

Magnesite formation from MgO and CO₂ at the pressures and temperatures of Earth's mantle

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ABSTRACT

Magnesite (MgCO₃) is an important phase for the carbon cycle in and out of the Earth's mantle. Its comparably large *P-T* stability has been inferred for several years based on the absence of its decomposition in experiments. Here we report the first experimental evidence for synthesis of magnesite out of its oxide components (MgO and CO₂) at *P-T* conditions relevant to the Earth's mantle.

Magnesite formation was observed in situ using synchrotron X-ray diffraction, coupled with laser-heated diamond-anvil cells (DACs), at pressures and temperatures of Earth's mantle. Despite the existence of multiple high-pressure CO₂ polymorphs, the magnesite-forming reaction was observed to proceed at pressures ranging from 5 to 40 GPa and temperatures between 1400 and 1800 K. No other pressure-quenchable materials were observed to form via the MgO + CO₂ = MgCO₃ reaction. This work further strengthens the notion that magnesite may indeed be the primary host phase for oxidized carbon in the deep Earth.

Keywords: Deep carbon, magnesite, carbon dioxide, polymorphism

INTRODUCTION

The geological significance of carbon is clear due to its key roles in magmatic, metamorphic, eruptive, and ore-formation processes (Ganino and Arndt 2009; Heijlen et al. 2008; Roberge et al. 2009; Tappert et al. 2009). Furthermore, the transport, cycling, and storage of deep-Earth carbon are areas of intense, long-standing interest in which the interplay between atmospheric, crustal, and mantle reservoirs is well appreciated (Hayes and Waldbauer 2006; Hirschmann and Dasgupta 2009; Ridgwell and Zeebe 2005; Sleep and Zahnle 2001). Due to its large volume (and therefore potentially large total carbon content), convective nature and direct connection to Earth's surface via subduction and volcanic exchanges, the mantle is particularly important for understanding the carbon cycle on a planetary scale. The relative abundance of oxidized carbon vs. reduced carbon is difficult to quantify and is likely to vary with depth (e.g., Dasgupta and Hirschmann 2010), and there is abundant evidence regarding its complex mantle speciation, as indicated by the occurrence of carbonates in mantle-derived xenoliths (Berg 1986; Hervig and Smith 1981; McGetchin and Besancon 1973), kimberlitic diamonds, carbonatite magmas (Bell 1989) and prolific volcanic outgassing of CO₂ (Oppenheimer and Kyle 2008; Werner and Brantley 2003) and, occasionally, CH₄ (Fiebig et al. 2004).

There have been many petrologic studies in the CaO-MgO-SiO₂-CO₂-H₂O system (or a subset or slight expansion thereof) (Berg 1986; Brey et al. 1983; Brey and Green 1977; Katsura and

Ito 1990; Koziol and Newton 1998; Kushiro 1975; Martinez et al. 1998; Newton and Sharp 1975; Olafsson and Eggler 1983; Wyllie 1977). The majority of these studies have been carried out in large-volume presses at pressures and temperatures (*P-T*) representative of the lower crust and upper mantle, where much of our current knowledge about the carbon-containing system lies.

A key conclusion from these early studies is the remarkable stability of carbonate minerals prior to decarbonation reactions at high temperatures. Although carbonate inclusions in xenoliths are fairly rare, likely due to decarbonation during exhumation (Canil 1990), numerous subsequent experimental studies have documented the striking stability of magnesite (MgCO₃) to very high *P-T* (Berg 1986; Biellmann et al. 1993; Canil 1990; Fiquet et al. 2002; Fiquet and Reynard 1999; Isshiki et al. 2004; Katsura and Ito 1990; Katsura et al. 1991; Litasov et al. 2008; Martinez et al. 1998; Ross 1997; Santillan et al. 2005). The high temperatures [3000 K at 85 GPa (Isshiki et al. 2004)] achieved via laser-heated DACs strongly support the stability of magnesite to extreme *P-T*. Accordingly, carbonate is generally considered to be the dominant oxidized carbon species throughout much of the mantle, with magnesite, specifically, as the most abundant carbonate.

The predominance of magnesite at deep lower-mantle conditions, however, is less clear, as a structural transition to magnesite II at pressures greater than ~115 GPa has been observed (Isshiki et al. 2004; Panero and Kabbes 2008; Skorodumova et al. 2005). Additionally, Boulard et al. (2011) demonstrated the formation of a (CO₄)⁴⁻-structured phase in iron-containing magnesite [(Mg,Fe)

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CO₂] at pressures greater than 60 GPa, which is similar to a computationally predicted phase (Oganov et al. 2008), yet this new phase is expected to coexist with magnesite and diamond to great depths, especially if transported within a relatively cold subducting slab (Boulevard et al. 2011). And, although pure magnesite has not been observed to dissociate into separate oxides such as MgO and CO₂ at the *P-T* of Earth's mantle, decarbonation reactions between magnesite and SiO₂, forming an assemblage of MgSiO₃-perovskite and CO₂, have been documented at lower-mantle *P-T* (Seto et al. 2008; Takafuji et al. 2006).

Furthermore, whereas some of the earlier large-volume press studies observed the formation of magnesite from CO₂ (typically introduced as Ag₂C₂O₄, silver oxalate), subsequent diamond-anvil studies on the high-pressure (>~10 GPa) stability and high-temperature decarbonation of magnesite have generally relied on magnesite as a starting material as opposed to observing its formation from oxides. An exception to this is the work of Boulevard et al. (2011) who reacted both pure and iron-bearing MgO with CO₂ at pressures greater than 80 GPa in their work demonstrating the formation of a high-pressure carbon-bearing phase based on CO₄⁺ units.

Notably, the phase diagram of CO₂ is complicated, and the high-pressure polymorphs are relatively dense and incompressible compared to "dry ice" (CO₂-I), which forms from the low-temperature deposition of gaseous CO₂ at ambient pressure or by the compression of liquid CO₂ to roughly 0.5 GPa at ambient temperature. Figure 1 shows an approximated phase diagram for CO₂ as it is currently known for the *P-T* relevant to this study, following the recent experimental results and compilation of earlier work as summarized by Litasov et al. (2011); the figure is supported by data and interpretations from Tschauner et al. (2001), Santoro et al. (2004), Giordano et al. (2006), Giordano and Datchi (2007), Iota et al. (2007), and Datchi et al. (2009). It is important to note, however, that kinetic effects can be quite

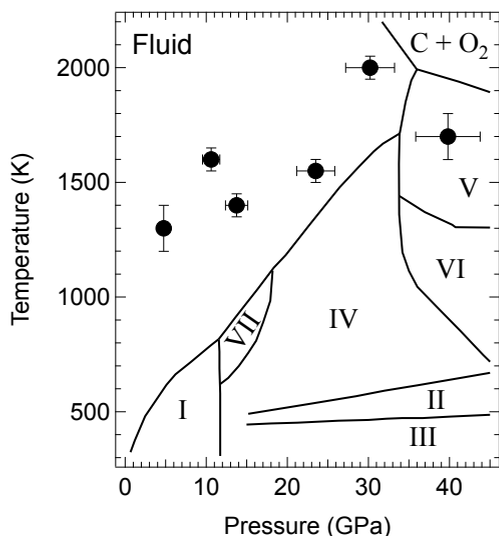


FIGURE 1. A simplified phase diagram of CO₂, after the summarizing figure of recent work presented by Litasov et al. (2011). The superimposed data points are the *P-T* conditions encountered during the laser-heating measurements of this study.

significant, and these are not necessarily equilibrium boundaries; Figure 1 is intended simply to summarize the CO₂ phases that one may expect in a study such as this, and some boundary locations, such as for the breakdown reaction from CO₂ to C + O₂, have been placed in different locations by previous workers. The reader is encouraged to refer to Litasov et al. (2011) or Sengupta et al. (2011) for a more thorough illustration of where these boundaries have been placed by different workers, or to Yoo et al. (2011a, 2011b) for extensions to even higher-pressure conditions.

It is conceivable that such high-pressure CO₂ polymorphism could hinder the formation of magnesite even at moderate pressures. The motivation of the present study was to investigate magnesite formation in situ from CO₂ and MgO well within its range of expected stability. It is important to recognize a key caveat regarding these experiments, however, which is that they were all performed under CO₂-saturated conditions, and it is certainly not realistic to portray the mantle as having an activity of CO₂ close to unity (*a*_{CO₂} = 1). That said, direct observations of the forward reaction, rather than the lack of a breakdown reaction, would strengthen the case for magnesite as a primary storage mineral for oxidized carbon in much of Earth's mantle.

EXPERIMENTAL METHODS

Symmetric-type diamond-anvil cells (DACs) were used to contain and compress samples for all experiments, and infrared lasers were used for heating. The DACs utilized type I anvils with either 350 or 500 μm diameter culets, depending on the desired experimental pressure, and 250 μm thick stainless steel (Type 301) was used as a gasket material. The gaskets were pre-indented to a thickness of between 60 and 80 μm, and sample chambers were drilled with diameters between 100 and 200 μm. For starting material, high-purity (>99.9%) Pt-black was added to pure MgO at the 3 wt% level to absorb infrared laser radiation, and this mixture was finely ground and pressed between two opposing anvils to form platelets approximately 20 μm thick.

A few grains of pure MgO were placed into the bottom of the sample chamber to hold the platelet above the anvil surface to avoid direct thermal contact with the anvil during laser heating. A piece of an MgO-Pt platelet, smaller than the diameter of the sample chamber, was then placed on the pure MgO grains in the center of the sample chamber. One or two grains of ruby of tens of micrometers in diameter were placed directly on the upper culet so that ruby fluorescence could be used to determine pressure during compression (Mao et al. 1978). The pressures reported in this paper, however, were determined by using X-ray diffraction and the equation of state (EoS) for MgO (Jacobsen et al. 2008); a 5% relative uncertainty in pressure was estimated for these measurements. Figure 2 shows a schematic illustration of the sample geometry and a photograph of an actual sample.

After loading and positioning the solid materials in the sample chamber, the entire cell was surrounded by liquid CO₂ using either the gas-loading system at the Carnegie Institution of Washington's Geophysical Laboratory in Washington, D.C., or the gas-loading system at the GeoSoilEnviro Consortium for Advanced Radiation Sources (GSECARS) at Argonne National Laboratory. This was accomplished at room temperature by placing a slightly open DAC into a vessel that was subsequently pressurized to ~0.06 GPa (~8000 p.s.i.) with pure CO₂. The DACs were then closed by remotely turning their screws to trap the CO₂. Raman spectroscopy was used subsequently to confirm the presence of CO₂ before further compression and heating.

X-ray diffraction measurements were conducted at the Advanced Photon Source (APS) of the Argonne National Laboratory at three different beamlines: 16-ID-B and 16-BM-D of the High-Pressure Collaborative Access Team (HPCAT) and 13-ID-D of GSECARS. The double-sided laser heating systems at 16-ID-B and 13-ID-D, both of which utilize Nd:YLF lasers with a wavelength of 1053 nm, were used to overcome potential kinetic barriers for the formation of equilibrium assemblages. The temperatures reported here are not intended to represent the location of equilibrium phase boundaries in *P-T* space and, in most cases, are simply the lowest temperature at which sustained and consistent heating could be pyrometrically measured.

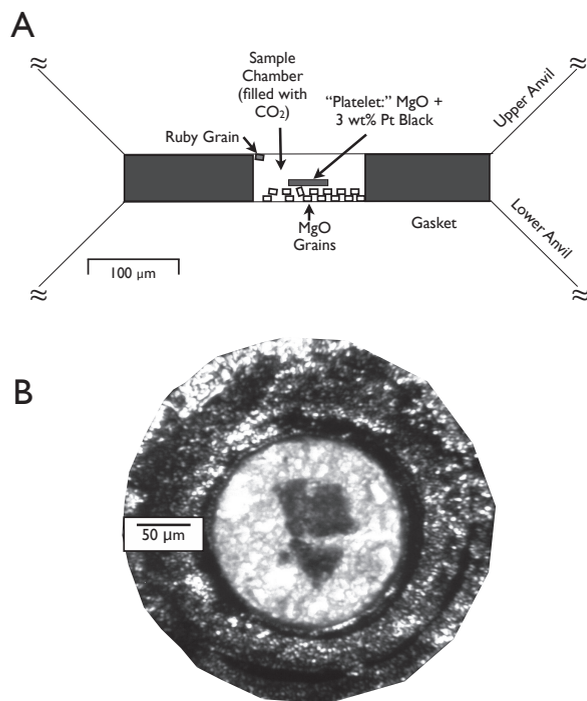


FIGURE 2. (a) Schematic for loaded sample chamber: Pt black was used to absorb infrared laser radiation and the pure MgO grains were used as a substrate to thermally isolate the MgO-Pt platelet from the anvils during heating. (b) A photograph of a loaded sample as viewed from above. This particular loading had two spatially separate platelets, one that is roughly square and the other roughly triangular; this readily allowed for two heating experiments to be conducted at different pressures.

Analyses of diffraction patterns, after they were converted to one-dimensional profiles using FIT2D (Hammersley et al. 1996), were conducted in two stages. First, peak positions and intensities for expected phases were approximated using previously published diffraction information: unit-cell data (i.e., crystal system and lattice parameters), Miller indices (hkl values), and observed relative intensities. For phases with well-known ambient-pressure structures and equations of state (K_{0T} and dK/dP), such as Pt, MgO, CO₂-I, and magnesite, the high-pressure lattice parameters, and therefore d -spacings for the expected hkl values, could be readily calculated. Because the equations of state for the high-pressure CO₂ polymorphs are less well constrained, estimates were made to calculate the lattice parameters at the pressure being considered. Unambiguous individual diffraction peaks attributed to each phase were then fit by Gaussian profiles to determine their d -spacings, and linear least-squares regression, coupled with their associated hkl values, was utilized to estimate the lattice parameters for each phase.

Subsequently, these estimated lattice parameters were used as starting values for whole-profile fitting with GSAS (Larson and Von Dreele 2000) utilizing the EXPGUI interface (Toby 2001). Due to the presence of multiple phases (up to five in some patterns) and some overlapping peaks, Rietveld structure refinement was found to be not feasible, but Le Bail whole-profile fitting was used to refine the lattice parameters for all phases satisfactorily.

RESULTS AND DISCUSSION

Overview of experiments performed

Six laser-heating experiments were conducted at pre-heating pressures of 4.8(2), 10.9(5), 14.4(7), 22.3(11), 30.3(15), and 41.7(20) GPa, with a measured pressure drop of less than 10% after heating for most experiments. Although multiple diffraction patterns were typically collected before, during, and after heating, these results will be discussed and presented as pre- and

post-heating diffraction pairs and grouped based on the CO₂ polymorphs observed prior to heating.

CO₂ initially as CO₂-I

The experiments conducted at 4.8 and 10.9 GPa are relatively straightforward because they are in the CO₂-I stability field at room temperature, in which CO₂ crystallizes in the cubic system [space group $Pa\bar{3}$ (Downs and Somayazulu 1998)] with a well-constrained EoS (Giordano et al. 2010; Liu 1984; Olinger 1982). All pre-heating diffraction peaks at both pressures can be readily indexed as CO₂-I, MgO, or Pt.

Measurable heating for the experiment at 4.8 GPa started at 1300 K, but a temperature measurement could only be made from one side of the DAC, and this heating was sustained for 40 min. Slightly higher laser power was used for the experiment at 10.9 GPa to facilitate pyrometric measurements from both sides of the DAC, and temperatures between 1550 and 1650 K were recorded and sustained for 45 min. In both experiments new diffraction peaks were observed immediately upon heating, and the extended heating was conducted to see if it had a noticeable effect, but it appears to have been unnecessary.

The formation of magnesite after heating at 4.8 GPa (and a subsequent pressure drop to 4.7 GPa) can be unambiguously confirmed based on the new diffraction peaks observed, and there is no evidence for the formation of any other phases. The experiment at 10.9 GPa (10.3 GPa after heating) produced largely similar results in terms of clear magnesite formation, but only the strongest peak from CO₂-I remained readily apparent, and several new, but low-intensity, peaks were present. Based on the pressure and heating history, either orthorhombic CO₂-III [space group $Cmca$ (Aoki et al. 1994)] or rhombohedral CO₂-IV [space group $R\bar{3}c$ (Datchi et al. 2009)] would be plausible candidate phases (e.g., Datchi et al. 2009; Park et al. 2003), but CO₂-IV is a much better match with the observed data. Table 1 presents a summary of unit-cell data for all phases observed in these experiments.

Decompression data were also collected for CO₂-I from the cell that was heated at 10.9 GPa, away from the heated region, to verify the EoS for CO₂-I. This was done due to an apparent discrepancy in the literature, with the earlier studies of Olinger (1982) and Liu (1984) producing similar EoS values [$K_{0T} = 2.2$ GPa, $dK/dP = 6.4$, and $V_0 = 197.9$ (Å³ per unit cell)] [our fit to the pressure-volume (P - V) data in Olinger (1982)] and

TABLE 1. CO₂ Initially as CO₂-I

P (GPa)	Phase	a (Å)	c (Å)	V (Å ³)
Pre: 4.8	MgO	4.1730(2)		72.67(1)
	Pt	3.904(3)		59.5(1)
	CO ₂ -I	5.1826(1)		139.20(1)
Post: 4.7	MgO	4.1734(2)		72.69(1)
	Pt	3.8989(3)		59.27(2)
	CO ₂ -I	5.2366(10)		143.59(8)
Pre: 10.9	Magnesite	4.6081(5)	14.717(5)	270.64(6)
	MgO	4.1287(1)		70.378(7)
	Pt	3.854(2)		57.24(9)
	CO ₂ -I	4.9426(9)		120.76(7)
Post: 10.3	MgO	4.1331(1)		70.602(4)
	Pt	3.8695(2)		57.94(1)
	CO ₂ -I	4.960(3)		122.0(3)
	CO ₂ -IV	8.7934(8)	10.603(4)	710.0(3)
	Magnesite	4.5403(3)	14.432(1)	257.66(2)

$K_{OT} = 2.93$ GPa, $dK/dP = 7.8$, and $V_0 = 208.6$ (\AA^3 per unit cell), respectively} compared to a more recent value of $K_{OT} = 12$ GPa reported by Yoo et al. (1999). Notably, Giordano et al. (2010) presented high-quality P - V - T EoS data with values quite similar to the earlier studies: $K_{OT} = 3$ GPa, $dK/dP = 8.4$, and $V_0 = 200$ (\AA^3 per unit cell). The results determined here are also similar: $K_{OT} = 3.5$ GPa, $dK/dP = 9.1$ GPa, and $V_0 = 197.9$ (\AA^3 per unit cell) [fixed based on the estimate of Olinger (1982)] and, as pointed out by Giordano et al. (2010), the variations in the studies with K_{OT} values near 3 GPa can be explained by the small differences in V_0 and covariance between K_{OT} and dK/dP . The much higher K_{OT} value reported by Yoo et al. (1999), however, is hard to interpret because it was not reported with values for V_0 or dK/dP .

CO₂ initially as CO₂-III

As expected, carbon dioxide was present entirely as CO₂-III [space group *Cmca* (Aoki et al. 1994)] in the pre-heating patterns of experiments conducted at 14.4 and 22.0 GPa. Temperatures between 1350 and 1450 K were recorded during a heating duration of 20 min for the cell at 14.4 GPa, and the cell at 22.0 GPa heated stably at 1550 K, which was sustained for 15 min. A summary of unit-cell data for all phases encountered both before and after heating can be seen in Table 2.

Post-heating, the cell initially at 14.4 GPa dropped to 13.1 GPa, whereas the cell initially at 22.0 GPa increased in pressure to 25.0 GPa—the only heating experiment observed to result in an increase in pressure after heating. As with the experiment at 10.9 GPa, temperature-quenched patterns from both of these experiments included CO₂-IV, rather than CO₂-III, but tetragonal CO₂-II [space group *P4₂/mnm* (Yoo et al. 2002)] is present in the post-heat pattern at 25 GPa as well.

The formation of magnesite was clear and unambiguous in both experiments, but the experiment at 14.4 GPa produced a few unaccounted for peaks at 1.669, 1.587, and 1.423 \AA . These peaks are low in intensity, but they are sharp and spotty, and clearly form distinct diffraction rings on the image plate. Notably, quenching from the liquid phase of CO₂ at this pressure would be consistent with the formation of orthorhombic CO₂-VII [space group *Cmca* (Giordano and Datchi 2007)], but with so much of the pattern obscured by stronger reflections from other phases, it is not possible to make a conclusive determination.

TABLE 2. CO₂ Initially as CO₂-III

P (GPa)	Phase	a (\AA)	b (\AA)	c (\AA)	V (\AA^3)
Pre: 14.4	MgO	4.1060(1)			69.220(5)
	Pt	3.884(1)			58.58(6)
	CO ₂ -III	4.393(2)	4.562(4)	5.838(3)	117.0(1)
Post: 13.1	MgO	4.1143(1)			69.649(5)
	Pt	3.8644(1)			57.710(5)
	CO ₂ -IV	8.699(1)		10.641(3)	697.3(2)
	Magnesite	4.5249(5)		14.304(1)	253.63(4)
Pre: 22.0	MgO	4.0609(2)			66.97(1)
	Pt	3.840(2)			56.63(1)
	CO ₂ -III	4.22(1)	4.55(1)	5.791(9)	111.3(1)
Post: 25.0	MgO	4.0446(2)			66.16(1)
	Pt	3.8493(3)			57.04(1)
	CO ₂ -II	3.546(1)		4.2405(8)	53.31(2)
	CO ₂ -IV	8.475(1)		10.479(4)	651.9(2)
	Magnesite	4.4876(4)		14.005(1)	244.25(3)

CO₂ initially as CO₂-IV

Prior to heating, only one diffraction line is visually attributable to CO₂ for either of the experiments conducted at 30.3 and 41.7 GPa, and in both cases it appears to be due to overlapping (024) and (220) reflections from the CO₂-IV structure. Given that these are pre-heating patterns one may expect the CO₂-III structure, but it should be noted that spatially separated locations in both DACs, from the same loading, had already been heated at lower pressures. Neither of the pre-heating patterns shows evidence for the presence of magnesite, yet it is likely that heating elsewhere in the cells (approximately 50 to 100 μm away) warmed the cells sufficiently to convert all of the CO₂ to the CO₂-IV structure.

During heating, however, more CO₂ diffraction lines became apparent from both cells, and some of the CO₂ transformed to the tetragonal CO₂-V structure [space group *I42d* (Datchi et al. 2012)], whereas some remained as CO₂-IV. The cell at 30.3 GPa was heated for approximately 60 min, with momentary peak temperature measurements of >2700 K for both sides, but sustainable measurements were in the range of 1900 to 2100 K. The post-heating pressure was 30.1 GPa. For the cell at 41.7 GPa, sustained temperature measurements were between 1600 and 1800 K with a similar duration of 60 min; upon temperature quench, the pressure was 37.8 GPa. Notably, this was the only heating experiment for which all of the CO₂ should have remained within the solid state, as can be seen in Figure 1.

As with the lower-pressure experiments, the heating experiments at 30.3 and 41.7 GPa both produced clear and unambiguous diffraction patterns indicative of magnesite, and a summary of all phases observed pre and post heating can be seen in Table 3. Figure 3 shows pre- and post-heating patterns for the 41.7 GPa heating experiment. This experiment was chosen to display representative patterns because, as the highest-pressure experiment conducted in this study, it is arguably the most significant in terms of demonstrating the formation of magnesite at mantle pressures.

The upper panels in Figure 3 show the results of fitting from within EXPGUI/GSAS, and the residuals are shown at the bottom. The lower panels show the same patterns, but they have been converted to d -spacing, the backgrounds have been subtracted, and the dominant MgO peaks have been truncated to better show peaks from the other phases. Furthermore, the vertical sticks on the lower panel have been scaled so that the most intense peak

TABLE 3. CO₂ Initially as CO₂-IV

P (GPa)	Phase	a (\AA)	c (\AA)	V (\AA^3)
Pre: 30.3	MgO	4.0170(1)		64.820(4)
	Pt	3.831(1)		56.22(6)
	CO ₂ -IV	8.410(3)	10.197(9)	624.5(7)
Post: 30.1	MgO	4.0179(1)		64.863(7)
	Pt	3.8164(6)		55.58(3)
	CO ₂ -IV	8.417(2)	10.210(3)	626.5(2)
	CO ₂ -V	3.608(1)	5.917(3)	77.03(2)
Pre: 41.7	Magnesite	4.4318(5)	13.650(2)	232.18(3)
	MgO	3.9643(1)		62.304(5)
	Pt	3.801(2)		54.93(6)
Post: 37.8	CO ₂ -IV	8.177(3)	10.193(9)	590.3(4)
	MgO	3.9817(2)		63.126(9)
	Pt	3.791(2)		54.49(8)
	CO ₂ -IV	8.130(2)	10.174(2)	582.4(2)
Post: 37.8	CO ₂ -V	3.568(1)	5.913(4)	75.28(3)
	Magnesite	4.4030(9)	13.463(5)	226.08(8)

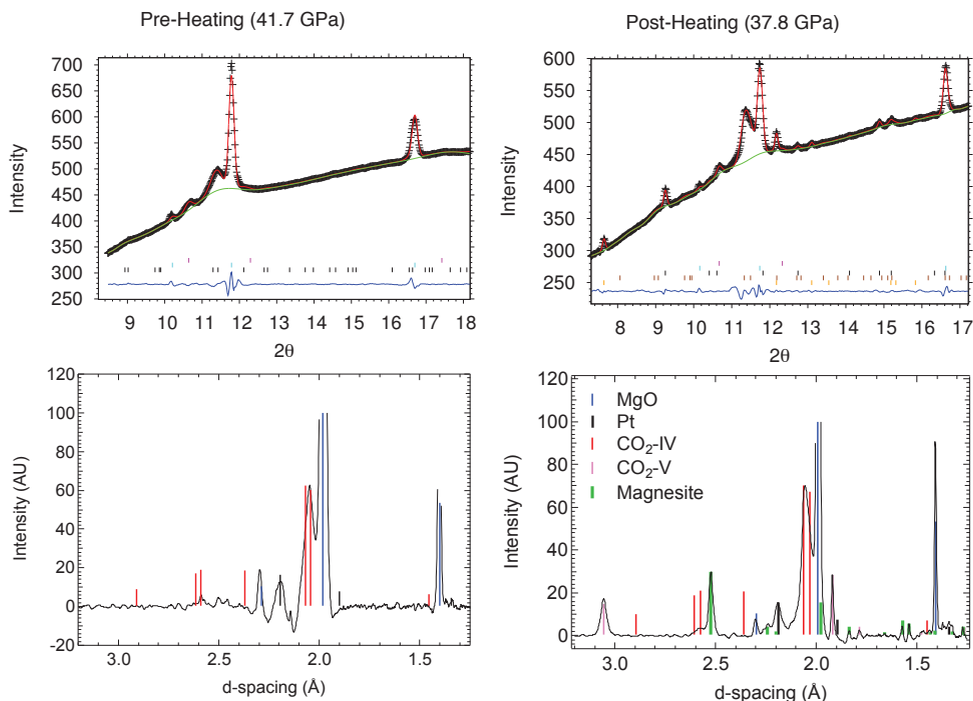


FIGURE 3. Diffraction patterns collected before and after laser heating at 41.7 GPa; a detailed description is provided in the text. (Color online.)

for each phase matches the greatest intensity observed for that phase. The remaining relative scaled intensities, however, are from the previously determined structures of other workers, and they are intended to provide an additional indication of the overall appropriateness of the phase identifications.

Although the integrated relative intensities for magnesite peaks show good agreement with those of previous workers, this is somewhat fortuitous because the data from the actual image plate are spotty as shown in Figure 4, which would be expected for crystal growth during laser heating. Although the most intense reflection from (104) can be identified as an individual ring, other reflections such as (202) manifest primarily as a few discrete spots with only very faint rings visible, if at all.

Both of these cells were subsequently decompressed, and all peaks in the ambient-pressure diffraction patterns can be readily indexed as magnesite, MgO or Pt. No peaks ascribed to CO₂ at high pressures persist to ambient pressure, and there are no unexpected peaks.

Anisotropic compression in magnesite

Because all of the magnesite observed in these experiments crystallized under simultaneous high P - T conditions, with the surrounding CO₂ above or very near its melting point, we would expect the deviatoric stress on the crystals to be fairly low. This is superficially supported by their narrow diffraction lines at all pressures. Our measured lattice-parameter axial ratios (i.e., c/a), along with those from previous workers (Fiquet et al. 2002; Fiquet and Reynard 1999; Ross 1997), are plotted against pressure in Figure 5 to compare our results with previous compressional data collected in both hydrostatic and non-hydrostatic conditions.

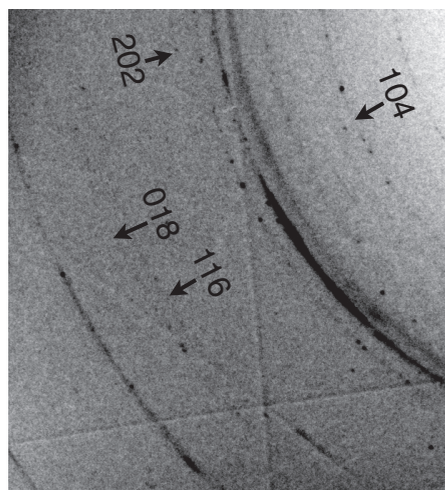


FIGURE 4. A portion of the detector image, for the post-heating X-ray measurement shown in Figure 3, prior to integration and conversion to a one-dimensional diffraction pattern. The labels are the hkl values for the magnesite reflections observed in the integrated patterns.

Fiquet et al. (1999) used various pressure-transmitting media to surround the magnesite; their data collected in argon and methanol-ethanol water (MEW) are essentially indistinguishable from the single-crystal data collected in methanol-ethanol (ME) by Ross (1997) over the pressure range for which they overlap, which is limited to pressures below about 9 GPa. Notably, this pressure is approximately where both ME and MEW solidify and, accordingly, non-hydrostaticity is expected. Fiquet et al. (2002), conversely, did not use any pressure-transmitting medium and,

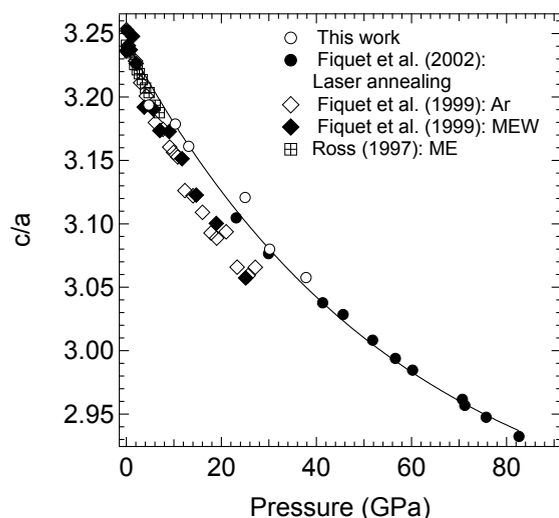


FIGURE 5. Ratio of lattice parameters, a and c , for the magnesite formed in this study compared to what has been observed by previous workers. The dashed vertical line roughly indicates the pressure below which the organic solvent pressure-transmitting media of previous workers would be in a liquid state and thus truly hydrostatic.

instead, used laser heating to thoroughly anneal the sample at each pressure step. An exponential trend can readily fit both their high-pressure data and the truly hydrostatic data points of Ross (1997) and Fiquet et al. (1999). The non-hydrostatic data points from Fiquet et al. (1999), however, clearly deviate to lower c/a values, strongly suggesting that deviatoric stress enhances the known anisotropic compression of magnesite. Our data show good agreement with this trend, but our datum at 25 GPa is at a higher c/a value than it would predict. Overall, however, it appears that the laser-annealing technique employed by Fiquet et al. (2002) does indeed relieve the deviatoric stress as intended.

Thermodynamics of magnesite formation

To accurately model the thermodynamic stability of magnesite relative to CO_2 and MgO at elevated P - T , one would need not only reference-state free energies, but also P - T dependent entropy and EoS data for all phases. Fiquet et al. (2002) created such a model using a combination of theoretically calculated and experimentally measured values to predict the equilibrium curve to pressures of 150 GPa and temperatures of 5000 K. They noted the importance of volumetric effects, particularly at high pressure, and they concluded that temperatures well above the geotherm would be necessary for the decarbonation of magnesite.

As a simple demonstration of this volumetric importance using the data presented in this study, Figure 6 shows the ΔV , in terms of molar volumes, of the reaction $\text{CO}_2 + \text{MgO} = \text{MgCO}_3$ upon temperature quench. The negative ΔV for calculations based on observations of quenched CO_2 -I and CO_2 -IV would indeed serve to drive the reaction toward magnesite production. However, the positive ΔV for calculations based on observations of quenched CO_2 -V would serve to inhibit the reaction. This does not, of course, suggest that the reaction should not proceed at high temperatures, but rather that an entropic term, or thermal

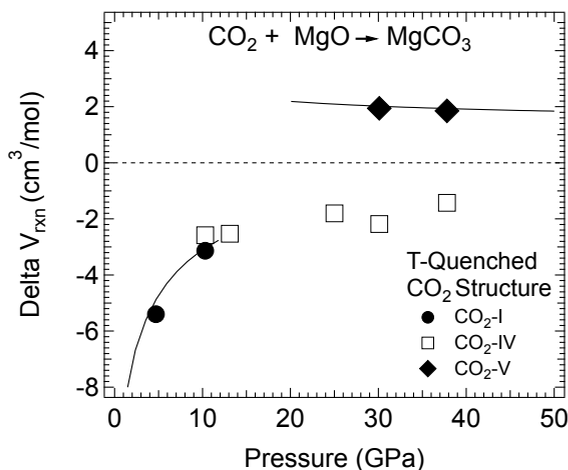


FIGURE 6. Change in molar volume (ΔV) for the magnesite-forming reaction upon temperature quench. The molar volumes were determined from the unit-cell volumes observed in this study using $Z = 4$ for CO_2 -I (Downs and Somayazulu 1998), CO_2 -V (Datchi et al. 2012), and MgO ; $Z = 6$ for magnesite; and $Z = 24$ for CO_2 -IV (Datchi et al. 2009). The solid curves were determined using EoS data for magnesite (Fiquet et al. 2002), MgO (Jacobsen et al. 2008), CO_2 -I (Giordano et al. 2010), and CO_2 -V (Datchi et al. 2012).

expansion, dominates in this case. That said, this is a significant observation because it does suggest that relatively dense high-pressure polymorphs in CO_2 , such as CO_2 -V, could serve to disfavor the magnesite-forming reaction at low temperatures.

CONCLUDING REMARKS

As expected based on a large body of previous work demonstrating its remarkable stability at high P - T , magnesite readily forms from CO_2 and MgO at pressures between 5 and 40 GPa and temperatures from 1400 to 1800 K. Indeed, even at the highest pressure of this study, and with CO_2 in the relatively dense CO_2 -IV structure, the reaction to form magnesite proceeded at temperatures below 1800 K.

The true significance of magnesite for carbon storage in the lower mantle remains a complicated issue that will, of course, depend on several critical parameters such as total carbon budget, relative partitioning between carbonates and silicates (Panero and Kabbes 2008) and oxygen fugacity (Frost and McCammon 2008). Indeed, the redox state at which carbonate would be reduced to diamond has been determined, based on buffered multi-anvil experiments, to be well above the iron-wüstite buffer (Stagno et al. 2011). Furthermore, recent work demonstrates that f_{O_2} decreases with increasing depth, at least in the upper mantle and, accordingly, carbon is likely present in reduced phases and oxidation does not occur until there is upwelling (Stagno et al. 2013).

Additionally, observations of high-pressure polymorphism (Isshiki et al. 2004) and newly discovered carbon-bearing phases (Boulard et al. 2011; Oganov et al. 2008) obscure the significance of the ambient-pressure magnesite structure at the greatest mantle depths. That said, the remarkable ease with which magnesite forms from MgO and CO_2 , as shown in this study, further supports the notion that it is indeed one of the dominant storage phases for oxidized carbon in much of Earth's interior.

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