

Book Reviews

Chemical Microstructure of Polymer Chains

Jack L. Koenig

J. Wiley and Sons, 414 pp.
including index + XVIII, £23.00

The value of polymeric materials derives not just from their high molecular weight, nor from the number of atoms of which the molecules are constituted though a large number is necessary of course, but from the identity of the elements concerned and from the way in which they are arranged relative to each other. A simple illustration of this is the organization of DNA, in which five elements (H, C, N, O and P) are arranged to form an extensive backbone and side chain system, whose function is to encode the information describing life at the chemical and biological level. The interrelation between the microstructure of a particular strand of RNA and the proteins derived from it is so well understood that to have the sequence of either is to have a full description of the sequences of both. The reproducibility of the structure of DNA during cell division and the variations which occur during meiosis concern us all, and have done so for some time, though this was little appreciated by our ancestors. Our present understanding of the structure and function of such biological molecules has developed in recent years simultaneously with the design and industrial production of plastic materials. The dichotomy between natural polymers such as wood and wool and products such as polythene may soon cease to be recognized if we can discover other living systems like that whose vital force, enshrined in its DNA microstructure, converts carbohydrate feed into poly(3-hydroxybutyrate). This product might be more useful if it had not a completely perfect tacticity.

For even a simple monomer such as vinyl chloride, the predominant head to tail addition reaction of propagation is supplemented by processes such as reverse addition and side chain production. Whereas in preparative organic chemistry the products of side reactions can be removed, when polymers are produced there is no such possibility: the irregularities are entrained in the polymer chain. The family of catalytic systems discovered by Zeigler and Natta create highly ordered structures and have been a most significant development in this respect. However irregularities in chemical placement and in stereochemistry are not necessarily regarded as imperfections in an ideal structure. They may be tolerated or even desired. What is required is that we can easily recognize them, that we understand their origin so that their prevalence can perhaps be controlled at the preparation stage, and that the link between their presence and the processability and properties of the bulk materials is understood. Koenig's book is devoted to the first and essentially more basic part of such developments in understanding polymeric materials.

The value of ^{13}C n.m.r. spectroscopy and of high field ^1H n.m.r. spectroscopy for

characterizing the microstructure of polymers in solution has been well established in recent years. Hetero atoms in either the main chain or in pendant groups influence chemical shifts in recognizable ways. In purely hydrocarbon polymers, the Grant and Paul parameter set or similar schemes may be applied to the shifts of C atoms in the neighbourhood of crosslinks or branch points to characterize the connectivity of the molecular framework. Provided that side chains are not too remote from each other, stereochemical effects may also be recognized in solution spectra. It is the relative placement of side chains, known well as tacticity, which controls the conformation of the main chain in the solid state and ultimately such bulk properties as the extent of crystallization. Most of the above microstructural features can also be detected in the solid state itself by Raman and infra-red spectroscopy which have also their more straightforward uses as an analytical tool. Moreover, the vibrational spectroscopy of polymers, to which this author has made a major contribution, is sensitive to long range order within one macromolecule such as is found in helical or other extended conformations, to intramolecular packing effects and to features at and on each side of the crystalline-amorphous interface. As well as these two spectroscopic methods, whose experimental techniques and power of discrimination are well illustrated, Koenig indicates the possibilities of e.s.c.a., u.v./visible spectroscopy and mass spectroscopy among others. Certain aspects of chain structure may also be revealed by chemical reactions concerning the polymer.

The structure of a macromolecule is determined by the conditions which prevailed during its preparation. The microstructure records in space what happened in time at the reaction site, and indeed some chemists value this feature alone for the information it provides them about their catalytic species. As we turn our attention from a one-dimensional to a three-dimensional macromolecule, the record of the order of events of preparation becomes less certain, the structural elements become more difficult to define and detect and the statistical apparatus required for analysis becomes more elaborate. The kineticist would wish to describe a preparation in terms of stoichiometry, mechanism and have values for the rate constants, but even for a linear molecule the business is complex enough. For the simplest case, a binary copolymerization to which Bernoullian statistics apply, the author describes the inherent difficulty, which arises from the structure of the errors, in finding the reactivity ratios from feed and copolymer compositions, and he provides the solutions of Tidwell and Mortimer and others. For such a copolymerization reaction, however, monomer sequence information obtained at different feed ratios is more sensitive than rate or polymer composition data obtained under similar conditions for distinguishing between the various possible models. Accounts of the theory of stochastic processes and of the statistical relations between the various possible sequences of repeat units are thus given prior to the treatment of Bernoullian and Markovian models of instantaneous

binary copolymerizations. Subsequently he examines the consequences of the drift of feed composition with conversion for binary copolymerizations, several special cases of multicomponent polymerizations and the influence of extra possibilities such as the formation of charge-transfer complexes between monomers and the occurrence of depropagation reactions near the ceiling temperature appropriate for the addition of one monomer. In a separate chapter the related but simpler topic of side chain stereo configuration is described. The tacticity of both homopolymers and copolymers may be produced by a simple terminal model, or may justify the development of penultimate, penultimate or two stage (Coleman-Fox) models.

We have here a well organized guide to three aspects of polymer microstructure: its definition and statistical description, its characterization by experimental methods and its use for elucidating the polymerization process. There are many, both old hands and newcomers who will find in this book succour and stimulus.

A. H. Fawcett

Developments in Polymer Stabilisation Volume 2

Editor G. Scott

Applied Science Publishers
Ltd, £18

The papers in this volume deal with the function of organotin compounds and metal soaps as thermal stabilizers in PVC. Ayrey and Pollar summarize the synthesis of alkyl tin stabilizers and describe the sporadic attempts which have been made to produce polymeric stabilizers. Their review of the toxicity of organotin derivatives should prove a useful introduction to this important subject. Like Ayrey and Pollar, Cooray and Scott consider in depth the stabilizing roles of organotin compounds. The latter authors point out that in mixing in the presence of small amounts of oxygen mechanochemical scission can give rise to the products, HCl, unsaturation and hydroperoxides, which decrease the thermal stability of PVC. How various stabilizers are effective in nullifying the catalytic effect of one or more of these products is a major topic of this paper.

After a brief review of the literature Guyot and Michel give a comprehensive account of work in their laboratory on the stabilization of PVC. From kinetic studies of the effects of metal chlorides, carboxylates and secondary stabilizers on the thermal dehydrochlorination of the model compound 4-chloro-2-ene an explanation of the stabilizing mechanism of each of these systems is proposed. For the more practical reader examples are given where colour, torque, infra-red, g.p.c. and coulometric techniques have been usefully employed to study PVC stabilization processes.

Pobedimskii, Mukemva and Kirpichnikov in giving an account of the effectiveness of organophosphorous stabilizers and tackling the complex question of how they function at the molecular level provide a valuable review of the considerable Russian work in this area.

The compatibility of a stabilizer with a polymer and how readily it can be extracted are factors which are no less important than chemical efficiency in determining the effectiveness of a stabilizer. The final paper of this volume by Luston is a well written and informative review on the physical loss of stabilizers from polymers. The importance of volatility, and how the structure of polymer and stabilizer affects the cohesion forces between them, and hence their compatibility, are discussed.

Luston presents a detailed account of the physical loss of stabilizers from polymers in which a good balance between experimental observation and theoretical explanation is maintained.

Although in the complex field of PVC stabilization science is still at least one step behind technology the view expressed by Professor Scott in the preface that major advances need a sound mechanistic foundation is valid. The papers presented in this volume make a solid and valuable contribution to this foundation.

A. Davis

Fatigue of Engineering Plastics

Richard W. Hertzberg and John A. Manson

Academic Press Publishers, \$35

One of the most striking features of this excellent monograph is that it is so up to date. Many of the references carry the same year of publication as the book itself yet the flow of the text has not been disrupted by the late insertion of new material, nor is there evidence of hurried and careless proofreading. This indicates admirable dedication on the part of the authors, and probably a great deal of sympathy from the publishers. Their reward will be to earn an extension of the period during which this book will remain essential reading for all research workers and graduate students in the field of mechanical properties and fracture of polymers, and advised reading for every designer working with plastics.

Approximately half of the references cited in the book have appeared since the publication in 1973 of a previous review on this subject by the same authors. If work continues apace it is difficult to see how it will be possible to confine any future review to a single volume if it is to retain the same degree of comprehensiveness and objectivity. Hertzberg and Manson have

conscientiously focused on the subject of the title and do not burden the reader with unnecessary background material or discussion of peripheral topics. The book is well-arranged, the treatment of the material is well-balanced, and it is for the most part easy to read. The subject of fatigue of plastics is of course most complex and the authors have quite properly made no attempt to gloss over those areas in which no clear pattern of behaviour has emerged. This sometimes leaves smooth presentation difficult to achieve, but leaves the reader with the task of forming his own conclusions, a valuable attribute for an advanced text such as this.

The authors have made numerous original contributions themselves and are able to write with authority on most of the topics covered in the book, but they properly refrain from emphasizing their own point of view in those areas in which there exists uncertainty. It is natural that they should use their own studies as a framework for much of the text, but this reflects the depth and breadth of their own work, and they give generous exposure to the contributions of others.

Chapter 1, 'Introduction to Fatigue' efficiently surveys stress systems, thermal and chemical effects, strain-rate dependence, notch-sensitivity and modes of deformation and fracture (including crazes and shear bands). Chapter 2, 'Cyclic Stress and Strain Fatigue: Unnotched Test Specimens', addresses the problem of choosing the most appropriate test variables when evaluating the fatigue properties of materials and considers the relevance of the results to field performance. Apart from alerting the designer to the possible problems that may be encountered when using plastics in engineering applications this chapter also gives some positive advice on component design. Chapter 3, 'Fatigue Crack Propagation', starts with an outline of fracture mechanics concepts and goes on to illustrate the degree of success that has been achieved when evaluating polymer crack growth data according to these principles. A useful, if short, section on design is again included. Chapter 4, 'Fatigue Fracture Micromechanics in Engineering Plastics' describes the micromechanisms of fatigue fracture that have been deduced from fractographic studies and indicates the kinds of features to look for when conducting a failure analysis. Chapter 5, 'Composite Systems', extends the discussion to toughened plastics (including copolymers such as ABS and modified polymers such as HIPS), fibrous composites, (with examples of boron and polyaramid filled materials as well as ones filled with glass-fibre and carbon-fibre), and contains short sections on particulate-filled plastics and adhesive joints.

Apart from the omission of a minus sign in one of the equations and some absences from the list of symbols the most serious criticism I have is that the authors have tended to reproduce the units used in the original

sources. Thus stress is expressed variously in ksi, (and KSI!), kg/mm^2 , lb/in^2 , lb(wt) in^{-2} , kg(wt) cm^{-2} , MPa and MN/m^2 . On p 100 the reader is asked to compare two graphs for which the units of the stress intensity amplitude, (plotted as y), are given as $\text{MPa m}^{\frac{1}{2}}$ and $10^2 \text{PSI IN}^{\frac{1}{2}}$, respectively. It is with pleasure that I predict that few people will be deterred by such minor faults.

J. R. White

Developments in Polymer Photochemistry

N. S. Allen (Ed)

Applied Science Publishers, Vol. 1, 1980, pp. 223; Vol. 2, 1981, pp. 278.

During the last few years *Applied Science Publishers* have made a significant impact on the review literature in polymer science with their 'Developments in' series which now cover subjects ranging from polymerization to polymer technology as well as many non-polymer subjects. These series are now joined by the present volumes which deal with the photochemistry of polymers.

The Editor has rightly taken a broad view of his subject and the seven reviews in each volume range widely covering topics as diverse as photo-initiated polymerization, luminescence spectroscopy, intermolecular energy transfer, photografting processes, photo-oxidation of polymers and photochemistry of dyes and pigments in polymers. The authors are all experts in their fields although it was surprising to find that only one of the fourteen reviews originated in an industrial laboratory, the remainder being from academic or government laboratories.

Of the fourteen reviews in these two volumes, seven are concerned with photodegradation and photostabilization of polymers which means that there is considerable overlap with two other series, *Developments in Polymer Degradation* and *Developments in Polymer Stabilization*. Indeed, many of the authors in the present series have also written for one or both of the others. In view of the relatively small number of people publishing in this field it remains to be seen whether there will be enough material to support three heavily overlapping series in the long term.

The reviews on these two volumes are authoritative and the quality of production is high. For 223 and 278 pages respectively the prices are not excessive and there is much here for anyone interested in polymer photochemistry.

N. C. Billingham