We found that the chiral cyanoethyl group bonded to the cobalt atom in a cobaloxime complex crystal was racemized by X-ray exposure without degradation of the crystallinity (Ohashi & Sasada, Nature(London), 1977, 267, 142-144). The mode of the racemization and the reaction rate explained quantitatively by the reaction cavity, which was defined as a void space around the reactive group (Ohashi, Acc. Chem. Res., 1988, 21, 268-274). If the chiral group was replaced with the bulkier methoxycarbonylethyl group, the racemization occurred in two stages, the conformational and configurational changes (Kurihara, Uchida, Ohashi, Sasada & Ohgo, J. Am. Chem. Soc., 1984, 106, 5718-5724). Recently the bulkiest dimethoxycarbonylethyl group was used as a chiral group. The structural change was so fast that the three-dimensional intensity data was obtained by the new type of diffractometer using the imaging plate. The analyzed structures at several intermediate stages revealed complicated disordered structures. Subtracting the original or final structures from the disordered ones, the unstable intermediate structure can be observed (Sakai, Ohashi, Arai & Ohgo, Mol. Cryst. Liq. Cryst., 1992, 219, 149-152). This indicates that if the reactive crystals were analyzed by X-rays more quickly, the dynamical process of reactions should be made clear on the basis of the structure change.

MS-06.01.08 CONFIGURATION AND NONLINEAR OPTICAL PROPERTIES OF DMIT DERIVATIVES. Q. Fang\*, Institute of Crystal Materials, Shandong Univ. 250100, China; Q.Zheng, Chem.Dept., Jiangxi Medical college, 330006, China.

Although it is well known that DMIT play an important part in the molecular conductor field, nonlinear optical phenomena of second harmonic generation (SHG) for DMIT derivatives has not been observed until we synthesized a-BNPT-DTT, B-BNPT-DTT and BNPT-DTO recently. The syntheses and the products' structural parameters are as follows:

triclinic Pl orthorhombic system orthorhombic P212121 P212121 group group P1 a(Å),a(\*) 12.18,96.5 b(Å),β(\*) 14.25,96.9 c(Å),γ(\*) 7.79, 72.3 Y,Z,De 1274,2,1.38 10.13 10.08 19.35 9.66 9.93 2005,4,1.76 1936,4,1.77 final R 0.038 0.027

final R 0.069 0.038 0.027
By using a YAG:Nd laser at 1064nm, only BNPT-DTO has SHG at 532nm, which is as strong as that produced by KDP. α-BNPT-DTT belongs to Pl, and of course has no SHG effect. β-BNPT-DTT and BNPT-DTO belong to same space group and possess nearly the same cell parameters. What make the differences? The charge distribution does not change much by substituting 0=< for S=< for all, respective bond lengths (except 0=C,S=C) and the respective NMR shifts of protons in the phenyl do not change. BNPT-DTO is in a stretching configuration whose dipole moment is considerably higher than that of β-BNPT-DTT. So it is the configurations that make the differences.

MS-06.01.09 SUPRAMOLECULAR ARCHITECTURES OF METALLOPOR PHYRINS IN CRYSTALLINE SOLIDS.

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Research efforts directed at the construction of microporous solids based on functionalized metalloporphyrins as building blocks uncovered a series of new materials with attractive structural features. This approach involves substitution of the rigid porphyrin core with polarized aryl groups, the polar functions being used to induce coordination between the molecular fragments, and thus form crystalline polymers with an extremely high degree of two- and three-dimensional cross-linking. Such structure-enforced supramolecular assemblies are then capable of incorporating smaller molecular entities into the lattice. Controlled variation of the microstructure in the resulting solids, and consequently of their structure-dependent properties, is effected by changing the type (and shape) of the sensor groups and their disposition in the porphyrin moiety.

Crystallographic investigations have confirmed that the intermolecular arrangement of the porphyrin materials is dominated by directional hydrogen bonds and other (e.g., Cl···Cl, and metal-nucleophile) interactions, as well as by the molecular shape, yielding continuous networks of strongly coordinated entities which resemble to a large extent the structural rigidity of a zeolite. Guest components can be absorbed in these solids in distictly defined sites of the porous lattice. The polymeric molecular organization, along with the rich chemistry associated with the porphyrin core, make these materials useful in a wide range of applications, including matrix isolation, molecular recognition and chemical separation, and mediation of ligand transport and controlled release. Several examples will be discussed to illustrate the potential for the design of crystalline lattices with different degrees of cross-linking, rigidity and polarity, and for the formation of polar tubular structures capable of aligning organic dipolar guest molecules in the crystal bulk.

MS-06.01.10 STRUCTURE AND PROPERTIES OF AN ALKENYL LIQUID CRYSTALLINE COMPOUND. By S. Gupta, A. Nath, S.Paul. Department of Physics, North Bengal University. Siliguri - 734430. India and H. Schenk, Crystallography Laboratory, University Of Amsterdam, 1018WV Amsterdam, The Netherlands.

We have undertaken the structural study of 4 (3"-pentenyl) 4" (cyano) 1, 1' bicyclohexane. Compounds with alkenyl end chains belong to different structural classes. Sheir common features are low viscosities as well as low optical anisotropies in the liquid crystalline state. The alkenyl compounds differ from each other in many of their properties with respect to systematic changes made in the position of the alkenyl double bond. The molecular arrangement in the crystalline state is one of the factors which sometimes predetermines the properties in the liquid crystalline state. The compound studied here has the following structure:

The transition temperatures of different phases as determind by X-ray diffraction and texture studies are given bellow :

We report here the crystal structure of the compound and try to explain the mesomorphic organization in relation to its solid state packing and molecular interactions. Intensity data were collected in a CAD-4 diffractometer. The compound crystallised in the monoclinic space group  $P2_{\star}/n$  with unit cell parameters