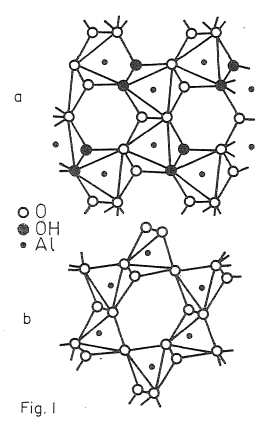
05.1-38 ON THE ORTHO-CLINO PHASE TRANSITION OF FERROSILITE(FeSiO₃) AND Fe-RICH PYROXENES. By <u>S. Sueno</u> and M. Kimata, Department of Geoscience, The University of Tsukuba, Ibaraki, 305, Japan.

Temperature induced orthorhombic(Pbca)-monoclinic(C2/c) pyroxene phase transition was studied using high-temperature precession camera method. The starting materials were synthetic ortho-pyroxenes with the composi-tion of (Fs), (Wo.02Fs.98), (Wo.05Fs.95), [Wo.05(Fs.75 En.25).95], (Fs.9En.1) and (Fs.8En.2) where Fs=FeSiO3, Wo=CaSiO3 and En=MgSiO3. Ortho-pyroxene with the Fs composition showed reversible and topotaxial transformation to the twined clino-pyroxene at 1020° C with the orientation (a^{*}_{ortho}//a^{*}_{clino}, cortho//c_{clino}). Wo free pyroxenes transformed to clino-phase topotaxially with the orientation same as those on Fs, but they did not transform reversibly to the ortho-phase on cooling. The pyroxenes with Wo composition transformed rapidly to twined clino-phase but the orientations between both phases were quite haphazard for each run. Smyth has suggested that on the coherent pyroxene transition M1 site in ortho-phase becomes M2 site in clinophase and M2 site in ortho-phase becomes M1 site in clino-phase(Smyth, Amer. Mineral.(1974) 59, 345). The results of the present study may support his suggestion as the coherent pyroxene transitions were observed only on pure Mg and Fe end-member pyroxenes in which both M-sites are occupied only by Mg and Fe. In ortho-pyroxene, Ca ion is in M2 site and it could not be entered to ML site in clino-phase because of its large ionic radius. This makes the coherent transition impossible and bring: about the "nucleus and growth type" transition. Whereas smaller Mg ion is in Ml site in orthophase and it may be allowed to enter M2 site in clino-phase, but may result in some distortion in clinophase structure which prevents the reverse transition to ortho-pyroxene on the cooling stage.

 $05.1\mathcal{39}$ THERMAL BEHAVIOUR OF MUSCOVITE - OD STRUCTURAL ASPECTS. By K. Dornberger-Schiff and K.-O. Backhaus, Central Institute of Physical Chemistry, Academy of Sciences of the G.D.R., Berlin-Adlershof, German Democratic Republic

Muscovite crystals are known to undergo irreversible changes when heated. These changes differ qualitatively when heated up to temperatures T <700° and T>700°. In order to elucidate the changes in structure, Weissenberg diagrams of heat-treated muscovite single crystals $2M_1$, quenched after heating, have been taken and discussed: <u>higher temperature treated (HTT) and lower tem-</u> perature treated (ITT) compared with 700°. (hkl) reflections (orthonexagonal) with k≠3n are diffuse (direction c^T) for LTT and HTT crystals, (hkl) with k=3n are different from the untreated crystals only for the HTT crystals.

Furthermore, IR investigations showed that no H atoms are present in the HTT crystals, indicating the loss of every second OH group together with the H atom of the other OH group, in agreement with earlier results (Peuker, private communication). The results of structure determinations of LTT and/or HTT crystals, carried out also with the help of OD theory, are to be presented. Fig. 1 Tentative structure of a) octahedral (idealized) sheet before, b) after dehydroxylation (5-coordinated).



05.1-40 RIGID-BODY ANALYSIS OF DIAMANTANE AT 7 K. By E.H.M. Evans, R. Hine and J.P.G. Richards, Department of Physics, University College, P.O. Box 78, Cardiff CF1 1XL, U.K.

Diamantane ($C_{14}H_{20}$) is a cage-like molecule which at room temperature crystallises in Pa3. It has been shown by Raman spectroscopy (Jenkins and Bates, J. Phys. C: Solid State Physics (1979) 12, 1003) that a phase transition occurs at 35 K involving a change of space group. Neutron Bragg intensities have been collected at 7 K, and the results of a rigid-body analysis of these will be presented. Preliminary results indicate no change of space group and no substantial recorientation about the three-fold axis.