THE ATOMIC STRUCTURE OF IONIC CONDUCTORS Li₃Sc₂(PO₄)₃ AND 07.3-4

Li₃Fe₂(PO₄)₃ AT 293 AND 593 K. By B.A.Maksimov, E.A.Genkina, L.A.Muradyan and <u>V.I.Simonov</u>, Institute of Crystallography, <u>USSR Acad.Sci.</u>, Moscow, USSR.

At 600 K the conductivity of $Li_3Sc_2(PO_4)_3$ and

Li,Fe,(PO), crystals, obtained by solid-phase synthesis reaches (2-5)·10⁻²Ω⁻¹cm⁻¹(M.Pintard-Screpel, F.d'Yvoire, E.Bretley, Solid State Chem.Proceed.(1982), 215-218)(E.A.Genkina, L. N.Demyanets et al., JETP letters(1983),v.38, N°5, p.257-259).According to electro-chemical measurements these compounds become fast ionic conductors at T~540 K. The X-ray intensity data were collected using MoK_L radiation on a CAD-4F diffractometer equipped with a graphite monochromator(step scanning method, W-techni-que, sin $\mathcal{O}\chi = 1.0 \text{ A}^{-1}$). measurements these compounds become fast ionic

At room temperature Li3Sc2(PO4)3 crystals are monoclinic, Fedorov group P2₁/n, a=8.851(1), b=12.271(2), c=8.800(1)Å, χ =90.02°, χ =953Å³, z = 4. In the fast-ionic state, at T=593 K, the symmetry is changed to orthorhombic (Fedo-rov group Pcan D¹⁴_{2h}, a=8.801(1), b=12.393(2), c=8.822(2)Å, χ = 962Å³, z=4. The Li₃Fe₂(P0₄)₃ unit cell have the following dimensions: a = 8.545(2), b = 12.003(3), c = 8.604(2) Å at T= =293 K; and a = 8.592(1), b = 12.129(2), c = = 8.637(2) Å at T = 593 K.

The basis of the structure of lithiumscandium phosphate and lithium-iron phosphate at room temperature (293 K) and at temperature 593 K, exceeding the phase transition point is three dimensional framework , formed by M^{2+} -octahedra (M^{2+} - Sc, Fe) and P-tetrahedra, connected by common vertices. The cavities in the framework are occupied by single-char-. ged lithium atoms. The analysis of the change in the struc-

ture parameters accompanying the considered phase transition to the fast-ionic state showed that $\{[\mathbb{M}_2(\mathbb{PO}_4)_3]^{3^-}\}$ frameworks are virtually identical both below and above the phase tran-sition point. After the phase transition the displacement of \mathbb{M}^{2^+} , P and O atoms does not exceed 0.2 Å. There are significant differen-ces only in the location of Li atoms at 293 and 593 K. At T = 293 K the 12 lithium atoms of a unit cell fully occupy three four-fold complexes. At T = 593 K in $\text{Li}_{3}M_{2}(\text{PO}_{4})_{3}$ struccomplexes. At T = 593 K in $\text{Li}_{3}\text{M}_{2}(\text{PO}_{4})_{3}$ structures Li atoms occupy also three, although 8-fold complexes: only one 8-fold site is fully occupied, the other two are occupied by Li with the probability of 1/4. The Li, atom is positioned inside a distorted tetrahedra, Li-0 distance being 1.91(2)-2.08(2) Å. Coordinational polyhedra of Li and Li atoms have five vertices, the distances being: $\text{Li}_{2}-0 = 2.15(2) - 2.43(2) Å$, $\text{Li}_{3}-0 = 1.99(2) - 2.60(2) Å$. The Li polyhedra are connected by common edges and vertices. common edges and vertices.

The mutual arrangement of Li atoms at 593 K indicates a highly probable and essential anisotropy of the conductivity.

07.3–5 PHASE TRANSITIONS IN SUPERIONIC

CONDUCTOR CSHSO4 STUDY BY NEUTRON DIFFRACTION

AND INELASTIC SCATTERING. By A.V. Belushkin, <u>I. Natkaniec</u>, N.M. Plakida, L.A. Shuvalov*, J. Wasicki, Joint Institute for Nuclear Rese-arch, 141980 Dubna, USSR, * Institute of Crystallography, 117333 Moscow, USSR.

Simultaneous measurements of neutron diffraction (ND) and inelastic incoherent neutron scattering (IINS) on a powder sample of CsHSO,

in the temperature range of 5+423 K have been performed at the IBR-2 pulsed reactor. Three phase transitions have been detected by ND phase transitions have been detected by ND when a sample was heated from the room tempe-rature up to 423 K. As it is seen from the LINS spectra the dynamics of protons signifi-cantly changes at the first transition near 325 K and at the superionic transition at 414 K as well. The phase upper 325 K has a metastable character and car her cardia compo metastable character and can be easily supercooled. Proton jumps in the superionic phase at 423 K should have a frequency lower than

10⁻¹¹sec, because we have not observe quasielastic broadening within our 8% resolution The broadening within our 3% resolution at the 4,9 meV elastic line. Using vibrational frequency obtained from IINS we estimate the diffusion coefficient for protons: $D = 2,7 \cdot 10^{-6} \text{cm}^2/\text{sec.}$ Necessary resolution to observe such a diffusion is of the order of 0,02 meV.

07.4-1 THE ORGANIC CONDUCTORS $(TMTSF)_2TaF_6$ AND $(TMTTF)_2TaF_6$. By N. Thorup and G. Rindorf, Chemistry Dept. B, Technical University of Den-mark, DK-2800 Lyngby, Denmark.

During the past few years extensive studies have been made on two series of molecular metals, namely (TMTSF)₂X and (TMTTF)₂X (for recent re-sults, see J. Physique (1983), Colloque C3, pp. 767-1136). TMTSF is tetramethyltetraselenaful-valene, TMTTF is its sulfur analogue tetrame-thyltetrathiafulvalene, and X is an anion, e.g. BF₄, ClO₄, ReO₄, PF₆ or TaF₆. These materials exhibit unusual physical features such as low-dimensional metallic transport properties, and superconductivity has been demonstrated in sev-eral cases including (TMTSF)₂TaF₆ at temperaeral cases including (TMTSF) 2 TaF6 at temperatures below 1 K and pressures above 10 kbar.

The crystals are all isostructural at room temperature and they consist of TMTSF cation-radicals stacked along the *a* axis. The stacks are connected (via Se-Se or S-S contacts) to sheets separated by anion layers parallel to the *ab* plane (see e.g. Thorup et al. Acta Cryst. (1981) <u>B37</u>, 1236).

The present work focuses on the structural details of the title compounds and, in particular, the differences between them. In both cases, however, the octahedral anions seem to be dis-ordered in a very similar fashion. Each octahedron can take one of two orientations, which are very close. The triclinic cell dimensions at 295 K are:

b а С α ß γ TMTSF salt 7.291 7.728 13.936 82.86 85.33 71.66 TMTTF salt 7.182 7.646 13.555 81.44 83.47 73.86