Fiber I
Synthetic Polymers

MS13.01.01 NEW ROUTES TO THE STRUCTURE OF DISORDERED POLYMERS THROUGH COUPLED SCATTERING AND MODELLING. GP Mitchell, Polymer Science Centre, JJ Thomson, Physical Laboratory, University of Reading Whiteknights Reading RG6 6AF UK

The local structure of disordered polymers remains a challenging area of polymer science. The interactions between chain segments lie at the heart of many central issues including miscibility in blends, crystallisation of non-periodic chains and in deformation processes. Although simulation procedures such as molecular dynamics are powerful tools, their complete reliance on the correctness of the force-fields employed limits their usefulness in many cases. We have developed atomistic molecular modelling tools which allow the tremendous level of information in a neutron or x-ray scattering pattern to be exploited directly in the model construction. The particular utility of broad Q neutron scattering with an extended Q range and the possibility of isotopic substitution will be highlighted. These tools are essentially Monte Carlo procedures in which the fit between calculated and experimental structure factors play the same role as the energy of the system in a conventional simulation. A particular feature of these procedures is the manner in which the chemical connectivity is directly involved. Examples demonstrating the power of these new procedures will be given. We have been able to extract the level of orientational segmental correlations has been obtained from experimental data alone for a series of polymers with differing degrees of backbone flexibility. The possibility of linking these procedures with energy based methods will be discussed.

MS13.01.02 THE CRYSTALLIZATION OF APERIODIC POLYMER CHAINS. Alan Windle, Department of Materials Science and Metallurgy, Pembroke Street, Cambridge, UK CB2 3QZ

Polymers with aperiodic chains have a quite remarkable ability to crystallise. However, there are a number of distinctly different mechanisms which enable a chemically aperiodic chain to contribute to the three dimensional periodicity of a crystal lattice. The paper will review several different systems as examples of mechanistic types, and in each case explore the diffraction conformation and sequence matching in the crystalline phases.

MS13.01.03 X-RAY FIBRE DIFFRACTION STUDIES OF STRUCTURAL VARIATION IN POLYMER MATERIALS. Watson Fuller, Physics Department, Keele University, Staffordshire, ST5 5BG, UK

Recent developments at the European Synchrotron Radiation Facility (ESRF) and the Daresbury Laboratory Synchrotron Radiation Source (SRS) have allowed a dramatic extension of the structural information which can be obtained in fibre diffraction studies of polymer materials. Three types of application will be described. These are: (i) The exploitation of the high brilliance of the two sources to record diffraction patterns with exposure times as short as 40 milliseconds with the facility for real time display of the accumulating pattern. A Photonics Science CCD detector linked to a Synoptic 1860 framegrabber is used to record the pattern and a purpose designed camera allows the development of orientation and crystallinity in polymer materials to be investigated under conditions comparable to those of industrial processing, i.e. draw rates up to 150,000% min, draw ratios up to 4.5 and temperatures up to 350°C. This x-ray camera has a video port which allows changes in the gross appearance of a specimen and hence the draw rate and draw ratio at the point in the specimen from which x-ray data is observed to be continuously recorded and displayed during the experiment. (ii) Beamline ID13 at the ESRF with focusing optics which provide an incident beam at the specimen with a diameter of ~2 microns has been used with an XZ stepping stage to investigate the variation in crystallinity and orientation in spherulites of organic polymers and across polymer interfaces and artifacts fabricated from polymer materials. (iii) Beamline 16.1 at the SRS has been developed to allow the simultaneous recording of the variation in small angle x-ray scattering (SAXS) and wide angle x-ray scattering (WAXS) during drawing and annealing of a number of organic polymers, thus allowing the development of lamellae to be correlated with changes in polymer chain conformation and packing.

MS13.01.04 ON THE STRUCTURE OF A NEW NYLON: POLY(2-METHYL-PENTAMETHYLENE TEREPTHALAL-MIDE). Francois Brisse, Annie Héroux, Département de chimie, Université de Montréal, CP 6128, Succursale Centre-ville, Montréal H3C 3J7, Quebec, Canada

The title aliphatic-aromatic polyamide (2MST nylon) is related to the nT nylon having an odd number, n = 3, 5, 7, of methylene groups. The X-ray diffraction patterns of these polyamides have a very poor resolution. However, that of poly(2-methyl pentamethylene terephthalamide) is very well resolved and yields thirty distinct diffraction spots to a dmax of 2.12 Å. The unit cell dimensions, obtained by a trial and error method using all the observed reflections, are: a = 4.96, b = 28.35, c = 26.82 Å (fiber repeat), and = 88.3°. The density measurement reveals the existence of twelve chemical units per unit cell but since the fiber repeat, p = c, indicates the presence of two chemical units in the c dimension, there are six polyamide chains per unit cell. The value of the fiber repeat precludes a fully trans conformation. The structure of the polymer chain was investigated by conformational analysis and the study of model compounds. The diffracted intensities were used to select and confirm the model most likely to represent the structure of the polyamide. Since the fiber repeat of both the 5T and the 2MST nylon are nearly identical, it is expected that they adopt the same conformation. That the two nylon have very similar diffraction patterns further confirms the above statement.

MS13.01.05 NEUTRON STRUCTURE ANALYSIS OF POLY(VINYL ALCOHOL). Yasuhiro Takashashi, Department of Macromolecular Science, Faculty of Science, Osaka University, Toyonaka, Osaka 560, Japan.

Two crystal structure models were proposed by Bunn and Sakurai et al., which were different in the azimuthal angle of the molecular plane and the hydrogen bonding network. In neutron diffraction, hydrogen atom contributes to the intensity more than X-ray diffraction. Therefore, it is possible to determine the azimuthal angle more accurately and to clarify the position of the hydrogen atom associated with the hydrogen bond. Commercially supplied polyvinyl alcohol) fiber was used for the sample. Neutron diffraction experiments were made at JAEA. The intensity distributions on the equator were measured at 100K, 200K, and room temperatures. The integrated intensity was estimated after the indices were assigned. The refinements were carried out under the rigid-body assumption. Here, the hydrogen atoms associated with the hydrogen bonds were not incorporated, since the hydrogen atom positions cannot be assumed because of the statistical distribution of the oxygen atoms due to the atactic configuration. The refinements by using the data at 100K, 200K, and room temperature gave R-factors, 26.8, 24.8 and 20.3 %, respectively. The