Rietveld analysis of dicalcium aluminate (Ca$_2$Al$_2$O$_5$)—A new high pressure phase with the Brownmillerite-type structure

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**ABSTRACT**

Dicalcium aluminate (Ca$_2$Al$_2$O$_5$) was prepared in a piston cylinder apparatus at 1250 °C and 2.5 GPa. The compound is orthorhombic with space group symmetry $P2_1/m$, $a = 5.2281$ Å, $b = 14.4686(2)$ Å, $c = 5.4004(1)$ Å ($Z = 4$, $D_{calc} = 3.481$ g/cm$^3$), and belongs to the brownmillerite structure family. Main building units are (1) layers of perovskite type corner connected AlO$_6$-octahedra perpendicular to [010], and (2) zweier single chains of AlO$_4$-tetrahedra running parallel [100]. The alternate stacking of the layers and sheets of single chains results in a three dimensional network in which the calcium ions are incorporated for charge compensation. The present structure is the first example for an alkaline earth aluminate with zweier single chains of tetrahedra.

**INTRODUCTION**

Since Hansen et al. (1928) studied the ternary system CaO-Al$_2$O$_3$-Fe$_2$O$_3$ the solid solution series Ca$_x$Fe$_{1-x}$Al$_x$O$_5$ ($C_xF_x$ and $C_xA_x$ using the nomenclature of cement chemistry) has been the subject of many detailed phase analytical and structural investigations (Bertaut et al. 1959; Smith 1962; Pobell and Wittmann 1965; Berggren 1971; Colville and Geller 1972). In addition to material scientific studies, brownmillerite type compounds have been recently investigated by geoscientists. For example, the defect clustering in oxygen deficient CaTiO$_3$ perovskites resulting from the substitution 2Ti$^{4+}$ + O$^{2-} \leftrightarrow 2$ Fe$^{3+}$ + with CaFeO$_{2.5}$ as a limiting case has been studied by Becerro et al. (1999). The behavior of the oxygen vacancies and their influence on the physical properties in this model system is an important source of information on the behavior of silicate perovskites, which are thought to be a major component of the Earth’s lower mantle.

According to Taylor (1997), the solid solution series Ca$_x$Fe$_{1-x}$Al$_x$O$_5$ exists at ordinary pressures in the compositional range between $x = 0$ and $x = 0.7$. The phases belong to two different space group types. The Fe rich compositions up to $x = 0.29$ adopt space group $Pnma$, whereas the samples beyond $x = 0.29$ crystallize in space group $I2m$. The pure aluminum end-member C$_2$A ($x = 1.0$) first described by Aggarwal et al. (1972) can be prepared at high pressures only, but it can be readily preserved at ambient pressure conditions. The crystals obtained by Aggarwal and co-workers were too small for X-ray single crystal investigations. However, selected-area electron diffraction indicated that the material has an orthorhombic body centred Bravais lattice with $a = 5.24$ Å, $b = 14.40$ Å, and $c = 5.44$ Å. Structural analyses have not been performed on this high pressure phase and therefore, we decided to re-investigate this compound as part of our program on brownmillerite type material.

**EXPERIMENTAL METHODS**

Sample preparation

CaCO$_3$ (Merck, 99%) and Al$_2$O$_3$ (Aldrich, 99.8%) corresponding to the molar ratio 2:1 were used as starting materials. The reagents were carefully mixed in an agate mortar under acetone and pressed into discs (10 mm diameter and 1 mm thickness) under a pressure of $6 \times 10^8$ Pa. The pellets confined in a platinum crucible were precalcinated at 1000 °C in air using a resistance heated furnace and subsequently sintered twice at 1250 °C for 24 h with intermediate grindings before final quenching in air.

Preliminary X-ray powder diffraction of the material obtained from the solid state reactions showed only two phases to be present: Ca$_2$Al$_2$O$_5$ (ICDD entry 38-1429) and Ca$_{12}$Al$_{14}$O$_{33}$ (ICDD entry 09-413).

The sintered precursor material was re-ground and used for high pressure experiments which were performed at the Bayerisches Geoinstitut choosing the same temperature and pressure conditions as described in Aggarwal et al. (1972). The experiments were accomplished in a 1/2 inch piston cylinder apparatus using talc pyrex cells with a tapered graphite heater. The samples were encapsulated in 1 cm long and 5 mm diameter welded Pt capsules. The experiments were performed at 2.5 GPa and 1250 °C for 2 h. Pressure was calibrated against the quartz-coesite and kyanite-sillimanite transitions as well as the melting point of diopside. A friction correction of 18% was applied to the nominal pressure on the basis of these calibration data. Temperature was measured with a Pt$_{10}$Rh$_{90}$-Pt thermocouple and controlled with a Eurotherm 818 controller. Temperatures are considered to be accurate within ±10 °C with a temperature gradient of about 5 °C from the top to the middle.
of the capsule (Shaw and Fliervoet, unpublished data). The experiments were carried out using the hot-piston out technique. The samples were quenched isobarically by turning off the power of the furnace. Quench rates were of the order of 75 °C/s.

**Powder diffraction data analysis**

Due to the small amounts of material obtained from the high pressure experiments, the data collection for the structure analysis was performed on a Stoe STADI-P diffractometer in flat plate transmission geometry. The diffractometer is equipped with a curved Ge monochromator in the primary beam yielding a strictly monochromatic CuKα-radiation. The sample was prepared between two thin foils and fixed with silicone grease (Baysilone). The effective value for the product μt (μ = linear absorption coefficient; t = sample thickness) used for the absorption correction was determined experimentally from the intensity ratios at θ = 0° with and without the sample. Details of the data collection, crystallographic data and definitions are given in Table 1.

A first inspection of the X-ray diffraction pattern revealed that all peaks could be indexed on the basis of an orthorhombic cell close to the one reported by Aggarwal et al. (1972). There was no indication of additional reflections from the starting material. For the Rietveld analysis the background was set by hand, because it could not be described by a simple function because of its irregular shape due to the scattering of the grease used for fixing the sample. Background values were determined by linear interpolation between consecutive breakpoints in the pattern. Intensities within eight times the full width at half maximum of a peak were considered to contribute to the central reflexion. Peaks below 2θ = 15° were corrected for asymmetry effects after Rietveld (1969). The pseudo-Voigt function was chosen for the simulation of the peak shape, with two parameters defining the Lorentzian and Gaussian character of the peaks as a function of 2θ. The angular variation of the line width was accounted for by using the Cagliotti function (Cagliotti et al. 1958). The powder diffraction data for Ca₂Al₂O₅ including calculated (based on the unit cell parameters of the Rietveld analysis), observed peak positions, and relative background corrected intensities are listed in Table 2. The figures of merit

### Table 1. Experimental conditions and crystallographic data

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**Notes:**

\[ R_{wp} = \left( \frac{1}{n} \sum_{i=1}^{n} \left( \frac{I_i - \mu_i}{\sigma_i} \right)^2 \right)^{1/2} \]

where \( I_i \) and \( \mu_i \) are peak intensities calculated with and without the sample, respectively; \( \sigma_i \) is the standard deviation of the peak intensity; \( p = 1 - \frac{\sum y_i}{\sum y_i} \)

\[ R_p = \frac{1}{n} \sum_{i=1}^{n} \left( \frac{I_i - \mu_i}{\sigma_i} \right)^2 \]

\[ R_1 = \frac{1}{n} \sum_{i=1}^{n} \left( \frac{y_i - y_i^c}{y_i^c} \right)^2 \]

\( y_i \) and \( y_i^c \) are observed and calculated profile intensities; \( p = 1 - \frac{\sum y_i}{\sum y_i} \); \( C \) = Scaling factor; \( w = \text{weight} = 1/y_i \)

Standard deviations for the lattice constants are given in parentheses.
STRUCTURE ANALYSIS AND REFINEMENT

The structure analysis as well as distance and angle calculations were performed with a modified version of the PC-Rietveld plus package (Fischer et al. 1993) adopted for transmission geometry. X-ray scattering factors in their respective valence states were taken from the *International Tables for X-ray crystallography Vol. IV* (Ibers and Hamilton 1974) and the values for O^2– from Hovestreydt (1983). For the drawing of the structural details, the computer programs ATOMS (Dowty 1995) and ORTEP-3 (Farrugia 1997) were used. Starting values for the fractional atomic coordinates were taken from Colville and Geller (1972) and transformed to the I2mb space group. This non-standard setting of space group Ima2 (no. 46) was selected to compare the coordinates between the body centered and the primitive brownmillerite structure type (Pnma setting) directly, without any transformation of coordinates.

The refinement was done in consecutive steps with the atomic coordinates and the thermal parameters held fixed in the initial calculations. They were allowed to vary after the scale factor, zero point, peak shape parameters, and lattice constants were close to convergence to their optimum values. For the final calculations a flat plate absorption correction was included. The graphical comparison between the observed and calculated powder pattern is given in Figure 1. The refined fractional atomic coordinates are listed in Table 3 and selected interatomic distances and angles in Table 4.

The two main building units of dicalcium aluminate are: (1) layers of perovskite type corner connected AlO_6-octahedra perpendicular to [010] and (2) zweier single chains of AlO_4-tetrahedra running parallel [100]. The alternate stacking of the layers and sheets of single chains results in a three dimensional network. Charge balance is achieved by calcium ions located in voids of the framework (compare Fig. 2).

The average value of the O-Al-O angle is very close to the ideal value of 109.4°, whereas the individual O-Al-O angles vary between 104° and 125°, respectively. This spread in the angles indicates a significant distortion of the tetrahedra and can be expressed numerically with the quadratic elongation \( l = 1.014 \) and the angle variance \( \sigma^2 = 62.3 \) as defined by Robinson et al. (1972). The individual octahedra within the perovskite type sheets are significantly less distorted \( l = 1.008; \sigma^2 = 8.5 \). They are tilted against each other by rotations of ±5° about [100] and ±1° about [010]. Both the tetrahedra and octahedra are primarily distorted along the crystallographic b axis. The calcium cations are coordinated by eight neighbors forming distorted bicapped trigonal prisms.
Figure 2. Projections of the whole structure of Ca₂Al₂O₅ (a) parallel to [100] and (b) parallel [010]. The large spheres in the interstitial sites = Ca atoms; smaller spheres = O anions. Light shading = tetrahedra. Dark shading = octahedra.

Comparison with other structures

A comparison of the main structural data of the end-members of the solid solution series C₂F-C₂A is given in Table 5. The trends in the mean cation-oxygen distances of the tetrahedra and octahedra are consistent with the substitution of Fe³⁺ by the smaller Al³⁺. Furthermore, the Ca-O distances are longer for dicalcium ferrite. The intra-chain Me⁵⁺-O-Me⁵⁺ angles are definitely larger in C₂A, indicating an increased stretching of the chains in the aluminate.

The value of 125° for an tetrahedral O-Al-O angle is a conspicuous feature of this structure, being close to the upper limit of observed O-Al-O angles in alkali- and alkaline earth-aluminates (compare Fig. 3). As is shown in Figure 4 this angle occurs at the O₁ ligands connecting the octahedral layers and the tetrahedral chains. A shortening of the O₁-O₁ edge of the tetrahedron makes the angle smaller, yielding a more regular tetrahedral geometry, but stretches the Al₁-O₁ bond in the octahedron. Distance least squares calculations using the program DLS-76 (Baerlocher et al. 1977) with prescribed distances for all cations and anions result in atomic positions with an O₁-Al₂-O₁ angle of 130°. Forcing the tetrahedron to assume a regular geometry by assigning a high weight to the O₁-O₁ bond causes a lengthening of the Al₁-O₁ bond from 2.05 Å in the unconstrained DLS-refinement (2.09 Å observed) to a crystal chemical unrealistic value of 2.23 Å in the DLS-refinement with essentially fixed tetrahedral geometry.

Likewise, the Ca ions are moved toward Al₁ yielding a rather short Al-Ca distance of 2.94 Å (Al-Ca distances for all the structural characterized calcium aluminates begin at about 3.10 Å). The relatively high DLS reliability index of 0.044 for the configuration resulting from the unconstrained refinement (essentially representing the actually observed structure) shows that
and Saalfeld 1968) and SrAl$_2$O$_4$ (Lindop et al. 1975) as well as Sr$_3$Al$_5$O$_{12}$ (Nadezhina et al. 1976). Furthermore, dicalcium aluminate is the first example of a pure aluminate with zweier single chains of corner connected tetrahedra.

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