superconductor at high pressure due to the high density of states at the Fermi level. Matsuoka et al. [2] followed Feng’s work and indeed found CaLi₂ to become a superconductor at 11 GPa. However, the crystal structure of CaLi₂ under pressure is still not well established. In this work, the crystallography of CaLi₂ at high pressure has been extensively investigated by the evolutionary algorithm USPEX. We explored several new phases of CaLi₂ at pressures in range 50 - 250 GPa. The chemical bonding and the electronic structures of these high-pressure phases CaLi₂ are studied in detail. The high-pressure structures of CaLi₂ and the physical mechanisms of phase transitions will be discussed.


Keywords: ab-initio structure determination, high pressure, electronic structure

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Effect of pressure on the crystal structure of α’-Ca₃Sr₈SiO₁₄
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One of the high-temperature polymorph of Ca₃SiO₄, α’ₚhase, which is stable above 973 K [1] is known to become quenchable to room temperature with incorporation of Sr [2]. It is reported that the phase transition between α’ₚ and α’₀ₚ is displacive and α’₀ₚ is a hettotype of α’ₚ which is isosctructural with β-K₂SO₄. In order to estimate the structural distortion of the α’ₚ phase from the α’₀ₚ phase, the tolerance factor (t) for the β-K₂SO₄ type structure was defined in this study. For the purpose of studying the effect of pressure on t, the crystal structure of Ca₃Sr₈SiO₁₄ was analyzed at 2.9 GPa and 4.6 GPa. A synthesized single crystal of 60 x 60 x 30 μm was placed in a modified Merrill-Bassett type diamond anvil pressure cell [3]. The pressure medium was a 4:1 mixture of methanol and ethanol and a stainless steel (SUS301) plate was the gasket. Pressure was calibrated by the ruby fluorescence method [4]. X-ray intensity measurements were performed using the BL-10A beam line at the Photon Factory (High Energy Accelerator Research Organization, Tsukuba, Japan). Judging from the observed bond compressibilities, t decreases with pressure and the structural distortion of Ca₃Sr₈SiO₁₄ increases with pressure.

References

Keywords: high pressure, single crystals, X-ray diffraction

P10.04.21

Rietveld refinement of natural fluorapatite from ultrahigh pressure eclogite
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Fluorapatite (FAP), found within eclogite from ultrahigh pressure orogen of China, is attracted for their structural response to pressure. Zhang et al. (2005) estimated the formation conditions of the ultrahigh pressure eclogite as >940°C and >4.5 GPa. Additional stress on FAP was kept away from crashing eclogite with hammer and grinding with agate mortar during the course of sample preparation. Powder X-ray diffraction method was used since the FAP aggregates containing amount of exsolutions on a scale of micrometer. The Rietveld method was performed for structure refinement. Polynomial background and the Pseudo-Voigt function were used for simulating measured profile. K-alpha 2 component was stripped and no smoothing was made. The final structure of the FAP is derived under the agreement index GOF=1.19. The space group of FAP is P63/m, and the refined cell parameters are a=6.8849(7) Å and a=9.3842(2) Å. This leads to a cell volume V=525.08 Å³. Metaptrism twist angle φ, defined as the (001) projected angle of O1-Ca1-O2, is measured as 23.9° from the refined structural model. The X’_ionic radius is measured as 1.313 Å from tunnel Ca-F minus Ca²⁺ radius. This X’_ionic radius is less than 1.3 Å an expected dimension of closest packing model when c/a=0.734. The formula of the FAP is, on averaged, derived as: (Ca₉₀Sr₀₀Na₀₀₀₂)₉₀(Ca₅₀Na₀₀Sr₀₀₀₂)₉₀(P₃₋₈₀Na₀₀₀₂)O₁₂F₁₋₁₀. As compression on flexible one-dimension tunnel in apatite leads to structurally adjustment through an increase in angle φ, it follows that higher pressure correlates with higher angle φ. The structure of the FAP from ultrahigh pressure eclogite of China possesses a small cell volume and a large twist angle and demonstrates a good record for the ultrahigh pressure processes underwent.

Keywords: apatite, Rietveld refinement, ultrahigh pressure

P10.04.22

Pressure-induced structural change of Ca and Mg alumosilicate melts
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Viscosity and density are important physical properties to control the transportation of magma in the Earth’s interior, which are strongly correlated to microscopic structure of magma. Magmas have been considered to be densified by changing its structure with increasing pressure. We have studied the static structure of the Ca₃Al₅Si₃O₁₄ and Mg₃Al₂Si₄O₁₄ melts up to 5 GPa by X-ray diffraction analysis, in order to clarify pressure-induced structural change in topology and aluminium coordination environment. We have found the degenerated T-O peaks splitting into two peaks up to 5 GPa, indicating the coordination change of aluminium with pressure. This observation is consistent with the results of NMR studies on the Ca₁₃Al₁₅Si₁₄ glasses quenched from high-P melts, showing the increase of population of highly coordinated aluminium.