17.5-02 EFFECTS OF HEAVY ATOMS AND SYMMETRY ON THE CUMULATIVE DISTRIBUTION FUNCTION OF NORMALIZED STRUCTURE AMPLITUDES By <u>G.D.Nigam</u> and Sikha Ghosh, Department of Physics, Indian Institute of Technology, Kharagpur, India.

Expansions for the probability density function of the normalised structure ampli-tude |E|, which account for the known contri-bution of heavy atoms in the unit-cell have been derived. The expressions for the cumu-lative distribution function N(|E|) for Edgeworth expansion have been derived by Mitra and Belgaumkar (Proc. Ind. Acad. Sci. (1973), 31,45), and for Gram-Charlier expansion by Shmueli (Acta Cryst., (1979), A35, 282), Shmueli and Wilson (Acta Cryst. (1981) under print). In the present investigation the expressions for N(|E|), using the above modified probability density function, have been obtained. The expressions are composition and symmetry dependent and lead to those obtained by Shmueli if the heavy atom contri-bution is absent. The polynomial series distribution is also used to derive an expression for N(|E|) in case of a hypercentric crystal. The asymptotic form of the distribution has been used to calculate the effect of heavy atoms on two phase structure semin-variants in P1. Numerical computation has been carried out to study the effect of com-position and symmetry on N(|E|). The results have been compared with those of Shmueli and Wilson and also tested on a few known crystal structures.

17.6-01 CONFORMATIONAL ENERGY ANALYSIS OF BENCE JONES PROTEIN RHE. By <u>A. Chatterjee</u>, B.C. Wang, W. Furey, Jr., C.S. Yoo and M. Sax Biocrystallography Laboratory, VA Medical Center, Pittsburgh, PA 15240 and the Department of Crystallography, University of Pittsburgh, Pittsburgh, PA 15260.

Conformational energy analysis of Bence Jones Protein Rhe starting with coordinates refined to 1.6 Å resolution is carried out by using a suitably chosen empirical conformational energy function and a set of derived energy parameters. In this process the atoms are moved so as to reduce the molecular potential energy in response to perturbation introduced in the system. The energy minimization makes use of the conjugate gradient method by searching along directions corresponding to the current local quadratic approximation to the energy function. The required computer programmes have recently been developed in this laboratory and results will be presented. **17.6-02** CRYSTAL STRUCTURE PREDICTION BY ENERGY MINIMISATION. By C.R.A. Catlow, A.N. Cormack, University College London, and F. Theobald, Laboratoire de Chimie Physique, Besançon, France.

This paper will show that calculations of crystal energetics for ionic solids coupled with efficient minimisation routines are capable of predicting the structure of complex inorganic solids, thereby promising to provide a valuable guide in structure refinements. The calculations we discuss are based on exact summations of both Coulomb and short-range components of the lattice energy. Energy minimisation may be performed with respect to all structural parameters. We illustrate the power of our method with two examples. The first concerns the recently discovered 'B' phases of VO and TiO (Marchand et al., Mat. Res. Bull. 15, 1129 (1980)). Energy minimisations were performed starting from an ideal model based on regular octahedra. The minimised structure shows distortions close to those observed in the X-ray refined model for VO_2 . A predicted structure for the analogous 'B' pha-se of TiO_2 is also presented; no experimental structure is yet available for this phase. Our second example is the mineral forsterite $(Mg_2 SiO_4)$ where again we show that distortions from idealised models are correctly predicted. We conclude with a discussion of potential future application of the technique to complex transition metal compounds and to minerals.

17.6-03 ADAPTION OF THE POTENTIAL ENERGY OF MOLECULAR AND CRYSTAL STRUCTURES TO POWDER DIAGRAMS. By <u>H. Bradaczek</u>, G. Barnickel and H. Labischinski, Institute of Crystallography, Free University Berlin, West Germany.

Recently a method has been developed to carry out potential energy calculations in Fourier space. Using reciprocal sphere potential functions the energy can be described by

$$P_{TOT} = {}^{J}FF_1 \quad dV + {}^{J}FF_2 \quad dV + \dots$$

sphere1 sphere2

where K = 1, 2... corresponds to the kind of interaction in a structure. The radii of the spheres are reciprocal to the interaction width and can be adapted to the first minimum

of the potential functions. FF_{K}^{\star} are the

structure factors corresponding either to interactions of the same atom type or of various ones. (Partial structures). The above equation allows the integration over spherical shells, the result of which is proportional to powder or amorphous structure diffractograms. The comparison opens possibilities to adapt parameters of the potential energy to experimental data. The method is usable for crystals as well as for amorphous structures.