that the solution chemistry remains constant, as does the ratio of forsterite and CO₂ consumed to the quantities of amorphous silica and magnesite precipitated. This is shown in Figure 12, where an excess of carbon dioxide eventually exhausts the reserve of forsterite, maintaining the stoichiometry of the reaction throughout. It must be stated that since the simplistic geochemical reaction path modeling is used in this study, many physical processes, such as material deformation, pore-fluid flow, heat transfer, advection, diffusion/dispersion, are ignored, although they should be considered in future work, as they are expected to be important during industrial-scale-up. In addition, because the chemical dissolution-front instability is also neglected in this study, many important factors, such as mineral reactive surface area, mineral dissolution ratio, solute dispersion, medium anisotropy, medium and fluid compressibility, have also been ignored. To consider these factors appropriately, more comprehensive chemical-transport modeling will be required.

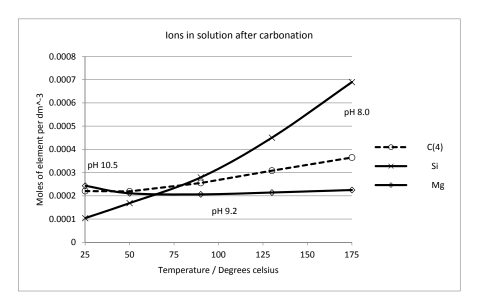


Figure 10. Solution chemistry predicted after reaction of excess forsterite (1 mol) with carbon dioxide gas (0.2 mol of CO_2) over the temperature interval (25 °C to 175 °C).

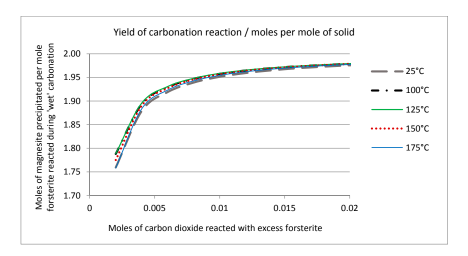


Figure 11. Solid phase chemistry predicted after reaction of excess forsterite (1 mol) with carbon dioxide gas over the temperature interval (25 °C to 175 °C) expressed as moles of magnesite precipitated/moles forsterite reacted, as a function of carbon dioxide consumed.

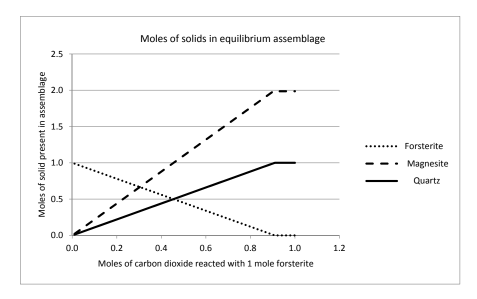


Figure 12. Solid phase chemistry predicted after reaction of one mole of forsterite with excess carbon dioxide gas at 25 °C, showing the stoichiometric relationship between moles of reactant (forsterite) consumed to moles of product (magnesite and quartz) precipitated.

3. Experimental and Modeling Section

3.1. Materials

Three kilograms of olivine mineral were excavated and collected from different depths of the Mont Tawai peridotite stratum in Malaysia. Although this source is local, the results can be considered as broadly representative of peridotite as the mineral composition is typical. The samples were crushed by a mechanical grinder and sieved into four different size ranges: <75 μ m, 75–125 μ m, 125–200 μ m and >200 μ m. Subsequently, they were dried to a constant weight at 120 °C for 2 h. Ten grams of each size fraction were mixed separately in 500 mL of 0.01 M hydrochloric acid (HCl, QRëC reagent grade, 37%) solution to remove impurities.

The olivine samples were characterized using scanning electron microscopy (SEM, S-4700 Hitachi, Tokyo, Japan) and X-ray diffraction (XRD, X'Pert powder, PANalytical, Almelo, The Netherlands), while elemental composition was measured by X-ray fluorescence (XRF, PW-1410, PANalytical, Almelo, The Netherlands). The carbonation yield of the olivine was determined using a total carbon analyzer (TCA, CS844, LECO Corp., St. Joseph, MI, USA).

3.2. Experimental Apparatus

The experimental equipment for CO_2 mineralization consisted of a CO_2 analyzer system mounted in a flow system connected to a cylindrical (500 mm \times 10 mm (d)) autoclave reactor with the means to supply diluted CO_2 and flushing with pure N_2 (Figure 13). CO_2 was introduced into the reactor at different partial pressures (up to 30%). Two flow meters (FM-1050, Matheson Tri-gas, Basking Ridge, NJ, USA) were used to control the flow rate of the inlet gases. The autoclave reactor was loaded with mineral and acid, then placed in a furnace where the temperature was measured and controlled using a thermocouple inserted directly into the reactor.

A water vapor generator was used to humidify the gas stream. CO₂ consumption was measured as the difference between supply and vent level at fixed flow (with integration over time) using an optical IR-sensor (GMP221, Vaisala Oyj, Helsinki, Finland), according to Equation (1):

$$CO_{2}\text{uptake }\left(\frac{mol}{g}\right) = \sum_{i}^{n} \frac{\left(pCO_{2\text{ in}} - pCO_{2\text{ out}}\right)_{i} \times \Delta t \times Q}{R \times T \times M}$$
(1)

where $pCO_{2 \text{ out}}$ and $pCO_{2 \text{ in}}$ are mean value of pCO_{2} (atm) at the inflow and outflow (equilibrium supplyP = 4.8 bar), Δt and Q are time interval (min) and flow rate (L/min), respectively, R and T are gas constant (0.082057 l.atm/mol.K) and temperature (K), respectively, and M is the mass of forsterite (g).

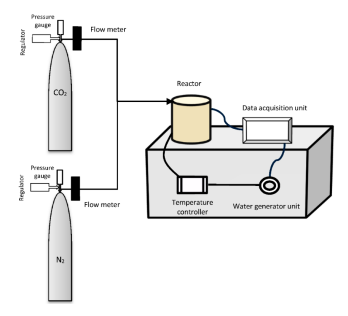


Figure 13. Schematic diagram of experimental set-up for CO₂ sequestration in chemically pretreated peridotite mineral from Mount Tawai, Malaysia.

3.3. Experimental Procedure

In this study, 1 M HCl was used for dissolution of Mg from the mineral matrix of peridotite and the dissolution experiment was conducted in a separate vessel. When a stoichiometric amount of HCl solution was added to olivine powder in a reaction vessel and stirred with a magnetic stirrer at 60 °C for two h, the entire mass formed a slurry. This slurry was then transferred into the autoclave reactor and neutralized by adding a base (NaOH: Fisher Chemicals reagent grade, with purity 99.999%) until the final pH was increased to above seven. Before heating the reactor, it was purged with nitrogen in order to replace the air inside the reactor, then the reactor was pre-heated to 175 °C for 1 h to dry the slurry and cooled to ambient. CO_2 gas (SFE grade, with a purity of 99.99% contained in a dip-tube cylinder and purchased from MOX Company, KL, Malaysia) was passed through the dried slurry at a typical flue gas level (15 vol %) or 4.8 bar pressure (P_{tot} = 32 bar) in the absence or presence of water vapor. The temperature of the carbonation process was studied over the interval from ambient to 175 °C.

The reactions involved in the extraction of Mg and carbonation are as follows:

$$4HCl_{(liquid)} + Mg_2SiO_{4 \text{ (solid)}} \rightarrow 2MgCl_{2 \text{ (aqueous)}} + SiO_{2 \text{ (solid)}} + 2H_2O_{(liquid)}$$
 (Rx. 4)

In this reaction scheme (Reaction (4)), forsterite dissolves in HCl, forming soluble $MgCl_2$ (Mg^{2+} remains in solution), and leaves behind insoluble SiO_2 . [15] Magnesium hydroxide [$Mg(OH)_2$] is precipitated by neutralization with NaOH (Reaction (5)). Then, by passing CO_2 , $Mg(OH)_2$ is converted to $MgCO_3$ in a gas-solid carbonation process (Reaction (6)).

$$MgCl_2 + 2NaOH \rightarrow Mg(OH)_2 + 2NaCl$$
 (Rx. 5)

$$Mg(OH)_2 + CO_2 \rightarrow MgCO_3 + H_2O$$
 (Rx. 6)

The effect of water vapor level on carbonation was studied in the range 5–20 vol % H₂O, corresponding to a range of relative humidity 18%–72% RH. After completion of the experiment,

the samples were collected and filtered using <75 μ m pore size Whatman filter papers. The MgCO₃ formed in the sample was quantified by titration against HCl. For the titration, a certain amount of solid (0.5 g) was weighed in a conical flask. Then, 20 mL of 1 M HCl was added into the flask and was allowed about two hours to react with the MgCO₃. The excess HCl was then back-titrated using 0.1 M NaOH solution. From the difference in titrant volume, the HCl consumed was calculated from which the content of MgCO₃ was deduced.

3.4. Estimation of Carbonation Yield

The extent of CO₂ mineral carbonation Y_{CO_2} was estimated using the TCA method, which is based on the mineralogy of samples tested and the capacity of carbon sequestrated $\left(\frac{1}{R_{CO_2}}\right)$ [64]. R_{CO_2} is considered as the weight fraction of CO₂ that can be trapped in a specific amount of mineral. According to Gadikota *et al.* [64] the capacity of CO₂ sequestration in forsterite (Mg₂SiO₄) can be expressed as follows:

$$\frac{W_{CO_2}}{W_{fo}} = \left(\frac{1}{R_{CO_2}}\right) = \frac{Y_{Mg}}{MW_{Mg}} \times MW_{CO_2} \tag{2}$$

where W_{fo} and W_{CO_2} are weights of forsterite before its mineral carbonation and CO_2 sequestrated in the solid phase (*i.e.*, magnesite), respectively. Y_{Mg} is the mass fraction of Mg^{2+} in the forsterite (*i.e.*, 34.55%) that can react with carbon dioxide to form stable magnesite. MW_{mg} and MW_{CO_2} are the formula weights of Mg^{2+} (48.61 mol/g in the forsterite) and CO_2 (44.01 mol/g), respectively, in the carbonated forsterite (2MgCO₃ & SiO₂—see (Reaction (1))). Therefore, Y_{CO_2} is the amount of carbon dioxide sequestrated (as magnesite) relative to the maximum capacity of CO_2 sequestration in forsterite ($\frac{1}{R_{CO_2}}$ = 31.27%).

$$Y_{CO_2} = R_{CO_2} \times \left[\frac{3.67 \times \text{weight fraction of carbon in MgCO}_3}{1 - \left(3.67 \times \text{weight fraction of carbon in MgCO}_3 \right)} \right] \times 100\%$$
 (3)

where 3.67 is the CO_2/C mass ratio. The carbonation yields of the forsterite and the effect on these of key empirical variables—reaction temperature, time, and particle size, are compared below.

3.5. Modeling System and Kinetic Analysis

Thermodynamic calculations were performed with the PHREEQC program software (version 2.18) with data from the LLNL [58]. Forsterite is the Mg-end member of the forsterite-fayalite solid solution series, and is included in the LLNL database. This program was used to estimate the dissolution and carbonation of forsterite samples in order to predict CO₂ uptake processes and potentials. Moreover, thermodynamic equilibrium constants for the mineral carbonation reactions of forsterite were provided by model databases. In doing so, reaction kinetics were implemented by using a BASIC interpreter. The possibility of implementing reactions kinetic into the code as BASIC statements was also used to predict the reaction progress over time. Consequently, the quality and validity of the model system and the determined rate and equilibrium parameters were verified against the results of carbonation experiments with forsterite samples. The data from a sequence of laboratory efforts were applied for that purpose, which were performed in the aqueous autoclave mini reactor.

The kinetic analysis of the forsterite dissolution rates was determined according to "standard integral analysis Levenspiel's method" [65] in Mg-rich solution using HCl. The results were set into several heterogeneous reaction models represented by integral rate equations and then the multiple regression coefficients (R) were calculated. The shrinking core model described by Dri et al. [66] was applied for the constant size of forsterite particles. Based on this method, reaction rates take place at the outer surface of the unreacted particles, and heterogeneous reactions are controlled by the product layer diffusion (Equation (4)), film diffusion (Equation (5)) and chemical reaction control (Equation (6)). In addition, the

possibility of having a compound effect of "chemical reaction control" and "product layer diffusion" was investigated using Equation (7).

$$kt = 1 - 3(1 - X_E)^{\frac{2}{3}} + 2(1 - X_E)$$
(4)

$$kt = 1 - 3(1 - X_E)^{\frac{1}{3}} \tag{5}$$

$$kt = X_E (6)$$

$$kt = \left[1 - 3(1 - X_E)^{\frac{2}{3}} + 2(1 - X_E)\right] + \left[1 - 3(1 - X_E)^{\frac{1}{3}}\right]$$
(7)

In these equations. "t" is time (s) and "k" (s⁻¹) and " X_E " denote the rate constant and the extent of reaction, respectively.

4. Conclusions

This work has experimentally and numerically modeled the process by which carbon dioxide gas may be sequestered, in situ by reaction with forsterite and/or its extracted intermediate brucite (ex situ) in the presence of moisture. In both cases, we have found that the reaction is favored resulting in a high carbonate yield; going almost to completion with the bulk of the carbon partitioning into magnesite and that very little remaining in solution. In the presence of water vapor, the degree of mineral carbonation was increased, we suggest, due to an alternative carbonation pathway providing for faster reaction kinetics [64]. Despite the observations made in other studies, we suggest that hydromagnesite is an intermediate in these carbonation experiments but is converted into magnesite on the (hours) timescale of reaction studied here, although the mechanism is as yet unclear. Hydromagnesite is less desirable as final product as it corresponds to only 80% sequestration (relative to magnesite) on a molar basis. Moreover, we recognize that this system is itself not at thermodynamic equilibrium, but maybe take a very long period and recrystallization as sepiolite (Mg₄Si₆O₁₅(OH)₂·6H₂O) is one possible outcome. However, we have found no evidence that this occurs over the period of one year. The clay mineral sepiolite is known to form in nature from dolomite/silica assemblages in the presence of water, but its formation kinetics are slow and we do not expect its formation to occur spontaneously in an industrial carbonation plant. Thus, we propose that the carbonation of readily available forsterite is a viable route for carbon sequestration.

From the computational perspective, a $\rm CO_2$ sequestration system with mineral carbonation can be treated as a fully-coupled problem between rock deformation, pore-fluid flow, heat transfer, mass transport, and chemical reaction processes. The results obtained from this study establish the viability of a geochemical model that can be used in to simulate the dynamic processes involved in $\rm CO_2$ sequestration in the Mount Tawai peridotite, Malaysia.

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