(c) hydrogens with H(D)...S > 3.1  $^{\circ}$  EH(D)...0 > 2.70  $^{\circ}$ J, far outside the usual hydrogen bond range.

Vibrational spectroscopic investigations isotopically dilute (H and D) single crystals, measured at 295 K and 75 K, showed uncoupled 0-H(D) stretching frequencies in the range  $\tilde{Y} = 2345-2665$  (3145-3620) Accordina to the Badger-Bauer rule these frequencies would correspond to hydrogen bond energies of  $5-40 \text{ kJ} \text{ mol}^{-1}$ . Reasonable assignments of the observed frequencies to the individual hydrogen bonds could be made. For the approximately linear O-H(D)...S bonds with H(D)...s = 2.35 Å (0...s = 3.3 Å)uncoupled 0-H(D) **¥** = 2400-2520 frequencies of (3220-3410) cm<sup>-1</sup> were observed, the specific values depending significantly on the chemical nature of the S acceptor atoms. Despite large H(D)...S distances (2.71-2.85 Å) the bifurcated 0-H(D)...S,S bonds show stretching frequencies which are similar to those of linear 0-H(D)...S bonds with H(D)...S = 2.35  $^{\circ}$ A. The type (c) hydrogen atoms observed in Na<sub>3</sub>PS<sub>4</sub>.8H<sub>2</sub>O and Na<sub>3</sub>AsS<sub>4</sub>.8D<sub>2</sub>O exhibit O-H(D) frequencies of  $\tilde{y}$  = 2665 (3620) cm<sup>-1</sup> which indicate some crystal field interactions with the surrounding atoms.

04.6-03 X-RAY AND NEUTRON DIFFRACTION STUDIES OF THE HYDROGEN BONDING IN HYDROXYBENZOPHENONES. By E.O. Schlemper and M.S. Hussain, Department of Chemistry, and Fred Ross, Research Reactor Facility, University of Missouri, Columbia, Missouri, U.S.A. In order to examine hydrogen atom positions in O--O hydrogen bonds of intermediate strength, we have undertaken diffraction studies of several hydroxybenzophenones. The crystal structure of 2,4-dihydroxybenzophenone (Liebich, Acta Cryst. (1979) B35, 1186) revealed an intramolecular O--O hydrogen bond (2.55Å). We have completed a neutron study of that compound and will present accurate hydrogen positions as well as an X-N deformation density study of the hydrogen bond. In addition X-ray studies of 2,2', 4,4'-tetrahydroxybenzophenone and 2,2'-dihydroxybenzophenone will be presented. In the former there are two symmetry independent molecular O--O hydrogen bonds involving the carbonyl oxygen with the 2 and 2' hydroxy groups. The O--O distances are 2.52, 2.58, 2.62, and 2.72 at the present stage of refinement (R=0.034). In addition there are intermolecular hydrogen bonds involving the hydrogen atoms of the 4 and 4' hydroxy groups. These involve O--O distances of 2.76, 2.76, 2.87, and 3.01Å, and O-H--O angles from 145 up to 175°. Whereas the two six membered aromatic rings in 2,4-dihydroxybenzophenone are nearly mutually perpendicular, those in 2,2',4,4'-tetrahydroxybenzophenone are more nearly coplanar (dihedral angles of 42 and 44°) because of the intramolecular hydrogen bonding between the carbonyl oxygen and both the 2 and 2' hydroxy groups.

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04.6-04 THE CRYSTAL STRUCTURE OF THE ADDITION COMPOUND OF FLUOROSILICIC ACID WITH UREA. By Ze-Ying Zhang, Mei-Cheng Shao, Xiao-Jie Xu and You-Chi Tang, Institute of Physical Chemistry, Peking University, Beijing, China; Yu-Qin Tu, Institute for Plant Protection Research, Chinese Academy of Agricultural Sciences, Beijing, China.

The addition compound ((NH<sub>2</sub>)<sub>2</sub>CO)<sub>4</sub>·H SiF<sub>6</sub> crystallizes in tetragonal space group P 4<sub>1</sub>2<sub>1</sub>2 with lattice parameters a=9.263(4) Å,c=17.898 (6) Å and Z=4. The structure was solved by direct method and refined for 1362 independent MoKα counter data. The R index is 0.052 and Rw=0.032. The result reveals that the crystal structure consists of octahedral fluorosilicate anions and ureated protons ((NH<sub>2</sub>)<sub>2</sub>CO···H+···OC(NH<sub>2</sub>)<sub>2</sub>). The structure formula is ((NH<sub>2</sub>)<sub>2</sub>CO···H+···OC(NH<sub>2</sub>)<sub>2</sub>)<sub>2</sub>·(SiF<sub>6</sub>). The two protons of a fluorosilicic acid molecule are captured by two pairs of urea molecule and form two strong hydrogen bonds 0-H-O respectively of 2.424 Å and 2.443 Å in length. The center of each such hydrogen bonds sits on a two-fold axis. All F and N atoms are involved in forming hydrogen bonds F···H-N. Such hydrogen bonds have certainly reinforced the framework of the crystal. This addition compound has been found to be an effective and practical agent for prevention and control of plant diseases such as wheat stem rust. The capture of the acidic protons by urea molecules through hydrogen bonding has made the fluorosilicic acid less harmful, more convenient to store and transport, and its effect more persistent.

A recent study (Knop, Westerhaus and Falk, 1980 Can. J. Chem., 58, 867) of infrared spectra of the ammonium ion in  $(\mathrm{NH}_4)_2\mathrm{TiF}_6$  at different temperatures shows that the stretching frequency of all N-H bonds increases with decreasing temperature. The negative values of  $\mathrm{dv}/\mathrm{dT}$  for the N-atoms have been explained by supposing that at low temperatures the hydrogen atoms form trifurcated bonds as required by the space group symmetry but that at higher temperatures the ammonium ion is rotationally disordered to form stronger hydrogen bonds with particular F atoms. In order to examine this hypothesis we have used x-ray diffraction to refine the structure of  $(\mathrm{NH}_4)_2\mathrm{TiF}_6$  at room temperature and at 153K to R = 0.02. The crystal is trigonal, space group P5ml with Z = 1, a = 5.920(2), c = 4.702(1)A (153K) and a = 5.972(2), c = 4.822(1) (290K).

Although hydrogen atoms were included with anisotropic temperature factors in both refinements, a difference map showing the electron density of the H atoms shows the inadequacy of the conventional model. At 153K the H atoms are well localised with one on the 3 fold axis and 3 on mirror planes but at room temperature the unique H atom appears to be disordered off the threefold axis. The relation of this disorder to that proposed by Knop et al. will be discussed.