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Jahn Teller Transition Phase in Cu_{0.5}TiO(PO₄) Oxyphosphate Pierre Gravereau^a, Saïd Benmokhtar^b, Jean-Pierre Chaminade^a, Abdelaziz El Jazouli^b, Eric Lebraud^a, Stanislas Péchev^a, ^aICMCB-CNRS, University of Bordeaux, Pessac, France. ^bLCMS, Faculté des Sciences Ben M'Sik, Casablanca, Maroc. E-mail: graver@icmcbbordeaux.cnrs.fr

In our research interest on titanium oxyphosphates, several compounds $M_{0.5}^{II}TiO(PO_4)$ have been prepared with $M^{II} = Mg$, Fe, Co, Ni, Cu, Zn. They all belong to the monoclinic P21/c structural type determined "ab initio" for Ni_{0.5}TiO(PO₄) from X-ray powder diffraction[1]. Thermostructural study of Cu_{0.5}TiO(PO₄) shows a phase transition $\alpha \rightarrow \beta$ (~800°C during heating, ~400°C during cooling).

Single crystals of the α phase have been obtained. Its cell presents a Jahn Teller deformation with Cu-O elongation in the (a,c) plane: a=7.5612(4); b=7.0919(4); c=7.4874(4)Å; $\beta=122.25(6)^{\circ}$ V=339.55(6)Å³. The single crystal structure confirms and states more precisely the previous powder model ($R_1=0.023$; $wR_2=0.063$).

The β phase has been obtained stabilised at room temperature in a powder mixture $\alpha(38\%)+\beta(62\%)$ without impurities. X-ray diffraction pattern of this powder can be indexed in a like α -type P2₁/c monoclinic cell : a=7.1081(10) ; b=7.7384(12) ; c=7.3013(10) Å ; $\beta{=}119.28(1)^\circ$; V=350.3(1)Å^3. As the $\alpha{-}structure~is~not~a~good$ starting model for Rietveld refinement (divergence), an "ab-initio" structure determination has been done. The refined structure corresponds to a "rocking" of the Jahn Teller elongation from the (a,c) plane to the **b** direction.

[1] Gravereau P., Chaminade J.P., Manoun B., Krimi S., El Jazouli A., Powder Diffr., 1999, 14, n.1, 10.

Keywords: oxyphosphates, phase transition, Jahn Teller effect

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A Temperature-Induced Phase Transition in Barbituric Acid Dihydrate

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In recent years the subject of phase transitions has become more and more popular for scientific investigation. Indeed virtually the whole of the January 2005 issue of Acta Crystallographica Section A: Foundations of Crystallography was devoted to the topic. By a combination of careful experimentation and some serendipitous good fortune we have determined a phase transition at low temperature in barbituric acid dihydrate.

The structure of barbituric acid dihydrate appears twice in the literature: an X-ray study [1] and a neutron study [2]. Both sets of experiments were carried out at room temperature and both report the crystal system and space group as orthorhombic Pnma. We have found that this only holds true at temperatures above ~220K. Below ~200K the structure is non-merohedrally twinned monoclinic $P2_1/n$, and at intermediate temperatures it appears to be a *mélange* of sorts of both crystal systems.

[1] Jeffrey G.A., Ghose S., Warwicker J.O., Acta Cryst., 1961, 14, 881. [2] Al-Karaghouli A.R., Abdul-Wahab B., Ajaj E., Al-Asaff S., Acta Cryst., 1977, B33 1655

Keywords: organic phase transitions, variable-temperature study, single-crystal structure determination

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An Order-Disorder Phase Transition in [Ag(bipy)NO₃]_n <u>Weenawan Somphon^a</u>, Kenneth J. Haller^a, A. David Rae^b, ^aSchool of Chemistry, Institute of Science, Suranaree Univ. of Tech., Nakhon Ratchasima 30000, Thailand. ^bResearch School of Chemistry, The Australian Univ., Canberra, ACT, 0200, Australia. E-mail: weenawanus@yahoo.com

The room temperature *Fddd* form of Ag(bipy)NO₃ contains a disordered NO3 ion and shows previously unreported planes of diffuse scattering indexed as $3k \pm l = 4n$ for the 296 K, a 12.8424(2), b 9.9429(1), c 34.4621(4) Å. Synthetic precession photographs showed the symmetry was Fddd at 160 K and above but a loss of systematic absences indicated F12/d1 (i.e. C2/c) at 150 K and below. The 100 K structure, C2/c, a 12.751(1), b 9.860(1), c 18.379(2) Å, β 109.98(1) ° has twin components related by a rotation around c*. The cell for F12/d1 has a' = a, b' = b, c' 34.547(2) Å, $\beta' 89.68(1)$ °. Refinement gave a 0.754(1):0.246 twin with no disorder and R(F) = 0.022 for 2294 obsd. rflns. Along a chain alternate, Ag atoms, (3b±c)/4 apart, are displaced 0.221(1) Å in opposite directions perpendicular to the chain. Chains are cross linked by Ag-Ag contacts of 2.958(1) Å and Ag-O contacts of 2.749(2) and 2.747(2) Å. The chains zig-zag so that Ag atoms avoid closer contact with the NO₃. The 200 K structure, Fddd, a 12.823(1), b 9.937(1), c 34.450(1) Å was refined as a 1:1 disorder of all atoms initiated by disordering the 100 K structure. Constrained refinement gave R(F) = 0.028 for 1004 obsd. rflns. The Ag displacements reduced to $\pm 0.123(3)$ Å. The alternative orientation of the nitrate gave three Ag-O contact distances indicating an intermediate step for a change of local ordering. The diffuse scattering indicates that mistakes in NO3 positions cause a localized straightening of the adjacent chains that moves substantial amounts of these chains along their lengths. In contrast, the actual position of a nitrate only affects the closest Ag atoms.

Keywords: polymorphic structure, phase transition, orderdisorder transition

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Metal-Insulator Transition in Hollandite Vanadate, K₂V₈O₁₆ Yutaka Ueda, N. Kouno, M. Isobe, T. Yamauchi, H. Ueda, MDCL, Institute for Solid State Physics, University of Tokyo, Kashiwa, Japan. E-mail: yueda@issp.u-tokyo.ac.jp

Vanadium oxides have attracted considerable interest in the novel properties such as metal-insulator transition [1], charge order transition [1], pressure-induced superconductivity [2] and so on. Some vanadium oxides with the Hollandite-type structure were also reported. Among them, $Bi_xV_8O_{16}$ is famous in its metal-insulator transition [3]. The structure of Hollandite vanadate $A_2V_8O_{16}$ consists of V₈O₁₆-framework and A-cation located at the tunnel sites of V₈O₁₆framework. The V_8O_{16} -framework is constructed from the double chains formed by sharing the edges of VO_6 octahedra. We successfully synthesized $K_2V_8O_{16}$ by a solid state reaction under 4 GPa at 1473 K for 30 minutes. Small crystals were also found in the sintered sample. K₂V₈O₁₆ shows a metal-insulator transition with the jump of reisitivity about three orders around 170 K, accompanied by the structure change from tetragonal to orthorhombic and the large reduction of magnetic susceptibility. Electron diffraction study reveals a superlattice of $\sqrt{2} \times \sqrt{2} \times 2$ in the low-temperature insulator phase. Taking these results into consideration, we propose a charge ordered model for the low-temperature insulator phase in which $V^{\bar{4} +} - V^{4+}$ spin singlet pairs are formed. We also first synthesized Rb₂V₈O₁₆ and discovered a metal-insulator transition around at 150 K.

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Re-examination of Phase Transitions in KNbO3

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As it is known, the physical properties of KNbO₃ are similar to the properties of BaTiO₃. Though the nano-size effects in BaTiO₃ are researched rather well, such studies of KNbO3 have never been held