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Carbonate stability in the reduced lower mantle

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ABSTRACT

Carbonate minerals are important hosts of carbon in the crust and mantle with a key role in the transport and storage of carbon in Earth's deep interior over the history of the planet. Whether subducted carbonates efficiently melt and break down due to interactions with reduced phases or are preserved to great depths and ultimately reach the core-mantle boundary remains controversial. In this study, experiments in the laser-heated diamond anvil cell (LHDAC) on layered samples of dolomite (Mg,Ca)CO₃ and iron at pressure and temperature conditions reaching those of the deep lower mantle show that carbon-iron redox interactions destabilize the MgCO₃ component, producing a mixture of diamond, Fe₇C₃, and (Mg,Fe)O. However, CaCO₃ is preserved, supporting its relative stability in carbonate-rich lithologies under reducing lower mantle conditions. These results constrain the thermodynamic stability of redox-driven breakdown of carbonates and demonstrate progress towards multiphase mantle petrology in the LHDAC at conditions of the lowermost mantle.

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1. Introduction

Carbonates are the major minerals responsible for transportation of carbon from the Earth's surface to its deep interior. Their behavior at depth is critical to the storage capacity and fluxes of the geologic carbon cycle. Recent estimates of the flux of carbon trapped in carbonate minerals that reach the deep Earth range from 0.0001 to 52 megatons annually (Dasgupta and Hirschmann, 2010; Kelemen and Manning, 2015). The uncertainty in this range hinges on poor constraints on the budget of carbon retained by subducting slabs. Transport of carbon to at least transition zone depths is demonstrated by carbonate inclusions in diamonds (e.g. Brenker et al., 2007; Wang et al., 1996), but whether any carbonates remain in the lower mantle is controversial. Most subducted carbon is expected to melt and/or break down and return to Earth's surface via volcanism (Dasgupta and Hirschmann, 2010; Kelemen and Manning, 2015; Thomson et al., 2016). However, relatively oxidizing conditions, low temperatures, and resulting slow kinetics within subducting slabs may result in transportation of

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carbonates magnesite, siderite, calcite, and their solid solutions to great depths (Martirosyan et al., 2016). Subducted carbonates in slabs that reach the base of the lower mantle will undergo multiple phase transitions and encounter reducing conditions, to be finally buffered by metallic iron at or near the CMB.

Carbon provides key constraints on the chemical evolution of the deep Earth through its role as a proxy for mantle redox conditions, as its speciation is largely governed by oxygen fugacity, or fO_2 (Frost and McCammon, 2008). Evidence of deep mantle chemistry (Walter et al., 2011) and redox state (Ryabchikov and Kaminsky, 2013; Smith et al., 2016) in local environments of diamond formation can likely be inferred from diamond inclusions. Diamonds and their inclusions indicate widely varying local fO_2 (Brenker et al., 2007; Walter et al., 2011; Ryabchikov and Kaminsky, 2013; Smith et al., 2016; Kaminsky, 2012), though fO_2 is expected to generally decrease with mantle depth (Frost and Mc-Cammon, 2008; Rohrbach and Schmidt, 2011). Both isotopic evidence (e.g. Harte, 2010; Tappert et al., 2005) and carbonate inclusions in diamonds (e.g. Wang et al., 1996) suggest that diamonds are (at least in part) formed by reduction of subducted carbonate minerals or carbonate melts. Reduction of carbonates to diamond takes place by interactions with mantle silicates (Pal'yanov et al., 2002; Stagno et al., 2011, 2013, 2015) or metal (Arima et al., 2002; Siebert et al., 2005; Rouquette et al., 2008; Pal'yanov et al., 2013).



These redox reactions contribute to the barrier to carbonate transport to the deep Earth (Rohrbach and Schmidt, 2011; Thomson et al., 2016). The key to determining the amount of oxidized carbon in the deep Earth is the dependence of both the thermodynamics and the kinetics of redox reactions in carbonates on temperature, pressure, and fO_2 .

Petrologic observations, experiments and computational studies on stability and breakdown kinetics of carbonates have determined that magnesite is the most stable carbonate phase throughout most of the mantle (e.g. Dasgupta and Hirschmann, 2010; Rohrbach and Schmidt, 2011), but calcite and siderite polymorphs have also attracted interest. Relative to aragonite (Spivak et al., 2011) or siderite (Tao et al., 2013), magnesite is more likely to subduct to the deep mantle due to its higher melting temperature (Isshiki et al., 2004; Katsura and Ito, 1990; Martinez et al., 1998; Solopova et al., 2014). Melting experiments on carbonated peridotite (Dasgupta and Hirschmann, 2010; Ghosh et al., 2014; Rohrbach and Schmidt, 2011) and eclogite (Thomson et al., 2016) systems confirm that calcium and iron carbonate components are more likely to enter a melt phase than magnesium carbonate in silicate lithologies. At sub-solidus temperatures, MgCO3 is also favored in carbonate-silicate mixtures by the reaction CaCO₃ + $MgSiO_3 \rightarrow MgCO_3 + CaSiO_3$ at lower-mantle pressures up to 80 GPa (Biellmann et al., 1993; Seto et al., 2008). Due to its broad solid solution with siderite (Lin et al., 2012; Liu et al., 2015), magnesite in the mantle will host iron. An (Mg, Fe)-carbonate in the lowermost mantle may have mixed redox state and coexist with diamond (Boulard et al., 2012). However, both (Mg, Fe)CO₃ (Brenker et al., 2007; Kaminsky et al., 2016; Phillips and Harris, 1995; Ryabchikov and Kaminsky, 2013) and CaCO₃ (Brenker et al., 2007; Meyer and McCallum, 1986) have been observed in inclusions in diamonds, including ultra-deep diamonds from the transition zone or possibly the lower mantle (Brenker et al., 2007). These inclusions demonstrate that both (Mg, Fe)CO₃ and CaCO₃ may be preserved by kinetic effects within carbonate-rich sediment in cold subducting slabs, consistent with slow breakdown kinetics observed in experiments at transition zone conditions (Martirosyan et al., 2016). The relative stability of (Mg, Fe)CO₃ and CaCO₃ may also change at depth due to polymorphism. If (Mg, Fe)CO3 or CaCO3 reaches the base of the lower mantle, density functional theory calculations (Oganov et al., 2008; Pickard and Needs, 2015) and experiments (Boulard et al., 2011; Ono et al., 2007) suggest these carbonates will transform to tetrahedrally-coordinated structures. If the transition in CaCO₃ occurs at a shallower depth than the analogous transition in MgCO₃, as predicted by Pickard and Needs (2015), CaCO₃ would be the denser and more energetically-favored carbonate in the deep lower mantle.

This study examines a Mg-Fe-Ca carbonate system in the petrologic context of a subducted carbonate assemblage in contact with metallic iron. Metallic iron may be present at the percent level throughout the mantle (Frost et al., 2004), and the outer core provides an unlimited reservoir of iron. Subducted carbonates may contact reduced core-mantle mixing regions if they reach depths within \sim 10s km from the base of the mantle (Frost and McCammon, 2008; Otsuka and Karato, 2012). Previous studies of carbonates at lower mantle pressures have investigated mineralogical stability of single phases as a function of pressure and temperature alone (Isshiki et al., 2004; Solopova et al., 2014; Boulard et al., 2011; Ono et al., 2005), or used mixtures of carbonates and silicates which complicate textural analysis of run products (Biellmann et al., 1993; Thomson et al., 2014). In contrast, we examine reaction interfaces between multiple carbonates and iron: similar previous studies have been limited to transition zone conditions (Martirosyan et al., 2016; Pal'yanov et al., 2013). We determine the relative stability and metastability of Mg-, Fe-,

Table 1

Pressure before heating (GPa)	Temperature (K)	Duration (min)
51	1800-2050	15
66	1900-2200	10
77	1900-2200	20
113	2400-2500	10

and Ca-carbonates as host phases for oxidized carbon storage in the lower mantle.

2. Methods

To bring experimental petrology of carbonates to the Mbar pressures and 1000s K temperatures of the core-mantle boundary, experiments must be miniaturized: sample geometry must be controlled within the \sim 50 \times 50 \times 10-micron chamber of the laser-heated diamond anvil cell, and analysis must be performed with \sim 100-nm-scale spatial resolution of typical grain sizes of run products. Dolomite, a major constituent of subducted marble, provides a unique advantage as a starting material as it has a mixed Mg-Fe-Ca carbonate composition homogeneous to <nm scale. Natural dolomite crystals with composition determined to be (Mg_{0.38}Ca_{0.59}Fe_{0.03})CO₃ by X-ray fluorescence spectroscopy and structure confirmed by X-ray diffraction were used as starting materials. Single dolomite crystals were polished to \sim 10-micron thickness and dried in a 120 °C oven overnight before loading. 3-micron thick 99.85% iron foil was machined into discs with a ps-pulsed 532-nm laser or cut with a razor blade. Iron foils were loaded sandwiched between dolomite crystals in symmetric diamond anvil cells. No other pressure standard or medium was loaded in order to prevent reactions with other components and contamination of the chemical system. Sample sandwiches were loaded within chambers cut in Re gaskets in diamond anvil cells. Diamond anvils with flat culets of 200 micron diameter or beveled culets of 150 micron diameter were chosen to generate pressures corresponding to the lower mantle.

Samples were compressed to pressures of 51, 66, 77, and 113 GPa, as determined by the equation of state of the iron foil (Mao et al., 1990) before laser heating (Table 1). Pressures were not measured during heating due to chemical reaction of the iron foil with the dolomite, but were likely \sim 7–10 GPa higher than those reported here due to thermal pressure. Laser heating was performed with *in situ* X-ray diffraction at the GSECARS sector of the Advanced Photon Source (Prakapenka et al., 2008). The laser spot was focused to a diameter of \sim 10 microns with a flat-top beam shape. Each sample was heated for 10–20 min at 1800–2500 K.

After recovery to ambient conditions, each heated spot was sectioned using a Zeiss NVision 40 dual-beam scanning electron microscope and focused Ga⁺ ion beam (Centre Interdisciplinaire Microscopie de Electronique (CIME), EPFL). Each sample was coated with 15 nm Au to reduce charging in the scanning electron microscope. To protect the samples from damage by the Ga⁺ ion beam, an additional \sim 1–2 micron thick layer of Pt or C was deposited across the centers of heated spots. Thin sections of each heated spot were extracted and polished to electron transparency (\sim 100 nm thickness).

Imaging of recovered thin sections was performed with scanning transmission election microscopy (S/TEM) and energy-dispersive X-ray spectroscopy (EDX) in a FEI Tecnai Osiris analytical TEM (CIME, EPFL). Accelerating voltage for these measurements was 200 kV. The Osiris TEM is equipped with four wide-solid-angle



Fig. 1. Backscattered scanning electron image recorded at 4 kV of dolomite-iron reaction interface in sample thin section recovered from 66 GPa and 1900–2200 K. The iron-rich region in the center appears brightest due to its high density. The extent of iron diffusion into and reaction with the Fe-poor carbonate is also evident based on density. A few micron offset in alignment of upstream and downstream lasers may be responsible for asymmetry of heated spot.

detectors for high-speed chemical measurements by EDX spectroscopy. EDX maps were scanned over 1024×1024 pixel areas with pixel dwell time of 50 microseconds. Typical count rates were \sim 40,000–50,000 counts per second. Chemical mapping prevents migration of elements due to damage by the electron beam. Uncertainties in compositions were determined from standard deviations of EDX measurements obtained from selected regions within multiple grains.

3. Results

Laser-heating dolomite-iron sandwiches from both sides produces a lens-shaped heated zone (Fig. 1), as predicted by models (Kiefer and Duffy, 2005; Rainey et al., 2013) and measurements (Campbell, 2008) of thermal diffusion in the diamond anvil cell. Laser absorption and diffraction indicate that the sample reaches a steady state within a few minutes. The hottest region of the sample is the laser-absorbing central iron layer. The insulating dolomite layers experience both axial and radial temperature gradients, with highest temperatures at the contacts with the iron foil and lowest temperatures at the diamond anvil surface. High-resolution TEM inspection of recovered thin sections indicates that grain sizes decrease away from the surface of the iron foil, consistent with this thermal gradient (Fig. 1). At the cool upper and lower edges of the heated spot, ribbons of magnesium and calcium carbonate exsolve from the dolomite (Fig. 2), as has been observed at lower pressures (Sato and Katsura, 2001; Shirasaka et al., 2002).

Both EDX analysis of recovered thin sections in the TEM (Fig. 3) and diffraction patterns obtained during and after sub-solidus laser heating (Fig. 4b) show that the metallic iron at the sample center reacts completely with the dolomite to generate a mixture of FeO and Fe₇C₃. The iron layer thus exhibits oxidation which must be compensated by reduction of the carbonate. Within the carbonate layer near the hot interface with the iron foil, three phases are found in all samples: diamond, Mg-bearing ferropericlase, and CaCO₃ (Fig. 3). The diffusion of iron from the sample center propagates breakdown of the magnesium carbonate to diamond and ferropericlase. Due to diffusion, this ferropericlase has much more MgO than the FeO at the sample center (Figs. 5–6). Unlike MgCO₃,



Fig. 2. Energy-dispersive X-ray image of a) exsolution of two carbonates from dolomite at 51 GPa and b) formation of $Fp + C + CaCO_3$ at 113 GPa. Elements color-coded as follows: red = iron, blue = calcium, green = magnesium, white = carbon. (For interpretation of the colors in the figure(s), the reader is referred to the web version of this article.)

CaCO₃ exsolves from the dolomite but is never observed to break down. EDX composition analysis confirms that the CaCO₃ phase retains both O and C within measurement error and there is no evidence for measurable Ca in any other phase (Fig. 5). At the pressures examined here, the stable polymorph of CaCO₃ is the postaragonite phase (Ono et al., 2005). Post-aragonite-type CaCO₃ is observed in diffraction patterns after heating (Fig. 4b), along with unreacted dolomite. These experiments show that a carbonate-rich rock such as subducted marble subjected to reducing conditions in contact with metallic iron, post-aragonite-type CaCO₃ is a more stable form of carbonate than MgCO₃ throughout the lower mantle pressure range.

4. Discussion

The mixture of (Mg, Fe)O, diamond, Fe_7C_3 and $CaCO_3$ is produced by redox coupling between the iron and carbon and breakdown of the MgCO₃ component. The following reactions explain these observations:

$$\begin{split} MgCO_3 + 2Fe &\rightarrow C + MgO + 2FeO \\ 3MgCO_3 + 13Fe &\rightarrow Fe_7C_3 + 3MgO + 6FeO \end{split}$$

In the center of the sample, the system is saturated in iron, leading to a mixture dominated by Fe_7C_3 and FeO. In the region where the carbonate breaks down, the carbon/iron ratio is higher, and reduced carbon is found in diamonds rather than carbides. The



Fig. 3. Composition measurements for dolomite-iron sample recovered from 113 GPa and 2400–2500 K. a) Energy-dispersive X-ray map with elements color-coded as follows: red = iron, blue = calcium, green = magnesium, white = carbon. b) Examples of EDX spectra obtained for each phase.

breakdown of MgCO₃ is favorable in all heated regions in the presence of iron. The stable host phase for carbon is controlled by the Fe/MgCO₃ ratio of reactants. Recent studies have largely focused on (Mg, Fe)CO₃ as a host phase for oxidized carbon in the deep Earth, as the melting point of magnesite or ferromagnesite is higher than that of CaCO₃ in the transition zone. However, (Mg, Fe)CO₃ is much more susceptible to redox breakdown than CaCO₃.

Redox breakdown will promote diamond formation from (Mg. Fe)CO₃ in carbonated eclogite and peridotite. This study and others have shown that diamonds nucleate and grow directly from carbonates at subsolidus, reducing conditions, and inclusion-bearing diamonds used as evidence that carbonates are stable to great depths in the mantle record incomplete redox breakdown (e.g. Wang et al., 1996). Reducing agents relevant to carbonate stability at different stages of Earth's chemical evolution include graphite (Pal'yanov et al., 1999), H₂ (Pal'yanov et al., 2002), Si metal (Siebert et al., 2005), and Fe metal. In a mixture of carbonate and Fe-Si alloy, Si and C were oxidized, producing a mixture in which diamonds coexist with metallic iron (Siebert et al., 2005). Other studies have suggested either that pure iron and pure carbon should not coexist in equilibrium (Rouquette et al., 2008: Scott et al., 2001) or that iron carbides and diamond compete as hosts of reduced carbon (lizuka et al., 1996). In this study, metallic iron reacts completely to carbide and oxide, indicating that the system is oversaturated in carbon. Because the mantle has been suggested to be a highly reducing environment, saturated in metallic iron (Frost et al., 2004; Frost and McCammon, 2008; Rohrbach et al., 2007; Rohrbach and Schmidt, 2011), carbonate in this system will be susceptible to redox breakdown at subsolidus temperatures. Reduction of subducted carbonates by interaction with metallic iron was previously explored at relatively modest depths (Martin and Hammouda, 2011: Martirosvan et al., 2015a, 2015b; Stagno et al., 2011). Most previous studies were conducted at pressures reaching \sim 6 GPa (Martin and Hammouda, 2011; Martirosyan et al., 2015a, 2015b), too low to model the depths at which metallic iron becomes available to react with subducted carbonate. Metallic iron may also be present in the lower mantle due to charge disproportionation in silicates (Frost et al., 2004; Rohrbach et al., 2007), mixing at the CMB (Knittle and Jeanloz, 1989; Otsuka and Karato, 2012), or residue of magma ocean solidification (Zhang et al., 2016). Recently, iron-magnesite mix-



Fig. 4. X-ray diffraction patterns observed before and after transforming samples of dolomite (d) and iron foil (Fe) to magnesiowüstite and ferropericlase (gold Mw and Fp), Fe₇C₃ (red), and post-aragonite-type CaCO₃ (blue) at pressures 51–113 GPa and temperatures 1800–2500 K in the laser-heated diamond anvil cell. Additional diffraction peaks from high-pressure polymorphs of dolomite are labeled dII and dIII. For these experiments $\lambda = 0.3100$ Å.



Fig. 5. Compositions for observed phases obtained from EDX spectra plotted on Mg-Fe-Ca ternary diagrams. At 49 and 64 GPa, Fp and Mw grain sizes are too small to resolve compositions by EDX. Ranges of compositions may also be due to grain overlaps.



Fig. 6. Observed unit cell volumes for Mw (black) and Fp (green) in this study plotted relative to previously-measured equations of state for Fp–Mw compositions with different Fe# = Fe/(Mg + Fe) (Fei et al., 2007; Fischer et al., 2011; Speziale et al., 2001). Pressures for data in this study are assumed to be the same as pressures obtained from the Fe foil before heating. (Mg,Fe)O diffraction peaks are split, indicating unit cell volumes consistent with coexistence of FeO at the sample center and \sim (Mg_{0.6}Fe_{0.4})O produced by breakdown of the dolomite. A decrease in unit cell volume observed in both oxides between 66 and 77 GPa is consistent with the spin transition in iron-rich Mw and structural change in FeO under these conditions.

tures were examined at 16–45 GPa and 1500–1700 °C, corresponding to transition zone to lower mantle depths (Stagno et al., 2011). At these conditions, diamonds are observed to coexist with magnesite, ferropericlase and carbon-free iron-iridium alloy. The Ir alloy was used to determine that fO_2 was \sim 3 log units above the ironwüstite buffer. The breakdown of magnesite observed here may imply lower fO_2 conditions, perhaps due to the sample environment in the diamond anvil cell or higher pressures. For subducted marble or carbonated silicate lithologies that reach a reduced zone near the core-mantle boundary, (Mg, Fe)CO₃ cannot be preserved.

Spin transitions in ferromagnesite have recently been studied for their potential to affect carbonate stability (Lin et al., 2012; Liu et al., 2015). The spin transition results in a ~8% unit cell volume collapse for (Mg_{0.35}Fe_{0.65})CO₃ (Lin et al., 2012) and would thus be expected to have a significant effect on phase equilibria. As the minimum pressure examined in these experiments is above the spin transition pressure, we are unable to evaluate whether the spin transition is related to the observed breakdown of (Mg, Fe)CO₃. The susceptibility of (Mg, Fe)CO₃ to disproportionation at low fO_2 suggests that low-spin iron-bearing carbonates are unlikely to be stable in Earth's mantle.

CaCO₃ is less sensitive than (Mg, Fe)CO₃ to redox breakdown, but melting, reaction with silicates, and buoyancy remain challenging barriers to its transport to the deep mantle. Although the melting point of CaCO₃ (Bayarjargal et al., 2010; Thomson et al., 2014) is well above a 1600 K adiabatic geotherm (Brown and Shankland, 1981), the eutectic melting point for carbonate mixtures is substantially lower and comparable to the geotherm (Thomson et al., 2014). Eutectic melting has been argued to effectively block carbonate transport to the lower mantle (Thomson et al., 2016). However, no evidence is observed for melting of the dolomite carbonate in this work. Moreover, geotherms both for the average mantle (e.g. Brown and Shankland, 1981; Andrault et al., 2011; Nomura et al., 2014) and subducting slabs (e.g. Andrault et al., 2014) remain highly uncertain. The key question is whether, for realistic subducted carbonates, the melting point of the mixture is high enough relative to the temperature in the slab to prevent diamond formation. With temperature uncertainties of at least 100s K, in addition to potential kinetic effects on the efficiency of carbonate reduction and melting (Martirosyan et al., 2016), preservation of carbonates in the lower mantle cannot yet be ruled out. CaCO₃ is also expected to be destabilized in the mantle by reaction with (Mg, Fe)SiO₃ in peridotite or eclogite to produce CaSiO₃ and (Mg, Fe)CO₃ (Biellmann et al., 1993; Seto et al., 2008). This reaction suggests CaCO₃ is most likely to be preserved in silicate-poor, dolomite-rich marble entrained with subducting slabs. Because metasediment is likely to be as much as \sim 5–15% less dense than surrounding peridotite, \sim 100 m packages of marble would likely rise to the surface through diapirism (Kelemen and Manning, 2015). Transport of CaCO₃ to the deep mantle would require carbonate to reside in layers or regions thin enough to be entrained with the cold slab, but thick enough that slow diffusion blocks reaction of carbonate with silicates.

Observations in this and other experimental studies challenge interpretation of oxide inclusions in identified in diamonds. While traditionally (Mg, Fe)O inclusions have been considered to be a smoking gun for formation in the Earth's lower mantle, as a leftover from the breakdown of Mg_2SiO_4 to lower mantle bridgmanite (Kaminsky, 2012), carbonate disproportionation is another possible mechanism for oxide formation (Tappert et al., 2005; Thomson et al., 2016). Direct observation of lower mantle silicates of appropriate compositions is necessary to confirm ultra-deep diamond origin.

5. Conclusions

Magnesite-siderite is not a likely host phase for carbonate in reduced regions of the deep lower mantle, but post-aragonite-type $CaCO_3$ may be stable in carbonate-rich lithologies. While $CaCO_3$ undergoes more pressure-induced polymorphic transitions than (Mg, Fe)CO₃ in the deep mantle and may be more likely to react with lower mantle silicates, relative to (Mg, Fe)CO₃ it is less likely to participate in redox reactions. A low fO_2 environment in the lower mantle (Frost and McCammon, 2008; Rohrbach and Schmidt, 2011) will promote breakdown of ferromagnesite but CaCO₃ will remain stable to pressure, temperature and redox conditions near the base of the lower mantle if isolated from silicates. A carbonate-rich heterogeneity near the base of the mantle could be a high-pressure marble, with CaCO₃ associated with diamonds formed by breakdown of MgCO₃.

Diamond-bearing marble in the lower mantle is not likely to be directly observable by seismic tomography, but could impact observations of heterogeneous redox conditions at depth due to mantle convection and subduction and/or signatures of Earth's chemical evolution. Heterogeneous oxygen fugacity at depth is evident in diamond inclusions (Brenker et al., 2007; Walter et al., 2011; Ryabchikov and Kaminsky, 2013; Smith et al., 2016; Kaminsky, 2012) as well as chemical variation in both mid-ocean ridge basalts due to relative contributions of depleted and primitive mantle sources (Cottrell and Kelley, 2013) and ocean island basalts potentially sourced from the core-mantle boundary (White, 2010). The interpretation of available redox proxies for the deep mantle over deep time, particularly presence of carbonate (e.g. Brenker et al., 2007) and speciation of iron (Frost and McCammon, 2008), must be evaluated in the light of the multivariable dependence of stability of MgCO₃ and CaCO₃ on temperature, bulk composition of the phase assemblage and fO_2 .

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Data reported in this paper are available by request from S.M. Dorfman at dorfman3@msu.edu.

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