Pressure-Induced Phase Transitions of Natural Brookite

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Supporting Information

ABSTRACT: The structures and stabilities of the high-pressure phases of titania (TiO2) are of great interest in the earth sciences, as these phases are the accessible analogs of component minerals in the earth’s mantle. Brookite is a natural titania mineral, whose bulk phase is hard to synthesize in the laboratory, and its phase behavior at very high pressure remains unknown. Thus, in this work, using phase-pure natural brookite as the sample, we studied the phase transition of bulk brookite under compression up to ~60 GPa in three different pressure transmitting media. Results show that, at room temperature and in near hydrostatic conditions, brookite undergoes a series of transitions, i.e., brookite (~0–12 GPa) → TiO2-II and minor rutile (~1–21 GPa) → baddeleyite (~12–60 GPa) → TiO2-OI (~33–60 GPa). Taking into account that all transitions occur at finite rates and that solidification of a pressure medium can influence transition kinetics, we show that the observed transition sequence is consistent with the expected high-pressure phase stability of TiO2 calculated from density functional theory. The knowledge obtained from this work may be used to infer the geological fate of brookite in nature.

KEYWORDS: titania, high pressure, phase transformation, nonhydrostaticity, pressure transmitting medium, shear modulus, equation-of-state, natural mineral

INTRODUCTION

As a technologically important material, titania (TiO2) has applications in photocatalysis, gas sensors, energy storage, biotechnology, and cosmetics, etc. Natural titania exists mainly in the three polymorphs, rutile (tetragonal structure), anatase (tetragonal structure), and brookite (orthorhombic structure). Studies show that, at ambient pressure, rutile is more stable than anatase at temperatures up to at least ~1000 °C. Anatase and brookite are metastable with respect to rutile, and hence, they can transform to rutile upon heating above certain temperatures. In nature, brookite often occurs in quartz veins of regional metamorphic rock series or as an accessory mineral in igneous rocks. Brookite is relatively rare compared to rutile and anatase. It is very difficult to synthesize bulk brookite in the laboratory, though nanocrystalline brookite can be prepared relatively easily in the laboratory using different methods. Previous high-pressure (HP) studies on TiO2 predominantly focused on the phase transitions of bulk or nanocrystalline anatase and rutile. Titania phases involved in the subsequent high-pressure phase transitions include columbite (TiO2-II, space group P63), baddeleyite (space group P21/c), TiO2-OI (space group Pmnn) and TiO2-OI (space group Pnma). We only found scarce reports on high-pressure behaviors of nanocrystalline brookite at pressures up to ~28 GPa and bulk brookite at pressures up to ~10 GPa. The phase behavior of bulk brookite at very high pressures (~>10 GPa) remains unknown possibly because of the unavailability of bulk brookite from laboratory syntheses. However, such knowledge is needed to understand the detailed formation mechanism and stability of brookite. Thus, in this work, using phase-pure natural brookite as the starting material, we studied the pressure-induced phase transitions of natural brookite at room temperature and at pressures up to ~60 GPa using in situ high-pressure X-ray diffraction (XRD) and Raman spectroscopy. We aim to obtain knowledge about the phase behavior and structural stability of bulk brookite under compression to very high pressures (and in subsequent decompression as well) so that insights into the history and possible fate of natural brookite minerals can be generated with new experimental data.

EXPERIMENTAL SECTION

Sample. The natural brookite sample used in this study was from the mineral collection no. 8115 of the Department of Earth and Planetary Science, University of California Berkeley.

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The mineral was sourced from Magnet Cove, AR, USA. It is similar in appearance to those described in refs 8 and 9. A few brookite crystals with well-developed morphologies were chosen. The crystals were immersed in a nitric acid solution (1 mol L\(^{-1}\)) to dissolve their surface stains and then washed with deionized water before drying and crushing into small pellets. Then, the pellets were ground with a mortar and pestle to very fine powders that were used for the high-pressure study in this work (below).

**X-ray Powder Diffraction.** To examine the crystal structures of the natural TiO\(_2\) sample, XRD patterns were obtained using an X-ray powder diffractometer (Malvern Panalytical) with Cu K\(\alpha\) radiation (wavelength, 1.54056 Å). The XRD patterns were collected from \(2\theta = 10\) to 80° at a scanning rate of 5°/min.

**Scanning Electron Microscopy and Energy Dispersive Spectroscopy.** The morphologies, particle sizes, and elemental compositions of the ground titania particles were examined using a scanning electron microscope (SEM; VERSA 3D, FEI) equipped with an energy dispersive spectrometer (EDS), which were operated under 5.0 kV/27 pA and 15.0 kV/0.21 nA, respectively.

**In Situ High-Pressure XRD and Raman Spectroscopy.** A diamond anvil cell (DAC) was used to generate the required high pressure of the sample. The cuvet size of the anvils was 300 µm in diameter. A stainless steel gasket was preindented to 23 µm in thickness, and then a hole of 150 µm in diameter was drilled through the center of the indentation using laser drilling (at HPSTAR), serving as the sample chamber. A tiny amount of the finely ground brookite powder was compressed into a thin foil at a pressure of \(\sim 8\) MPa using a powder pressing machine. Then, one piece of the thin foil \(\sim 15\) µm thick and an appropriate size for the sample chamber were chosen for loading into the DAC together with a few grains of ruby as the pressure transmitting medium for the HP-XRD measurements; a methanol mixture (4:1 in volume) was used as the pressure transmitting medium for the HP-XRD measurements. In situ HP-XRD measurements were performed at the 12.2.2 beamline station of the Advanced Light Source (ALS) at room temperature.\(^5\)\(^,\)\(^\textit{12,32}\) HP-XRD patterns were collected at an X-ray wavelength of 0.4959 Å using a MAR345 image plate detector. The sample-to-detector distance was calibrated using a CeO\(_2\) standard. After compressing the sample near the required pressure value and equilibrating the system for \(\sim 10\) min, the sample pressure was measured using laser fluorescence (excitation laser wavelength, 447 nm). Then, the HP-XRD measurement at the given pressure was conducted in the beamline station, taking \(\sim 5\) min to complete. The HP-XRD and the pressure measurements were repeated at all pressures in the full compression−decompression cycle.

Additional HP-XRD data were collected at the 13-BM-C beamline station of the Advanced Photon Source (APS) at room temperature with an X-ray wavelength of 0.4340 Å. At 13-BM-C, a MAR165 CCD detector was used to collect the diffraction image and a NIST standard LaB\(_6\) sample was used to calibrate the sample-to-detector distance and the detector tilt. The pressure measurements were similar to those at ALS-12.2.2 (above) except that the excitation laser had a wavelength of 532 nm. The sample pressure was measured after equilibrating the system for \(\sim 6\) min, and the HP-XRD measurement at each given pressure took \(\sim 2\) min to complete.

The diffraction images were integrated using the Dioptas program\(^23\) to yield numerical intensity vs 2θ data. Rietveld-fitting analyses were performed on the collected XRD patterns using GSAS software\(^24,25\) to identify and/or verify the phases present and to obtain the phase contents of the titania phases as a function of pressure.

High-pressure Raman data were collected from 100 to 800 cm\(^{-1}\) at room temperature using a Raman spectrometer (inVia Reflex, Renishaw) with a 532 nm excitation laser at \(\sim 10\)% of the laser power (83.3 mW). The integration time was 0.029 s for each incremental step of 1 cm\(^{-1}\). At a given pressure, the collection of one Raman spectrum took \(\sim 20\) s.

**COMPUTATIONAL SECTION**

The enthalpies of the concerned TiO\(_2\) phases (brookite, anatase, rutile, baddeleyite, TiO\(_2\)-II, and TiO\(_2\)-OI) at different pressures were calculated using density functional theory (DFT) implemented in the Quantum Espresso (QE) package.\(^26\) The computations were conducted using an integrated graphic user interface and a running environment of QE, Burai (v.1.3.2).\(^27\) The projector augmented-wave (PAW) pseudopotential functions\(^28\) of Ti and O with an energy cutoff of \(\sim 52\) Ry and charge density cutoff of \(\sim 575\) Ry, and PBE exchange−correlations\(^29\) at the GGA+U approximation\(^30\) were used for the calculations, where the Hubbard correction parameter \(U\) was taken as 5.0 eV for both the Ti and O atoms.\(^31\) At a given pressure, the crystal structure of a Ti=O phase was optimized and the enthalpy of the phase was obtained. The enthalpy relative to rutile at the same pressure could then be calculated and used to determine the relative phase stability of various titania phases at different pressures.

The enthalpy of a high-pressure titania phase (baddeleyite or TiO\(_2\)-OI) was also calculated under compression with unequal pressures acting on different crystallographic orientations of the crystal (see Figure S1 and Table S1, Supporting Information (SI)). First, the crystal was hydrostatically compressed in all orientations at a pressure \(P\), generating a geometrically optimized structure and enthalpy. Then, the crystal was non-hydrostatically compressed under an axial pressure of \(P_{\text{ax}}/\text{GPa} \simeq P + S\) and two perpendicular radial pressures of \(P_{\text{rad}} \simeq P\) under the constraint that only the axial dimension was free to change. This produced another geometrically optimized structure and enthalpy. The enthalpy of the non-hydrostatic case (at \(P_{\text{ax}}/\text{GPa} = P + S\) and \(P_{\text{rad}} = P\) relative to that of the hydrostatic case (at \(P_{\text{ax}}/\text{GPa} = P + S\)) reflects the effect of the non-hydrostaticity on the enthalpy of the high-pressure phase.

**RESULTS AND DISCUSSION**

**Sample Characterization.** A photograph of the natural brookite crystals is shown in Figure 1a. The single crystals are black in color and have pseudo-hexagonal bipyramidal shapes with sizes ranging from \(\sim 2.2\) to 4.5 mm. One of the crystals is twinned with another smaller one.

The XRD pattern at ambient pressure of the brookite sample is shown in Figure 1b. Rietveld analysis confirmed that the sample is a single-phase brookite (ICSD No. 154095), since all of the diffraction peaks can be indexed as reflections from the brookite structure in the \(P\text{bc}a\) space group.\(^33\) The XRD pattern of the brookite thin foil for loading into the DAC is
allows us to investigate the influences of their hydrostaticity and elastic properties on the phase transition of natural bulk brookite.

The XRD patterns of the bulk brookite under compression in helium are shown in Figure 3a. A series of phase transitions occurred during compression of the sample. By comparing these with the calculated XRD patterns of reference TiO₂ phases (bottom of Figure 3a) and using Rietveld analyses (representative fittings are shown in Figure 4), the present phases were identified. Rietveld analyses show that small amounts of TiO₂-II phase (space group, Pbcm; ICSD No. 153282) and rutile (space group, P4/mmm, ICSD No. 916135) appeared at pressures as low as 0.8 GPa (Figure 4a). At 9.4 GPa, the majority of brookite had transformed to TiO₂-II and minor rutile. At 14.3 GPa, brookite disappeared. Meanwhile, the formed TiO₂-II and the minor rutile started to transform to baddeleyite (space group, P2₁/c; COD No. 901535535), and the transformations completed at ~23.7 GPa (Figure 3a). The baddeleyite phase remained a single phase between 23.7 and 29.7 GPa, above which it started to transform to and coexist with TiO₂-OI (space group, Pca₂₁) up to the maximum experimental pressure (59.7 GPa) (Figure 3a). The coexistence of the two phases in such a wide pressure range (~35–60 GPa; Figure 3a) may be attributed to the similarity between the two structures. 36–39 During decompression, TiO₂-OI converted back to baddeleyite (Figure 3a). When decompressed to 13.6 GPa, TiO₂-OI disappeared, while TiO₂-II and small amounts of rutile reappeared, coexisting with baddeleyite. Upon full decompression to ~0.1 GPa, baddeleyite converted almost fully to TiO₂-II coexisting with a small amount of rutile (Figure 3a). From the Rietveld analyses of all the XRD patterns in Figure 3a, the variations of the phase contents of various titania phases with pressure are depicted in Figure 5a. This diagram shows clearly the phase behavior of bulk brookite under compression in a helium medium.

Similarly, the phases present during compression and decompression in the neon pressure medium (Figure 3b–c) or methanol–ethanol mixture (Figure 3d–e) were identified using the same methods as the one used for Figure 3a. The variations of the phase contents with pressure in these two media are illustrated in Figure 5b–c, respectively. The three diagrams in Figure 5 show that the phase behaviors of bulk brookite in both helium and neon are similar at all P, indicating that their (quasi-) hydrostaticities are nearly equal and affect the phase behavior of brookite in the same way. Phase transitions of bulk brookite in the methanol–ethanol mixture (Figure 5c) are also similar to those in helium (Figure 5a) and neon (Figure 5b) in the hydrostatic pressure range (~11 GPa36). However, they deviate from those in the latter two media significantly at ~>11 GPa: in methanol–ethanol, the baddeleyite phase fully formed from brookite at ~25 GPa could remain up to ~50 GPa before transforming to TiO₂-OI (Figure 5c), while, in helium and neon, the baddeleyite also fully formed at ~25 GPa but started to transform to TiO₂-OI as low as ~30 GPa (Figure 5a,b). Moreover, during decompression, baddeleyite could also retain its single phase between ~30 and ~15 GPa in the methanol–ethanol medium (Figure 5c), while, in helium and neon, no single baddeleyite phase was observed (Figure 5a,b). These differences demonstrate that the hydrostaticity and the related elasticity of a pressure medium can modify the phase behavior of titania markedly: non-hydrostaticity favors stabilization of the
baddeleyite phase as a single phase. This phenomenon may be interpreted as a result of the solidification of the methanol–ethanol mixture, as the solidified glassy and stiff mixture can introduce lattice strains in the titania phases, impacting the transition kinetics. Because baddeleyite has a lower symmetry (monoclinic) than the other occurring phases (either orthorhombic or tetragonal), it can accommodate more lattice strains (due to less geometric constraint on the unit cell) than them (due to more geometric constraints on the unit cells). Thus, non-hydrostaticity (and related medium properties) tends to slow or even inhibit the transition from baddeleyite to TiO2-OI.

Figure 5 should be treated as non-equilibrium (i.e., kinetic) state diagrams as they deal with the behavior of a metastable phase (brookite), show the path-dependent behavior (e.g., difference between compression and decompression), and do not conform to the equilibrium phase rule. If the TiO2 system were to be in equilibrium at room temperature under compression, the maximum number of coexisting phases is P = C + 1 − F = 1 + 1 − 0 = 2. However, in our experiments, three phases occurred concurrently at some pressures (e.g., at 9.4 GPa in compression in Figure 5a). This happened because non-hydrostatic stress provides an additional degree of freedom at some of the experimental conditions, allowing the occurrence of competing phase transitions from one phase (or two phases) to the other two phases (or another phase). For instance, at 9.4 GPa in compression, brookite → TiO2-II and brookite → rutile (Figures 3a and 5a).

High-Pressure Raman Spectroscopy. High-pressure Raman measurements of the natural bulk brookite (Figure 6) were conducted to confirm the phase transitions observed by XRD (Figure 3a). As shown in Figure 6a, the Raman peaks observed at 1.1 GPa are in good agreement with the predicted vibrational bands of brookite:40 A1g (127, 155, 192, 246, 414, and 643 cm−1), B1g (136, 155, 213, 322, 414, and 504 cm−1), B2g (366, 464, and 586 cm−1), and B3g (172, 286, 547, and 449 cm−1).40,41 Among which the band at 155 cm−1 is the strongest. Upon further compression to 5.8 GPa, the Raman bands of the TiO2-II phase42−44 became obvious, while the Raman bands of brookite became weak and almost disappeared at 11.1 GPa. When the pressure reached 19.8 GPa, all TiO2-II Raman peaks disappeared, while the baddeleyite Raman peaks42−44 appeared and then became significantly broader and weaker under further compression to 45.0 GPa. From 49.3 to 59.6 GPa, a sharp Raman peak at ~295 cm−1 appeared, indicating the TiO2-OI phase, and hence, the Raman spectra consisted of contributions from both the baddeleyite and TiO2-OI phases.

Table 1. Major Chemical Composition of the Natural Brookite Sample

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<thead>
<tr>
<th>element</th>
<th>av wt percent (%)</th>
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<tr>
<td>O</td>
<td>38.64</td>
<td>65.72</td>
</tr>
<tr>
<td>Ti</td>
<td>59.23</td>
<td>33.65</td>
</tr>
<tr>
<td>Nb</td>
<td>2.13</td>
<td>0.63</td>
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<td>total</td>
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Figure 2. SEM image of the natural brookite particles produced from crushing and grinding a brookite crystal (a); chosen sample locations for the EDS analyses (b); EDS spectra from the chosen locations (c, d).
During decompression from 59.6 to 33.0 GPa (Figure 6b), the TiO$_2$-OI Raman peaks became progressively weaker while those of baddeleyite became stronger. On further decompression to 13.2 GPa, the Raman spectra were dominated by baddeleyite. When the pressure decreased to 2.9 GPa, the TiO$_2$-II Raman bands increased significantly, which were retained at 1.0 GPa. Rutile Raman bands were not observed in decompression to near the ambient pressure, as would be expected from the XRD data (Figure 5c). We interpret this as being due to the very low content of rutile. Overall, the high-pressure Raman spectroscopy data of the phase transitions of natural brookite in a methanol—ethanol $P$ medium are in good agreement with the XRD results (Figure 5c).

In the Raman measurements, laser illumination on a sample might produce heat, causing thermally activated phase transitions in brookite. However, as the applied laser power was low and the exposure time was short (see above), any heat generated could be easily dissipated into the gasket via the pressure medium and then to the DAC set. In addition, the thermally activated brookite-to-rutile transition requires very high temperature (over 800 °C).45 This temperature is unlikely to have been achieved by the Raman laser illumination in our

Figure 3. High-pressure XRD patterns of natural brookite under compression using pressure transmitting media of helium (a), neon (b and c), and methanol—ethanol (4:1 in volume; d, e). Data in panels a–c were collected at ALS-12.2.2 beamline station, and those in panels d and e at the APS-13-BMC beamline station. In panel a, prefix "D" in the pressure values indicates decompression. The calculated XRD patterns under ambient pressure are shown at the very bottom of each diagram for reference. Asterisk symbols (*) denote rutile (110) peaks.
experiments. Thus, the phase transitions observed in the HP-Raman measurements can all be attributed to the pressure effects.

**Sequences of Phase Transitions of Bulk Brookite under Compression.** Our experimental results (Figure 5) show that, under compression, bulk brookite follows the phase transition sequence of brookite → TiO$_2$-II (+minor rutile) → baddeleyite → TiO$_2$-OI, whereas during decompression it is TiO$_2$-OI → baddeleyite → TiO$_2$-II (+minor rutile). In comparison, nanobrookite transforms to baddeleyite directly.\textsuperscript{20} Such a difference may be related to the increased free energy of nanoparticles, which is size- and morphology-dependent, causing modified phase behavior and transition sequence in nanoparticles.\textsuperscript{46}

Previous experimental and theoretical works show that, under compression, bulk anatase or rutile follows the phase transition sequence of anatase/rutile → TiO$_2$-II → baddeleyite → TiO$_2$-OI → TiO$_2$-OII.\textsuperscript{18,19,47–49} In this respect, the pressure-induced phase transitions of bulk brookite resemble those of bulk anatase and bulk rutile. However, the observed temperatures for the TiO$_2$-OI phase formation are quite different, as discussed below.

Theoretical calculations predicted the presence of TiO$_2$-OI in the pressure range of $\sim$45–70 GPa (300 K),\textsuperscript{18} above $\sim$40 GPa (0 K),\textsuperscript{18} or in $\sim$20–40 GPa (0 K).\textsuperscript{30} Experimentally, TiO$_2$-OI occurred at 1300–1500 K by laser heating and in the pressure range of 28–32 GPa.\textsuperscript{18} In another experiment, TiO$_2$-OI was observed at 2200–2500 K in laser heating and in the pressure range of 35.3–36.2 GPa.\textsuperscript{18} In contrast to these, TiO$_2$-OI formed at room temperature at $\sim$33 GPa in helium/neon media (Figure 5a,b) or $\sim$50 GPa in methanol–ethanol medium (Figure 5c) in our experiments using bulk brookite as the starting material. Moreover, the formed TiO$_2$-OI was stable at $P$ up to the maximum experimental pressure of $\sim$60 GPa. The formation pressure of bulk TiO$_2$-OI in quasi-hydrostatic conditions ($\sim$33 GPa) is far less than the predicted pressure of 45–70 GPa.\textsuperscript{18} Yet, it is close to the experimentally observed pressures in the laser heating experiments of $\sim$28–32 GPa\textsuperscript{18} or $\sim$35–36 GPa.\textsuperscript{51} These indicate that using brookite as
and the starting material can alter the phase transition kinetic route so that the kinetic barrier can be reduced significantly.

Figure 5 reveals that, in the methanol–ethanol medium at high pressure, the non-hydrostaticity and the change in the medium elasticity (see below) can inhibit the baddeleyite → TiO2-OI transition (Figure 5c vs Figure 5a,b), making the transition pressure (~50 GPa) much higher than those in more hydrostatic conditions (~33 GPa). In previous laser heating experiments, no pressure medium or NaCl medium was used, which would produce less or no hydrostatic conditions. Again, this would introduce high lattice strain in baddeleyite, stabilizing baddeleyite at high pressures (see above). Consequently, much higher temperatures (>1000 °C) were needed to trigger the baddeleyite → TiO2-OI transition with laser heating in these works.

Understanding the Phase Transition Sequences of Brookite Using Computations. The phase stabilities at both ambient and high pressures of various titanias phases have been studied extensively using different kinds of computational methods, achieving different levels of success in terms of consistency with the available experimental data. It was proven that it is nontrivial to correctly model the phase stability of the titania system as a function of either or both of the temperatures and pressures. For example, to correctly reproduce the observed high-pressure phase transitions among anatase, rutile, and TiO2-II using the DFT+U method, the Hubbard parameter U should equal 5 eV. Based on this, we chose U = 5 eV in our computations using the DFT+U method. Figure 7 shows our calculated enthalpies of the concerned titania phases at different pressures. Using the calculated lattice parameters and, hence, the unit cell volumes as a function of pressure, we also derived the bulk moduli of several TiO2 phases by fitting the data to the equations-of-state (EOS) (see Figure S3). Results in Table S3 show that the DFT-derived bulk moduli agree fairly well with those derived from the experimental data (Figure S4). By contrast, the bulk moduli from a previous DFT work without the Hubbard correction deviate appreciably from the experimental results (except for TiO2-II).

Our DFT calculation results show that, at ambient temperature and pressure, ΔH(rutile→anatase) = 3.0 kJ mol\(^{-1}\) and ΔH(rutile→brookite) = 5.5 kJ mol\(^{-1}\) (Figure 7). Using the experimentally determined entropies of rutile (50.4 J mol\(^{-1}\)K\(^{-1}\)), anatase (49.8 J mol\(^{-1}\)K\(^{-1}\)) and brookite (52.6 J mol\(^{-1}\)K\(^{-1}\)), the entropy contribution to the free energy change is \(\Delta S(rutile→anatase) = 0.2 \text{ kJ mol}^{-1}\) and \(\Delta S(rutile→brookite) = -0.7 \text{ kJ mol}^{-1}\), accounting for 6.2% and 14.6% of the total free energy change (i.e., \(\Delta G = \Delta H - \Delta S\)) respectively. As the pressure increases, the entropy contribution (\(\Delta G / (\Delta U + \Delta P \Delta V - \Delta S)\)) decreases. Based on this, and considering that experimental entropies of other involving phases are unavailable, we used the enthalpies of the TiO2 phases to analyze the phase transitions of TiO2.

Figure 7 shows possible phase transitions following the enthalpy curve of brookite at 30 GPa is due to its spontaneous change to a new structure (phase) in the DFT geometric optimization.
brookite is metastable with respect to rutile and TiO2-II. Thus, brookite can transform to rutile and TiO2-II via competing reactions. This is consistent with the experimental data (Figures 3 and 5) that show the formation of TiO2-II and rutile from brookite. The experimentally formed rutile has a lower content than TiO2-II, while in Figure 7 at P < 18 GPa, rutile has a lower enthalpy and, hence, it should be more stable and have a higher content than TiO2-II. If the entropies of the involved phases are considered, \( \Delta G(\text{brookite} \rightarrow \text{rutile}) = -5.5 - 0.3 \times (51.4 - 52.6) = -5.1 \text{ kJ mol}^{-1} \) and \( \Delta G(\text{brookite} \rightarrow \text{TiO2-II}) = (5.4 - 5.5) - 0.3 \times (48.4 - 52.6) = 1.16 \text{ kJ mol}^{-1} \) at ambient pressure. This means that, around ambient pressure, the transition from brookite to rutile is even more thermodynamically favorable than brookite to TiO2-II. Thus, the difference between the theoretical expectation and the experimental data suggests that there is a faster transition kinetics from brookite to TiO2-II than to rutile due to the similarity between the TiO2-II and brookite structures, as they differ only in the stacking arrangement of similar structural layers.4,59 On compression between 13 and 20 GPa, unreacted brookite continues to transform to TiO2-II and rutile, and to baddeleyite, as the enthalpy of the latter now becomes lower than that of brookite (Figure 7). This also agrees with the experimental data in Figure 5, though experimentally the contents of rutile and TiO2-II diminish with the increase of \( P \). The fast increase of the baddeleyite content above ~15 GPa (Figure 5) suggests that there is a faster transition kinetics from brookite to baddeleyite than to TiO2-II and rutile.

Upon further compression above 20 GPa up to the maximum \( P \) of 60 GPa, TiO2-OI phase has a lower enthalpy than baddeleyite (Figure 7). Thus, the previously formed baddeleyite should transform to the TiO2-OI phase. However, experimentally, this transition was not observed until \( P \) above ~30 GPa (Figure 5) or ~50 GPa (Figure 5c). Such a hysteresis is related to the shear modulus of the pressure medium.60 The onset transition pressure deviates more from the equilibrium value in a pressure medium with a lower shear modulus (\( G \)). According to ref 61, under compression, the shear modulus of helium has a magnitude that is almost the same as that of the pressure itself (e.g., at \( P = 30 \) and 50 GPa, \( G \approx 30 \) and 50 GPa). Using the bulk modulus data of methanol measured by Zaug et al.,62 we estimated that the shear modulus of methanol is \( G \approx 14 \) and 22 GPa at \( P = 30 \) and 50 GPa, respectively (see the SI). If the elastic behaviors of helium and neon are similar and those of methanol and ethanol are similar, then we would expect the onset pressure of the baddeleyite \( \rightarrow \) TiO2-OI transition in the helium/ neon medium to be lower than that in the methanol–ethanol mixture. This deduction is consistent with the experimental results shown in Figure 5.

It is also possible that the non-hydrostaticity of a pressure medium (after its solidification) may contribute to the hysteresis in the baddeleyite \( \rightarrow \) TiO2-OI transition. Results from our DFT calculations show that the non-hydrostaticity can indeed delay the transition, making baddeleyite more stable by ~0.1–0.2 kJ/mol than at hydrostatic conditions (see Table S1). However, this effect is minor compared to the effect of shear modulus of a pressure medium (above).

Similar analyses can also be applied to the decompression process, which also shows a good agreement between the predicted phase transition sequences (Figure 7) with the experimental ones (Figure 5a,b). Experimentally, metastable TiO2-II dominated during decompression below ~15 GPa possibly because of the higher transition kinetics from baddeleyite to TiO2-II than to rutile.

The above analyses show that the calculated relative enthalpy diagram of TiO2 is consistent with observations when additional degrees of freedom (from non-hydrostatic pressure) and their influence on kinetics are considered in conjunction.

CONCLUSIONS

At room temperature and under compression up to ~60 GPa, natural brookite undergoes a series of phase transitions at attainable kinetics, driven by a decrease in the free energy. Below ~21 GPa, brookite exhibits similar transition behaviors in three kinds of pressure media, helium, neon, and a methanol–ethanol mixture: brookite (~0–12 GPa) \( \rightarrow \) TiO2-II and minor rutile (~1–21 GPa) \( \rightarrow \) baddeleyite (~12–21 GPa). A single phase of baddeleyite exists from ~21 to 33 GPa (in helium/neon) or from ~21 to 50 GPa (methanol–ethanol mixture). Above ~33 GPa (in helium/neon) or ~50 GPa (methanol–ethanol mixture) up to ~60 GPa, baddeleyite transforms to TiO2-OI. The differences in these onset pressures of the baddeleyite \( \rightarrow \) TiO2-OI transition arise from the differences in the shear moduli of the pressure media after solidification. The non-hydrostaticity of a pressure medium at high pressure makes the baddeleyite \( \rightarrow \) TiO2-OI transition energetically less favorable than at hydrostatic conditions. This work provides new knowledge of the natural brookite phase transitions at very high pressures (up to ~60 GPa), offering insights into how hydrostaticity and the elastic properties of a pressure medium can affect the phase transition behavior under compression. The obtained fundamental understanding of the brookite system may be transferable to other natural mineral systems.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsearthspacechem.8b00213.

Effect of non-hydrostaticity on enthalpy, XRD pattern of thin foil brookite, lattice parameters from representative Rietveld fitting, bulk moduli of TiO2 phases, and shear modulus of methanol at high pressure (PDF)

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