12-Amorphous, Imperfectly Ordered and Quasi-periodic Materials

334 OCM-12.01.04

INCOMMENSURATE PHASES IN CHARGE-DENSITY WAVE SYSTEMS. J.C Bennett¹ and F.W. Boswell², ¹Dept. of Physics, University of Alberta, Edmonton, AB, Canada T6G 1K7 and ²Dept. of Physics, University of Waterloo, Waterloo, ON, Canada N2L 3G1.

Charge-density waves (CDW), a coupled modulation of the conduction electron density and crystal lattice, have been reported in a large number of structurally anisotropic compounds. These CDW modulations, which are in general incommensurate with the periodicity of the undistorted lattice, occur in materials encompassing a wide range of chemical and physical properties. However, in spite of this diversity, many aspects of the observed CDW phenomena are remarkably similiar. In this paper, we will attempt to highlight this through an examination of two model CDW systems: the TaTe₄-NbTe₄ system and NbSe₃.

Extensive experimental and theoretical investigations of TaTe₄ and NbTe₄ have demonstrated that these compounds constitute a nearly prototypical CDW system. The compounds possess a quasi-one-dimensional crystal structure in which chains of metal atoms are centered within extended cages of Te atoms in square antiprismatic coordination. The observed lattice distortions, involving mainly longitudinal motions of the metal atoms along a chain, correspond to that of the classic CDW model. In addition, the compounds exhibit the full spectrum of possible CDW-driven phase transitions: commensurate to incommensurate (C to IC), commensurate to commensurate (C to IC), commensurate (IC to IC). Despite this apparent complexity, it has recently been shown that the various phases occuring among the tetrachalcogenides are rather simply interrelated with each phase representing a member of a series of long period structures. In addition, transmission electron microscopy studies have revealed the important role of microstructure, including discommensuration arrays and antiphase boundaries, in this CDW system.

The crystal structure of NbSe₃ is considerably more complex than that of the tetrachalcogenides and, until recently, it had not been possible to precisely determine the nature of the IC modulations. In analogy with the tetrachalcogenide system, we have developed models for the incommensurate distortions found in NbSe₃. The implications of the new structural models for the observed sliding of CDW under the application of an electric field will be discussed.

OCM-12.01.05

A MODULATION WAVE APPROACH TO COMMENSURATELY MODULATED STRUCTURES. By R.L.Withers*, J.G.Thompson, S.Schmid, Research School of Chemistry, Australian National University, GPO Box 4, Canberra City, ACT 0200, Australia and A.D.Rae, School of Chemistry, University of New South Wales, P.O. Box 1, Kensington, N.S.W. 2033, Australia.

Commensurately or incommensurately modulated structures can always be described in terms of an underlying parent structure (characterized in reciprocal space by a set of sharp Bragg reflections G) plus compositional and/or displacive modulations thereof associated with independent modulation wave-vectors $\{mq_1 + nq_2 + ...\}$ (and characterized in reciprocal space by a set of sharp Bragg reflections at $G \pm \{mq_1 + nq_2 + ...\}$,where q1,q2,... represent independent primary modulation wavevectors and m,n,.. are integers). Given knowledge of the underlying parent structure, structure determination reduces to the determination of the compositional and/or displacement modulation patterns (amplitudes and phases for each atom in the asymmetric unit of the parent structure) associated with each independent modulation wave-vector $\{mq_1 + nq_2 + ...\}$. The symmetry-allowed structural degrees of freedom associated with each independent modulation harmonic are determined by the resultant space group (or the super-space group if the primary modulation wave-vectors are correctly chosen) in the case of a commensurately modulated structure or by the super-space group in the case of an incommensurately modulated structure.

An additional important feature of many such modulated structures is that the amplitudes associated with each independent modulation harmonic often fall off monotonically and fairly rapidly with increasing harmonic order (provided the correct choice of the primary modulation wave-vectors is made) so that there is a natural hierarchy as regards the symmetry allowed structural degrees of freedom. It is demonstrated, via Fourier decomposition of previously reported example superstructure phases, that a modulation wave approach to such superstructures almost invariably provides a much simpler structural parameterization than conventional superstructure refinements using independent atom-based parameters and that the latter, because they fail to take advantage of this natural hierarchy, are often grossly overparameterized.

The use of such a modulation wave approach to the structural parameterization of commensurately modulated structures is reviewed and recent developments in the understanding of such superstructures highlighted. Using examples, the advantages and possible disadvantages with respect to a conventional superstructure approach are discussed as regards the information content of reciprocal space, the possibility of false minima in conventional superstructure refinements and the possibility of homometric (i.e. indistinguishable but non-identical) structure solutions.

OCM-12.01.06

SUPERSPACE SYMMETRY OF COMPOSITE CRYSTALS. By AKIJI YAMAMOTO National Inst. Res. Inorg. Mat. Namiki 1, Tsukuba, 305 Japan

A composite crystal structure is a generic name for misfit layer structures, intergrowth compounds, vernier structures, and chimneyladder structures, which have mutually interpenetrating two or more substructures with incommensurate (or commensurate) period along some (one or two) directions. It has been found in minerals, organic compounds, metals, and in particular in many sulfides. Each substructure is modulated by the interaction to the others. Therefore this is a general case of modulated structures and superspace groups introduced for usual modulated structures can be applied to its symmetry (Janner, A. and Janssen, T., Acta Cryst. 1980, A 36, 408-415). Recently several structure analyses were made on the basis of the superspace symmetry (for examplc, Kato, K., Acta Cryst. 1990, B 46, 39-44, Smaalen, S. van, J. Phys. Cond. Mat. 1991, 3, 1247-1263) and the efficiency of the superspace group in the structure analysis has been proved. The diffraction pattern is indexable with more than 3 vectors (for example $ha_1 * + kb * + lc * + ma_2 *$ etc.) as in modulated structures but is different from that of modulated structures. In the composite crystal, there are several sets of main reflections corresponding to several substructures (Fig. 1). The main reflections of each substructure are at the same time the satellite reflections of the other substructure. The superspace group can be expressed by a pair (or triplet) of the superspace group symbols, each of which specifies the symmetry of a modulated substructure. They are not independent but equivalent to each other as higher-dimensional space groups. The symbol of a superspace group depends on the setting, in particular, the selection of the wave vector of modulation waves. If we employ the wave vector of each modulated substructure within