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Shock-induced transformation of olivine to a new metastable
(Mg,Fe)₂SiO₄ polymorph in Martian meteorites

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Introduction

Ferromagnesian olivines form the major silicate fraction of the mantle of terrestrial planets that also predominates in chondritic meteorites. The $(\text{Mg,Fe})_2\text{SiO}_4$ phase diagram at high-P,T conditions is well established from experiments [1, 2]. The Mg_2SiO_4 end-member (forsterite; Fo) transforms first into wadsleyite- Mg_2SiO_4 and then spinel-structured ringwoodite- Mg_2SiO_4 at $P = 12\text{-}22$ GPa depending upon the T, before disproportionating into MgSiO_3 perovskite + MgO at $P = 25$ GPa. Fe_2SiO_4 olivine (fayalite, Fa) transforms directly into the γ -spinel polymorph at $P = 6\text{-}7$ GPa. High-P,T transformations of Fo,Fa olivine solid solutions lead to the high-P minerals wadsleyite or ringwoodite (i.e., β -, γ - $(\text{Mg,Fe})_2\text{SiO}_4$). These phase transitions explain the major seismic discontinuities observed within the Earth's upper mantle and transition zone [3, 4]. Wadsleyite and ringwoodite are also documented within chondritic meteorites, where they are formed by shock metamorphism of $(\text{Mg,Fe})_2\text{SiO}_4$ olivines. Melting is also reported to occur in certain meteorite samples. Here we studied olivine grains within the newly-described Martian meteorites NWA2737 and NWA1950 [5, 6]. Meteorite NWA1950 is a lherzolitic shergottite with a gabbroic cumulate texture. It contains 55% olivine (Fo_{66} to Fo_{75}) crystals identified by optical microscopy. NWA2737 is the second example of a chassignite meteorite, found in the Western Sahara. It is an olivine-cumulate (dunite) containing minor pyroxenes and feldspar glass. It differs slightly from Chassigny by the dark colour of its olivine crystals (Fig. 1). The olivines in NWA1950 also appear black in hand specimen.

Petrological observations

From optical and Back-Scattered Electron (BSE) microscopy, individual olivine grains in the two meteorites are clearly outlined by secondary calcite that fills grain boundaries as well as fractures within the grains. The shocked olivine crystals also contain light-coloured stripes.

Some grains exhibit sets of subperpendicular stripes, crosscut by calcite-filled fractures (Fig. 1). In BSE images, these stripes appear dark, indicating a density and/or compositional contrast with the surrounding "olivine" crystals. Micro-Raman spectra of the clear stripes exhibit sharp peaks characteristic of $(\text{Mg,Fe})_2\text{SiO}_4$ olivine (Fig 1). The Raman spectra of the dark zones are different. Although they exhibit main features of the normal olivine spectrum, the peaks are broadened. A broad background extends from 100 to 600 cm^{-1} , and also under the high frequency region (800-1000 cm^{-1}). Such broad features could be attributed to highly disordered olivine crystals formed during the shock process. In a few crystals, a weak peak observed at 760 cm^{-1} might be assigned to formation of SiOSi linkages or highly coordinated (SiO_5 , SiO_6) species as "defects" within the structure [7]. However, most grains examined did not show such a feature. Instead, the spectra showed additional broad bands at 650 and 686 cm^{-1} that cannot form part of the olivine spectrum, because the 600-800 cm^{-1} region corresponds to a well-defined gap in the vibrational density-of-states (VDOS) function [8, 9]. Weak peaks are observed at the same position in experimentally shocked olivine at 59 GPa [10]. Also, the characteristic Raman mode of olivine at 600 cm^{-1} is missing (Fig. 1). Micro-Raman mapping experiments showed that the "altered" olivine spectrum was homogeneous within both meteorite samples, both for individual grains and among different crystals. The results indicate the presence of a new phase that is structurally related to but is different from olivine, produced metastably during the shock process. We term this new orthosilicate polymorph ζ - $(\text{Mg,Fe})_2\text{SiO}_4$ for convenience in the discussion below.

HRTEM observations

We determined the chemical composition of the dark zones and clear stripes within the shocked olivines using a combination of electron microprobe and nanoscale X-ray Energy-Dispersive analysis (ED) in the TEM studies. No difference in composition could be

established. For HRTEM studies we selected a single dark grain from NWA2737 and prepared 50-250 nm thick sections by diamond microtome (Diatome, Switzerland). The HRTEM study confirmed that the sample was entirely crystalline. Within a very few regions, selected area electron diffraction (SAED) patterns could be analysed within space group $Pbnm$ with lattice parameters $a = 0.4777$, $b = 1.026$, $c = 0.601$ nm consistent with olivine. However, most HRTEM images indicated considerable distortion of the lattice planes, and the Fourier transformed (FT) images could no longer be indexed as olivine, although the O^{2-} anion sublattice maintained hexagonal (*hcp*) symmetry (Fig. 2). Similar results were reported for olivine crystals shocked in the laboratory and also in laser-heated Diamond Anvil Cell (DAC) experiments [11, 12]. We observed 5-30 nm metallic inclusions distributed among the shocked olivine crystals (Fig 2). EDX analysis within the TEM showed that the inclusions consisted of Fe_xNi_y alloys with their structure and composition dependent upon particle size. Asymmetric $Fe-K_\beta$ and $Ni-K_\alpha$ peaks indicate the presence of a few amount of Co (too low to be quantify) which is consistent with Co enrichment in metal with respect to olivine [13]. The nanoparticles cause the black colour of the shocked olivines observed in hand specimen and thin section. The appearance of Fe_xNi_y in the olivine samples results metal ion reduction and migration during the shock process (well detailed in [14]). The HRTEM observations show that the olivine structure immediately surrounding the metal nanoparticles is highly ordered (Fig. 2). So far, the clear stripes observed within the black olivines do not contain metallic particles.

Molecular Dynamics simulations

We carried out Dynamics simulations (MD) to identify the nature of the new metastable Mg_2SiO_4 phase and its formation from metastably compressed olivine by shock transformation. An Anisotropic-Ion Model (AIM) is employed [15], in which induced ion

moments and short-range size and shape deformations of O^{2-} are included to quadrupolar level. The AIM is parameterised using high-level electronic structure calculations [16]. We show calculated $V(P)$ relations for α -, β -, γ - and ζ -phases of Mg_2SiO_4 generated within a simulation box with 672 ions under constant P (variable V and cell dimensions including angles) at $T=300K$ (Fig. 3). The new ζ - Mg_2SiO_4 polymorph has a density intermediate between α - and β - Mg_2SiO_4 . It is produced in the simulation by applying a rapid P increment to metastably compressed α - Mg_2SiO_4 at $P\sim 50$ GPa, mimicking the actual shock conditions. The new structure maintains an *hcp* O^{2-} sublattice and isolated SiO_4^{4-} units: it can be correlated with the olivine structure by a model proposed by Hyde et al. [17]. These authors noted that the metal cation sublattice in olivine corresponds to an alloy structure Ni_2In , with $Si(Mg,Fe)_6$ trigonal prisms linked by face-sharing to form a corrugated-prismatic structure. The γ - $(Mg,Fe)_2SiO_4$ spinel structure instead gives rise to truncated-tetrahedral $Si(Mg,Fe)_{12}$ units as found in the alloy $MgCu_2$. Both structures maintain isolated SiO_4^{4-} units and *hcp* O^{2-} packing. The ζ - Mg_2SiO_4 structure produced by MD shock compression is related to olivine by reorganisation of the metal sublattice resulting in an increase in the next-nearest-neighbour cation coordination (Fig. 4). During natural and experimental shock conditions, substantial disorder in the cation packing can result from the solid-state transformation, consistent with the HRTEM and Raman results. The AIM-MD simulations show that ζ - Mg_2SiO_4 persists metastably upon decompression to $P = 1$ atm. For comparison with experiment, we calculated the Vibrational Densities Of States (VDOS) and the SiO_4^{4-} stretching and bending mode contributions that dominate the Raman spectrum (Fig. 3). The VDOS are calculated both from the dynamical matrices (an instantaneous normal mode analysis) and by constructing time correlation functions which directly probe the underlying silicon-centred local modes of vibration [18]. The VDOS for the simulated ζ - Mg_2SiO_4 is similar to that of olivine but is

broadened so that features appear in the range 600-800 cm^{-1} , as observed experimentally (Fig. 1, 3).

Discussion

The conditions attained during shock are constrained by the absence of high-P wadsleyite or ringwoodite phases. The peak shock pressure in the analogous Chassigny meteorite was estimated as $P \sim 35 \pm 5$ GPa from planar fractures within olivine grains, and is consistent with the observation of small (250-500 nm in size) crystals of wadsleyite and ringwoodite, indicating likely similar shock conditions for NWA2737 and NWA1950. The absence of melting indicates that T did not exceed 2500 K. Using known Fe-Ni-Mg interdiffusion coefficients for olivine we can constrain the time-temperature (t-T) conditions associated with the nanoparticle formation. The average spacing observed between nanoparticles determines a characteristic diffusion length $x = 50 \pm 10$ nm. Using $x = (Dt)^{1/2}$, we calculate diffusion timescales on the order of $t \sim 10$ ms for $T = 2100 \pm 100$ K and ~ 1 s for $T = 1600 \pm 100$ K. These values correspond well with the expected duration of typical thermal events following an initial shock of this magnitude [19].

The mechanisms of transformations between α -, β - and γ -polymorphs of $(\text{Mg,Fe})_2\text{SiO}_4$ have been studied extensively [20-23]. A series of "spineloid" structures exists for AB_2O_4 compounds between γ -spinel and a hypothetical structure designated the ω - or ϵ^* - phase, formed by shifting spinel blocks relative to each other along crystal planes to create Si-O-Si linkages [17, 24]. β - Mg_2SiO_4 corresponds to one spinelloid polymorph: others have been discovered in the Ni_2SiO_4 - NiAl_2O_4 and NiGa_2O_4 system, among Mg-gallogermanates and in fayalite-magnetite solid solutions [25-28]. Formation of the ω - or ϵ^* - phase provides an intermediate step in α -(β , γ) phase transformations and leads to other spinelloid structures [17, 24]. However, the appearance of the distorted olivine phase observed following shock

compression and described above cannot be explained by the same transformation mechanism. The spinelloids are based on cubic packing of the O^{2-} sublattice, rather than the *hcp* symmetry observed here, and they contain SiOSi linkages between polymerised SiO_4 units that would give rise to strong Raman peaks both at $600-700\text{ cm}^{-1}$ and $950-1100\text{ cm}^{-1}$ [29, 30]. Instead, the Raman spectra of the shocked olivines remain characteristic of an orthosilicate structure containing isolated SiO_4^{4-} units (Fig. 1). It is thus likely that the new ζ - $(Mg,Fe)_2SiO_4$ polymorph might be present in many shocked ultrabasic rocks including meteorites, but its presence has remained undetected due to its structural and spectroscopic similarities with olivine (Fig. 3).

We calculated stable and metastable $G(P)$ relations for α -, β -, γ - and ζ - polymorphs of Mg_2SiO_4 ($G=H-TS$ is Gibbs' free energy; H and S are enthalpy and entropy) (Fig. 4). The AIM-MD model predicts α - β and β - γ transitions at $P = 12$ and 13 GPa (compared with $P=9$, 13 GPa from experiment after linear extrapolation to $T = 0\text{ K}$ [1, 2]). The high kinetic barriers associated with these transitions allow olivine to be compressed metastably beyond these limits at low T [7]. The predicted ζ - polymorph becomes stabilised relative to olivine at 38 GPa . At ambient P , the free energy of the ζ -phase lies between those of α - and β - Mg_2SiO_4 so that metastable back-transformation of wadsleyite can result in formation of the ζ -polymorph. This analysis agrees with experimental studies by Raman spectroscopy and X-ray diffraction, that show formation of a new metastable structure after "gentle" heating of metastably-decompressed β - Mg_2SiO_4 recovered from high- P,T synthesis experiments [31-33].

We have thus discovered a new metastable polymorph of $(Mg,Fe)_2SiO_4$ orthosilicate within Martian meteorites in which the olivine phase was subjected to $P > 35\text{ GPa}$ shock pressure at low T . The conditions were not sufficient to cause melting or transformation to stable high- P polymorphs. Such shock conditions are common among natural impacts [34].

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Figure captions

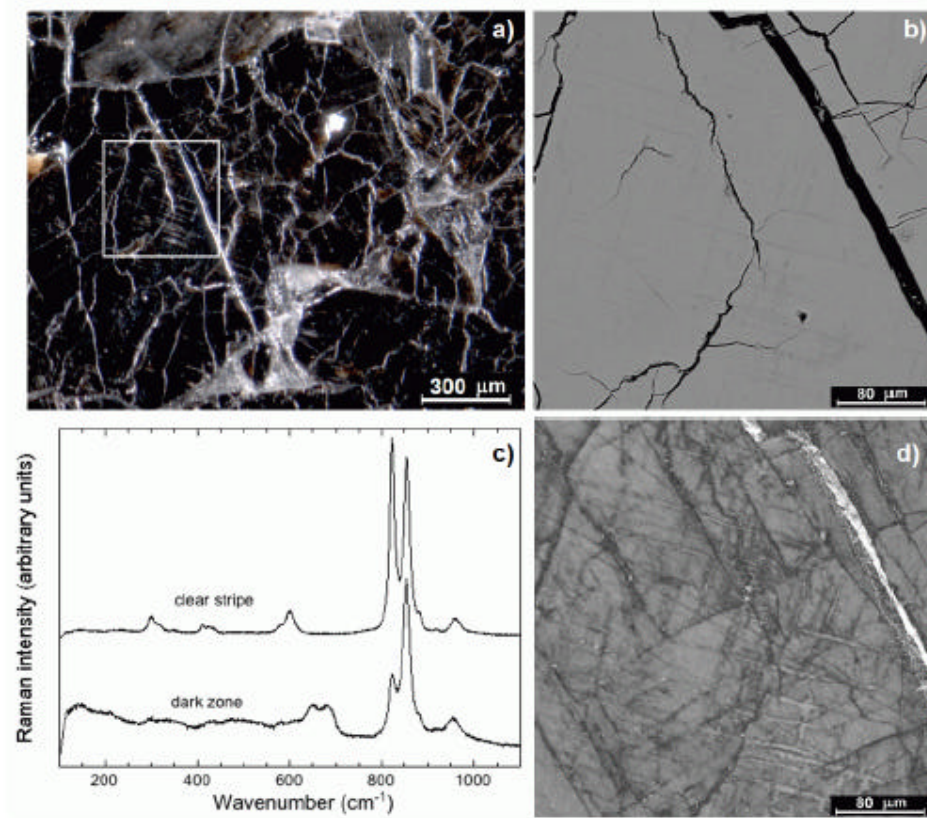
Figure 1. Optical and electron microscopy of shocked olivine samples and Raman spectroscopy results. (a) Optical micrograph of a polished section of NWA2737 illustrating the unusual dark colour of the olivines in these rocks. Some clear bands are observed within the black olivine grains, with widths up to about 50 μm , oriented along two subperpendicular directions corresponding to $\{021\}$ crystallographic planes. (b) Back-scattered electron (BSE) images show contrast between the clear bands and the majority black grains that reveals an electron density difference between the two. (c) Raman spectra recorded within the clear stripes correspond to pristine olivine. The spectra obtained within the dark zones resemble olivine upon first examination but they contain unusual features that can not be assigned to an olivine structure. In particular there are broad bands in the 600-700 cm^{-1} range that corresponds to a forbidden region within the VDOS of olivine crystals; also the characteristic 600 cm^{-1} peak of olivine is absent. The spectrum is reproduced throughout the majority dark crystalline material found in shocked meteorite samples NWA2737 and NWA1950, and appears constant from micro-Raman mapping studies. The spectrum likely indicates the presence of a new $(\text{Mg,Fe})_2\text{SiO}_4$ polymorph. d) band contrast image from EBSD mapping on the same area as (b). The bright lamellae consist of well-oriented olivine ($< 2^\circ$ difference from point to point) and the darker majority zones to olivine-like pattern with large misorientations ($5\text{-}10^\circ$) consistent with hcp structure domains.

Figure 2. High Resolution TEM images of shocked olivines within the Martian meteorite sample NWA2737. (a) HRTEM images of the new orthosilicate ζ -polymorph derived from $(\text{Mg,Fe})_2\text{SiO}_4$ olivine by shock compression. The Fourier transform of the HRTEM image is readily indexed to an hcp lattice of O^{2-} ions and disordered cation positions with lattice

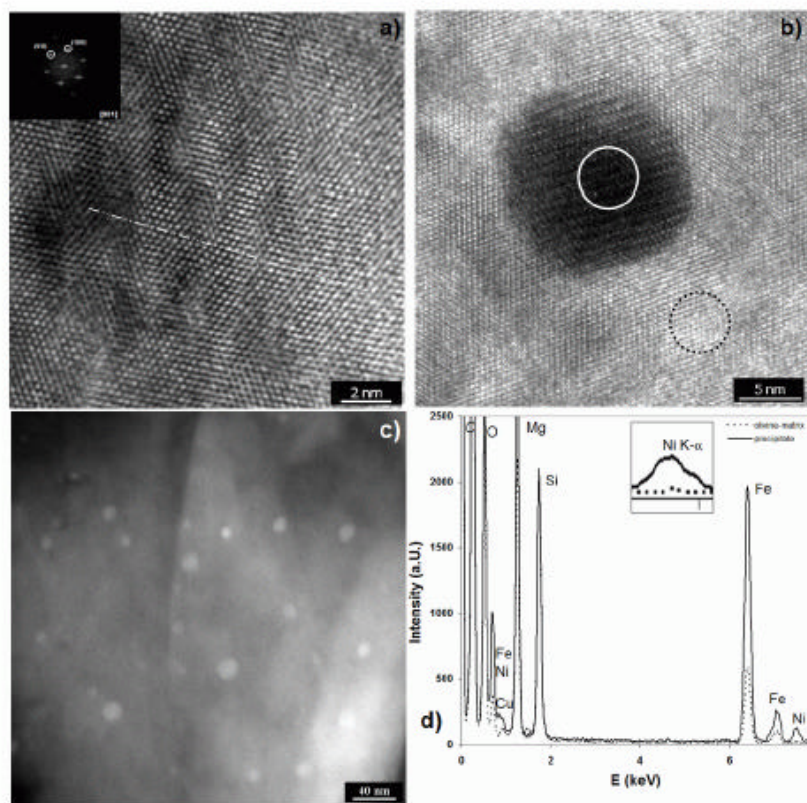
parameters $a \sim 0.3$ nm and $c \sim 2a$. Such a lattice has been proposed for high-P low-T transformations of olivine following DAC experiments[12]. (b) HRTEM image of a metallic Fe_xNi_y particle (15 nm diameter) contained within the shocked olivine. (c) Annular dark field TEM image of sub-spherical metal inclusions contained within the shocked olivine grains. (d) EDX spectra (normalized to the Si- K_α peak) in the precipitate and olivine matrix analysis spots (probe size 5 nm) depicted in (b). Deconvolution of analytical data yields a composition $\sim \text{Fe}_{0.93}\text{Ni}_{0.07}$ for the inclusion.

Figure 3. MD results. (a) V(P) relations showing the evolution of the high pressure ζ -phase. (b) Static (0 K) energy/volume relations for the α , β , γ and ζ phases. The figure also shows the dynamic (300 K) evolution of the system energy with the arrow highlighting the location of the pressure-driven phase transition. (c) Vibrational densities of states calculated from time correlation functions constructed to reflect the vibrational modes of the SiO_4 tetrahedra. From bottom to top, the first three curves show the effect of pressure on the α -phase, whilst the upper two curves show the high and low pressure states for the ζ -phase. (d) X-ray diffraction patterns calculated from the simulated structure factors for the α -phase starting material (lower curve) and the high pressure ζ -phase (upper curve).

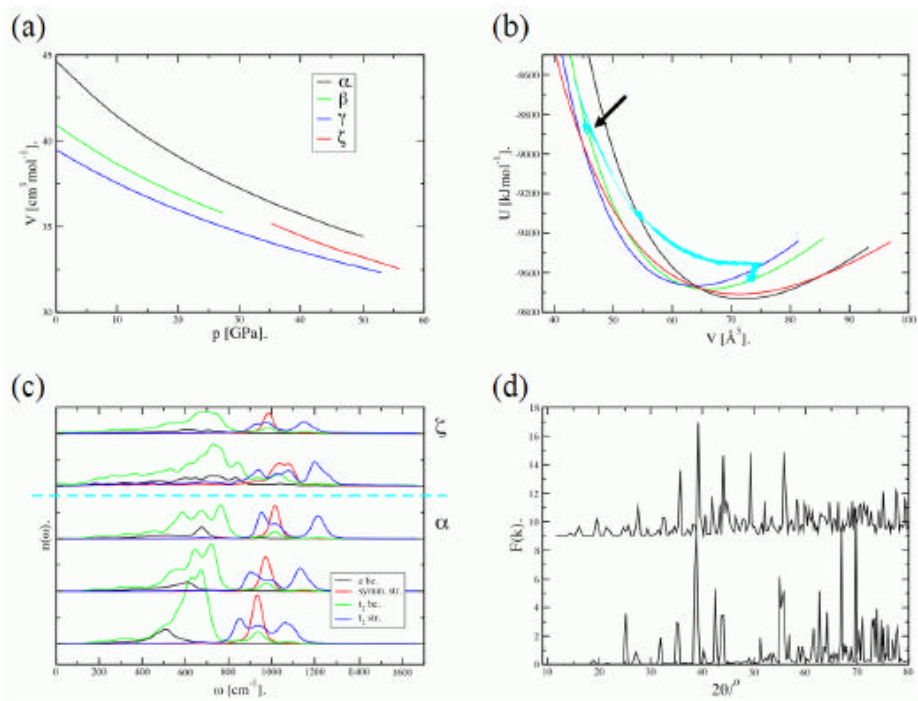
Figure 4. (a) Simulated free energy curves calculated as a function of pressure. The free energies are expressed relative to that of the α -phase in order to emphasize the thermodynamic phase relations. (b) Molecular dynamics snapshot of the high pressure ζ -phase shown in the ab plane. The Mg, Si and O atoms are shown as the red, blue and magenta circles respectively. The green box highlights the underlying unit cell.



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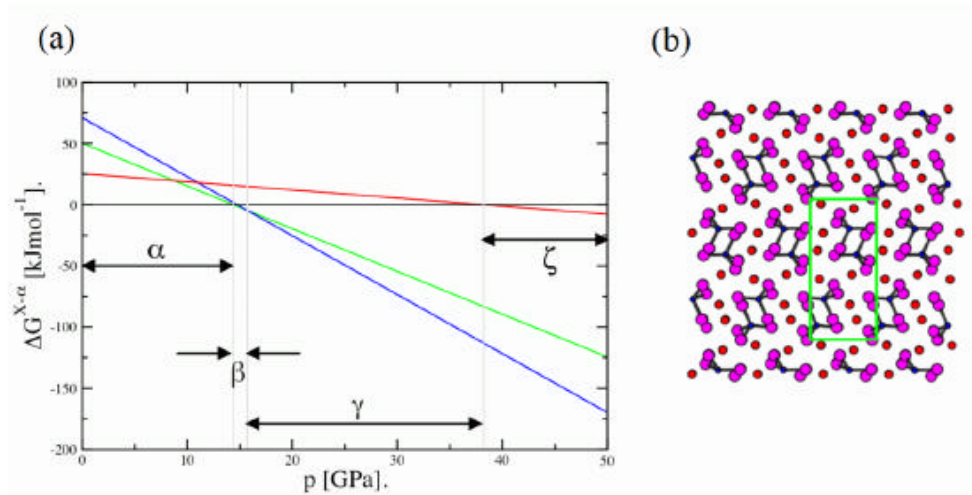


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