Feature Article

Deformation mechanisms in polymer fibres and nanocomposites

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Abstract

This review covers the development of the understanding of the deformation micromechanics of both synthetic and natural polymeric fibres using spectroscopic and X-ray diffraction techniques. The concept of fibres as composites, where hard and stiff phases are combined with softer polymeric materials is also discussed. Starting with the first discoveries on the molecular orientation and morphology of polymeric fibres, the widely used concepts of uniform stress and strain are examined for the analysis of fibre deformation. The use of advanced techniques such as Raman and infrared spectroscopies to follow molecular deformation in both rigid-rod (e.g. PpPTA, PBO, PBT, polyethylene) and natural (e.g. cellulose, collagen, silk, chitin) polymer fibres is presented. A clear distinction between fibres that have structures that are subjected to uniform stress or strain is presented, with the evidence that is detected from the response of the molecules (by Raman spectroscopy) and the crystalline fraction (by X-ray diffraction). It is suggested that natural fibres, such as cellulose, silk and others, may have different types of microstructures that are subjected to a uniform strain, which could have potentially led to incorrect determinations of crystal moduli. It is also demonstrated that the Raman and X-ray techniques have been influential on our development of fibres, and have shown that the morphology plays a critical role in mechanical properties. In addition to this, the use of X-ray diffraction using microfocus synchrotron sources is also reviewed. This approach allows a more complete picture of both molecular and crystalline deformations to be developed, and with the advent of nanocomposites it is shown that a combination of the two techniques will be vital for our understanding of their exploitation in technological applications.

Keywords: Fibres; Nanocomposites; Deformation

1. Introduction

Polymer fibres and composite materials thereof have made a major contribution to the well-being and technological development of the human race. Aerospace and other advanced industries are driving the need to develop lightweight materials with high stiffnesses, strengths and fracture toughness. With these properties ultimately in mind, progress over the last 30 years into the production of high-performance polymeric fibres has been rapid. In tandem with this progress has been the development of the necessary tools to follow the deformation processes in polymeric fibres to better understand how their performance may be improved. Particular highlights have been on the development of Raman spectroscopy and X-ray (and more recently from synchrotron sources) diffraction, with which the molecular and crystal deformations of polymeric fibres can be followed in detail. Non-polymeric high-performance fibres, such as carbon-based materials, have also been studied widely using these techniques, but this article will primarily concentrate upon the polymeric forms. The work presented has been particularly useful for studying the micromechanics of nanocomposites, which have potential as a new type of high-performance materials. A clear distinction between fibres derived by man-made routes and those of natural origin is made in terms of their structural uniform stress and strain responses to external deformation.

In this article the deformation mechanisms of polymer fibres and nanocomposites using Raman spectroscopy and X-ray diffraction, we will cover the theoretical basis for such investigations, from the early stages of inception upon synthetic fibres to more recent studies of natural polymer-based systems.

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2. Deformation mechanisms in polymeric fibres

It has been well-known for some time that there is a good correlation between optical properties and molecular orientation in polymeric materials. Treloar first discussed the effect of the reorientation of a polymeric material (rubber), and the subsequent changes observed in birefringence measurements [1]. Later, others showed experimentally [2–4] that the optical birefringence in a wide range of polymeric materials could be used to follow the deformation of oriented chains, and could also be related to their crystalline/amorphous structure. Hermans [5] used a similar approach to these investigations for cellulose fibres, and derived orientation functions that could be obtained from simple birefringence measurements. Ward subsequently developed models [6] that interrelated structure, orientation and elastic modulus in idealised semicrystalline polymeric materials. Ward also showed in his seminal paper [6] that the relationship between mechanical anisotropy and orientation was much more complex than that between the static birefringence and orientation, and derived an expression relating the birefringence and the orientation function in a similar manner to Hermans [5]. This expression for birefringence and orientation was also confirmed experimentally for polyethylene filaments [7].

2.1. Fibre mechanics

A large number of publications, before and after these initial discoveries, dealt with the nature of the semicrystalline structure of fibres and filamentous materials, and it is beyond the scope of this review to cover this literature in full. However, it is worth mentioning the work of Halpin and Kardos [8], who for the first time gave upper and lower bands for the predictions of mechanical stiffness of semicrystalline polymeric materials based on composite theory. The lower band due to Reuss [9] assumes that the elements in the structure (i.e. the monomer and polymer molecules) are lined up in series and experience the same stress. The modulus of the polymer in this case, \( E_p \), will be given by an equation of the form:

\[
1/E_p = V_c/E_c + (1 - V_c)/E_m
\]

(1)

where \( V_c \) is the volume fraction of crystals and \( E_c \) and \( E_m \) are the moduli of the crystals and matrix (monomer), respectively. The second model due to Voigt [9] assumes that all the elements are lined up in parallel, and so it experiences the same strain. The modulus of the polymer for the Voigt model is given by the rule of mixtures as:

\[
E_p = E_cV_c + E_m(1 - V_c)
\]

(2)

These upper and lower bands are schematically represented in Fig. 1, where the upper band reflects a uniform strain structure and the lower band a uniform stress structure. This study by Halpin and Kardos [8] allowed the mechanical stiffness of a polymer to be derived on the basis of high and low aspect ratio crystals in an amorphous matrix, with different mechanical properties (i.e. high-modulus crystals and a much lower modulus matrix) without the need for the presence of tie molecules. Thermodynamic equilibrium between the crystalline and amorphous regions was assumed in their models [8], which had initially been considered as a conceptual problem with such morphologies.

Aggregate models, where crystalline domains are considered in series with amorphous domains, were first derived by Ward [6], and developed later by Northolt and vanderHout [10] for highly crystalline fibres. In this series arrangement, the stress in the crystals and amorphous regions is assumed to be uniform. Takayanagi et al. [11] reported the use of a combined parallel-series model in which the stress in the crystals and amorphous regions is non-uniform, with the former overstressed and the latter understressed. In this sense, one could consider, in a fibre with such a morphology, that a stress-transfer process is taking place, and given the dimensions of the crystals, that the structure is that of a nanocomposite. Schematic diagrams of the Takayanagi model are shown in Fig. 2.

2.2. Raman spectroscopy and X-ray diffraction

To better understand the nature of local stress and its development during deformation in polymeric fibres, a number of non-contact techniques with high spatial resolution have been utilised. In this article we will focus on two such techniques; namely Raman spectroscopy and X-ray diffraction. The former is useful for determining molecular deformation and orientation, whereas the latter is sensitive to crystal and amorphous interactions. Orientation analysis, of the crystalline fraction only, is also possible using X-ray diffraction, and this will be covered in more detail later. It will be shown, however, that it is essential to use a combination of both techniques to obtain a complete understanding of the micromechanics of deformation of both semicrystalline polymeric fibres and nanocomposites.

Fig. 3a shows the Raman spectrum of a single aramid fibre obtained using a HeNe laser and Fig. 3b shows the effect of tensile stress upon the 1610 cm\(^{-1}\) Raman band. It has been established [12] that this change in Raman wavenumber, \( \Delta \nu \), during the deformation of high-performance fibres is a result of chain stretching and related directly to the stress on the...
crystalline reinforcing units, $\sigma_r$, such that for an increment of stress

$$d\Delta \nu \propto d\sigma_r$$  \hspace{1cm} (3)

If it is assumed that the stress on the reinforcing units is the same as that on the fibre as a whole (uniform stress) then $\sigma_r$ equals $\sigma_f$, the fibre stress, and so Eq. (3) becomes

$$d\Delta \nu \propto \frac{d\sigma_f}{d\sigma_f} = E_f$$  \hspace{1cm} (4)

by dividing by an increment of fibre strain, $\varepsilon_f$, where $E_f$ is the Young’s modulus of the fibre. Moreover in the case of uniform stress since $\sigma_r$ equals $\sigma_f$ it follows from Eq. (3) that $d\Delta \nu/d\sigma_f$ should be independent of fibre modulus.

An alternative situation that can be employed to model the behaviour is the uniform strain model in which the strain on the reinforcing units, $\varepsilon_r$, is the same as the overall fibre strain, $\varepsilon_f$, which leads to

$$\frac{\sigma_r}{E_r} = \frac{\sigma_f}{E_f}$$  \hspace{1cm} (5)

where $E_r$ is the Young’s modulus of the reinforcing units. Rearranging and using Eq. (3) gives

$$d\Delta \nu \propto \frac{E_f}{E_r}$$  \hspace{1cm} (6)

Since it can assumed that the modulus of the reinforcement, $E_r$, is constant then

$$d\Delta \nu \propto \frac{1}{E_f}$$  \hspace{1cm} (7)

Hence it is predicted that in the situation where there is a uniform strain in the fibre, $d\Delta \nu/d\sigma_f$ should be proportional to the reciprocal of the fibre modulus, $1/E_f$. If the band shift is measured as a function of strain then using Eq. (3) and since the strain is uniform

$$d\Delta \nu \propto \frac{d\sigma_f}{d\varepsilon_f} = E_f$$  \hspace{1cm} (8)

Hence since $E_f$ is constant, then $d\Delta \nu/d\varepsilon_f$ is constant. The results of these predictions for Raman band shifts in the cases of uniform stress and uniform strain are summarised in Table 1.

It is possible to make an analogous set of predictions for experiments upon the deformation of polymer fibres using X-ray diffraction. Fig. 4a shows a wide-angle X-ray diffraction pattern obtained from a single PBO high-modulus polymer fibre and Fig. 4b shows the shift of the (005) and (006) layers that lie in a fibre subjected to tensile deformation. In this
case the shift of the layer lines towards the main beam indicates that the lattice parameter in the chain direction, \( c \), increases on the application of a tensile stress. The crystal strain \( \varepsilon_c \) is defined as \( \Delta c/c \), and assuming that the crystals are the reinforcing units in the structure, and they undergo elastic deformation, it follows that for an increment of stress \( \varepsilon_c \) is related by:

\[
\varepsilon_c \propto \Delta \sigma_f
\]  

(9)

This relationship is analogous to Eq. (3), and so it is possible to make a similar set of predictions for the variations of \( d\varepsilon_c/d\sigma_f \) and \( d\varepsilon_c/d\sigma_f \) with fibre modulus, and \( E_f \) for the situation of uniform stress and uniform strain. These predictions are shown in Table 2. It should be noted that \( d\varepsilon_c/d\sigma_f \) is not usually measured for a fibre but the reciprocal of \( d\varepsilon_c/d\sigma_f \) is normally determined, and a uniform stress assumption made. It is then termed the crystal modulus, \( E_c = d\sigma_f/d\varepsilon_c \).

### 3. Raman spectroscopy and molecular deformation

The Raman effect was first discovered by the distinguished Indian scientist, Chandrasekhar Venkata Raman in 1924 [13]. The secondary radiation that was observed in his experiments was a consequence of inelastic scattering of light by the molecules [14]. In the early developmental stages of Raman spectroscopy, because of low intensity sources of radiation compared to the modern laser systems, and because the effect itself is very weak, infrared spectroscopy of polymers was far more popular as a technique of choice for the characterisation of polymers [15–19]. However, even in these early stages of development, Raman spectra were reported from polymeric materials such as polystyrene [20]. Until the development of laser technology, and particularly the emergence of new optical rejection filters [21] for enhancing Raman Stokes and anti-Stokes spectral peaks, and the microscope or microprobe system for high spatial resolution [22], detailed studies of polymers were not possible. Interest then began into using vibrational spectroscopy to follow local changes in the structure of polymer fibres. Further interest in the use of Raman spectroscopy to characterise polymers has increased with the development of polymers with conjugated backbones which undergo resonance Raman scattering and give strong, very well-defined spectra. After the initial studies on the characterisation of static samples, people began to use these techniques to understand deformation processes and the progress of this work will now be reviewed.

#### 3.1. Background and theory

Perhaps the predecessor for the observations of shifts in the positions of Raman bands of high polymer fibres was the discovery of a similar effect using infrared spectroscopy [23–26]. It became clear during the early 1970s that orientation changes in polymeric materials could be followed using infrared and \textit{in situ} deformation [27]. In addition, and during the same period, Wool showed that the intensity maxima in number of polymers shifted towards a lower wavenumber upon external deformation [28,29]. Initially Zhurkov et al. proposed that the rate of the shifts of different bands with strain within a polymeric material was constant [23]. However, Wool showed that this was not true, and that this factor depended on the “molecular stress distribution via the morphology of
the sample” [29], which is a point which will be addressed at
the later stage in this article.

In 1977 Mitra et al. [30] made the discovery that when con-
jugated polydiacetylene single crystal fibres were deformed in
tension, Raman bands located at 1498 cm\(^{-1}\) (corresponding to
the \(-\text{C}==\text{C}\) backbone moiety) and 2104 cm\(^{-1}\) (correspond-
ing to the \(-\text{C}==\text{C}==\text{C}\) backbone moiety) shifted the position to-
wards a lower wavenumber [30]. The shifts are shown in Fig. 5
[30]. This effect was also predicted theoretically [30] by using
an analysis based on anharmonic force constants of bonds
within an idealised polymeric chain and Badger’s rule [31],
which interrelates the force-constant with the equilibrium sep-
paration of neighbouring atoms [31]. In this sense, Mitra et al.
[30] showed that the shift was due to direct stretching of the
polydiacetylene polymer backbone, and the subsequent
change therefore in the force constants of the bond was asso-
ciated with this structural form. This was argued over the
counter explanations of changes in the local refractive index
of matrix material surrounding a chain, which would also result
in decreased delocalisation and potentially in a band shift
[30]. Batchelder and Bloor also predicted the band shifts
in polydiacetylene based on a model of point masses and
anharmonic spring constants [32].

Previous studies on the plastic deformation of crystals of
bis(p-toluene sulfonate) 2,4-hexadiene-1,6-diol (TSHD) had shown
that the predominant stress-relieving process in such
materials was a twinning or kinking of the polymer chains
[33]. However, since this would not occur in tension, it was as-
sumed by Batchelder and Bloor [32] that polydiacetylene single
crystals could in theory sustain large tensile stresses along
their backbones without the occurrence of plastic deformation
[32]. It is worth noting at this point that Raman spectroscopic
studies of the molecular deformation of all types of polydiac-
etylene single crystals reveal that the band shift rate per unit
strain \(\frac{d\Delta \nu}{de}\) is invariant \((\sim 20 \text{ cm}^{-1}\% \text{ GPa}^{-1})\), irrespective of
the side groups that are present along the main chain [34].

The polydiacetylene crystal or fibre modulus is controlled by
the size of these side groups, and this observation that \(\frac{d\Delta \nu}{de}\) is independent of \(E_t\) means that, with reference to Table
1, the reinforcing units (the polydiacetylene backbones) are
subjected to the same strain as the fibre as a whole (i.e. uni-
form strain).

Inspired by these observations upon polydiacetylene single
crystals, initial attempts to follow band shifts in rigid-rod poly-
mers, such as Kevlar, were, however, unsuccessful, and the
lack of an effect was put down to only a small part of the structure
being under sufficient stress [35]. In 1985 band shifts were
observed for the first time in stressed Kevlar fibres [36], and after
some initial evidence to the contrary [37] it has been shown conclusively that this method is viable for de-
termining local deformation provided that precautions to avoid
radiation damage (e.g. the choice of laser type and power) are
taken into account [38].

3.2. Characterisation of synthetic polymeric fibres

After the initial studies on diacetylenes, Raman band shifts
in high-performance polymer fibres such as poly(p-phenylene
benzobisthiazole) (PBT) fibres were reported [39], although
infrared band shifts had already been observed for these mate-
rials [40]. Significant breakthroughs were then made in obtain-
ing morphological information from stressed polymers, the
most notable around this time being the work of Grubb and
Prasad on high-modulus polyethylene fibres [41]. Band shifts
were not only observed for the 1063 cm\(^{-1}\) peak, corresponding
to the C–C asymmetric stretch mode, but also observed to broaden asymetrically, which was something not seen
for the X-ray diffraction layer lines for the same material under
tensile deformation. This asymmetric broadening was at-
tributed to ‘taut-tie’ molecules within the amorphous regions of
the fibre, and it was found to be only a partially reversible
effect, with relaxation occurring over time. The crystals were
found to go into compression when the load was removed from
the fibres, suggesting that the relaxation of the taut-tie chains was slow. Ultra-high molecular weight gel-spun fibres
were also investigated using the same techniques by another
research group [42]. A later paper [43] found that 40% of
the Raman-active crystalline phase was subjected to high
strain and the mechanisms and morphology proposed were
different to those of Grubb and Prasad [41].

Perhaps the most important contribution to understanding the
deformation of polymeric fibres using Raman spectroscopy was
from a series of papers on rigid-rod materials; namely
PpPTA, PIPD and others [39,43,44]. The first of these studies
detailed only PBT and PBO [39,43] but a later paper outlined
molecular deformation in more detail for poly-aromatic rigid-
rod fibres [44]. This paper [44] showed that for a range of
different fibres (Kevlar, Twaron, Technora, PBO, PET) the
stress-induced Raman band shift for the 1610 cm\(^{-1}\) peak, cor-
responding to the aromatic ring stretch was invariant and equal
to a value of roughly \(-4.0 \text{ cm}^{-1}\text{ GPa}^{-1}\) (shown in Fig. 6a for
Kevlar fibres [44]), although the band shift rates per unit strain
were reported to vary (shown in Fig. 6b for Kevlar fibres [44]).
This result was in spite of the fact that some of the fibres had different chemical structures and were produced using different procedures resulting in different microstructural morphologies. Stress-induced band broadening was also reported for all fibres (Fig. 6c reports this result for Kevlar fibres [44]), which was shown to decrease with increasing fibre modulus for the PpPTA (Kevlar and Twaron) fibres, approaching zero for a fully crystalline structure. Symmetrical broadening of the 1610 cm\(^{-1}\) peak was reported for Kevlar, Twaron, Technora and PBO fibres, whereas PET showed asymmetric broadening. This difference was reported to be due to the fact that some of the bonds in PET must be overstressed, as had been detailed previously [45].

Finally, the deformation of all fibres was shown to follow the continuous chain model of Northolt et al. [46] wherein the structure deforms by a chain extension, orientation and sequential plastic deformation due to shear between the crystal-lites. This model is similar to an approach by Davies et al. [47] who showed that a number of thermotropic liquid crystalline polyesters and polyamides deformed by this mechanism. One outcome of this type of deformation is the increase in the fibre modulus as the deformation proceeds into the high strain region [46,47], which was seen for all the rigid-rod fibre samples analysed by Young and Yeh [44]. This type of strain stiffening has been seen more clearly from the molecular deformation of a number of cellulose fibres by Kong and Eichhorn [48]. The molecular deformation in PBO fibres was also reported [49,50] wherein it was shown for a range of fibres (AS, HM and HM\(+\)) that band shifts in the 1618 cm\(^{-1}\) band, corresponding to the aromatic stretch, could be used for this purpose. It was also found [49] that reduced changes in the FWHM (full-width-half-maximum) of the 1618 cm\(^{-1}\) peak for HM\(+\) fibres could be accounted for due to their more homogenous structure. A second paper in this series [50] showed that less hysteresis in the HM\(+\) fibres supported this hypothesis. Skin-core orientations were also inferred from the distributions of the radial 2\(\theta\) intensity profiles by SAED (Selective Area Electron Diffraction) where the core of the fibres was found to be less oriented than the skin. This type of skin-core orientation had been previously reported for PpPTA fibres [51] where it was shown that different Raman band shift rates could be obtained from fibres with different skin orientations as the Raman signal is recorded only from this region of the fibres.

Galiotis et al. in a recent paper [52] have reported a more detailed description of the effect of stress dependent shifts of two Raman optical phonons \((\nu_1 = 1611\text{ cm}^{-1} \text{ and } \nu_2 = 1648\text{ cm}^{-1})\) for PpPTA fibres at different temperatures. The fibres were deformed inside a furnace, and spectra were recorded under isothermal conditions. It was found that \(\nu_1\) was moderately anharmonic and \(\nu_2\) was harmonic and that both soften with axial deformation, under isothermal conditions, irrespective of the fibre modulus. However, under increased temperatures (above 25\(^\circ\)C) the phonons appeared to harden. This result was also predicted, based on a theoretical approach that treats the fibres as molecular wires with negative expansion coefficients. This paper [52] represents the first

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**Fig. 6.** Raman band shifts as a function of (a) tensile stress, (b) tensile strain for a range of Kevlar fibres (29, 49 and 149); (c) Stress-induced Raman band broadening as a function of fibre modulus for a range of Kevlar and Twaron fibres (data replotted from Ref. [44]. Original data [44] was not fitted with an error weighted linear line. If this is done then the predicted modulus is reduced to 180 GPa).
study of molecular deformation using Raman spectroscopy with environmental control.

3.3. Characterisation of natural polymeric fibres

Raman and infrared spectroscopic characterisations of natural fibres in recent times have developed into an area that is gaining interest worldwide. It must be noted, however, at this point that spectroscopic characterisation of natural materials is fraught with difficulty. It is found that light scattering from such samples is often not very intense and they sometimes undergo fluorescence, and hence longer exposure times are required if one is to obtain well-resolved Raman spectra. Resonance Raman also does not occur for natural polymers such as cellulose, and hence the high intensities seen in polymers such as the diacetylenes are not possible. Nevertheless there have been developments in laser technology that have led to the availability of high-power near infrared lasers that significantly reduce fluorescence.

One of the first reported Raman (and infrared) studies on undeformed plant materials (cellulose) was the work by Blackwell et al. in the early 1970s [53]. Since 1940s people had been using infrared spectroscopy to conduct structural characterisations of hydrogen bonding in plant based cellulose [54]. It is interesting to note that even at this very early stage, Ellis and Bath [54] were able to see changes in the hydrogen bond system of the cellulose structure before and after chemical treatment. After this initial period, a number of detailed papers were published on using infrared spectroscopy for structural characterisation of celluloses by Marchessault and Liang [55–57].

Blackwell et al. [53] obtained Raman and infrared spectra from the cell walls of the algae Valonia ventricosa using an argon ion laser and a Spex-1400 double monochromator, and assigned a number of bands to particular parts of the structure. Later on in 1970s Koenig and Lin also reported Raman spectra for silk, wool, hair and feather [58]. A great deal of effort was made during 1970s by Atalla and Wiley to use Raman spectroscopy to characterise cellulosics, most of which has been extensively reviewed elsewhere [59]. Equipment in those days was reliant on photomultiplier tubes for recording the spectra, but the development of CCD (charge-coupled devices) for recording signals from spectrometers has greatly reduced exposure times [60]. Finally, perhaps the most important part of work on the characterisation of natural fibres has been the work of Edwards et al., including detailed band assignments of cellulosics [61], silks [62], wool [63] and hair [64]. However, it was only in the late 1990s people began to look at the deformation of natural fibres using spectroscopic techniques.

The first study on the molecular deformation of cellulose fibres using Raman spectroscopy was on regenerated forms by Hamad and Eichhorn [65]. This showed for the first time that band shifts, as reported for synthetic fibres, were observable for cellulosic materials. However, there were problems in resolving the shifts with lower modulus fibres, and indeed this study [65] showed that larger band shifts per percent tensile strain applied could be obtained with stiffer fibres. This was indeed achieved with natural plant fibres (flax and hemp) a few years later [66] (Fig. 7) and it was shown that the rate of band shift with respect to strain was directly proportional to the stiffness of the fibres. More studies on these effects were then followed [67,68] to investigate the deformation of wood and other plant fibres in detail. In recent times these types of experiments have been repeated by two independent groups at the University of Halle and at the Max Planck Institute [69–71]. The first of these studies showed that it was indeed possible to detect band shifts in plant material with different chemical and enzymatic treatments [70]. Unlike the original work by Eichhorn et al. [66,67] it was not necessary to pre-bleach the fibres using hydrogen peroxide to overcome problems with fluorescence, as an FT (Fourier Transform) Raman system was used.

Other studies on the deformation micromechanics of cellulose fibres have used infrared spectroscopy. Of particular note has been the work of Salmen and co-workers on the use of dynamic infrared spectroscopy to elucidate the complex hydrogen bond interactions during the loading of the cellulose chains during deformation [72,73]. Without recourse to a full review of the area, the role of the –OH moiety in cellulose spectra is notoriously difficult to interpret. We now know, from detailed studies on the structure of cellulose polymorphs [74–77], that the disorder along the chains and the interdigitation of disorder may play a key role in mechanics and structural interconversion [78], and hence our understanding of the mechanics of this aspect of the structure is vital. Indirect evidence of the breakdown in hydrogen bonding has recently been obtained from Raman spectroscopic studies of deformed cellulose-II fibres [48], but it is difficult to ascertain within which part of the structure (crystalline or amorphous) this is occurring [48]. However, distinct plateaus in the band shift were observed for both the 1095 cm⁻¹ and 1414 cm⁻¹ bands of fibres produced using different draw ratios (Fig. 8), which was attributed to a breakdown in the hydrogen bonding.

Some work has been reported on the use of Raman spectroscopy to probe the deformation of collagen (in the

![Fig. 7. Raman band shifts from the maximum for a flax fibre. 1 = 1122 cm⁻¹ peak and 2 = 1095 cm⁻¹ peak [66] (data replotted from Ref. [66]).](image-url)
The work on collagen fibres showed that it is possible to obtain Raman band shifts, and therefore indications of molecular deformation. However, the deformation on silk, using both silkworm and spider forms of the material, reported that the structure responded in a near uniform stress fashion. The exact conformational structure of silk fibres is still in debate, but most people accept that there are β-pleated sheets (hard segments) within an entangled amorphous network (soft segments) as originally suggested by Termonia [82]. This structure has been confirmed by others in light of further experimental evidence [83,84]. The exact location of the amorphous segments relative to the crystalline β-pleated sheet segments will determine whether the material undergoes uniform stress or strain deformation, but Termonia’s model [82] would appear to be a uniform strain structure.

Various attempts to understand the morphology of silk have been made, starting with modelling of the chain stiffness. The chain stiffness of an alanine–glycine di-peptide structure has been found to be \( \sim 135 \text{ GPa} \) [85]. In spite of this, experimental determinations of the crystal modulus have been comparatively lower in magnitude (in the range 20–28 GPa) [86,87]. The deficiency of the uniform stress assumption for silk fibres has in fact been recently highlighted [86], where it was found that the crystal modulus varied as a function of the crystallinity for a range of fibres (Fig. 9). The values of crystal modulus reported in this study [86] are apparently comparable with theoretical values when using seemingly improved computational analysis, and the acknowledgement of relaxation of the fibre stress is taken into account [88]. However, differences could still be due to the fact that there is an inherent assumption that the structure is subjected to a uniform stress during the measurement of crystal modulus. If this were not true, and the structure was more akin to a nanocomposite, subjected to uniform strain, then one might expect that the crystals would be overstressed, with an understressed matrix component. This has recently been experimentally proven for silk fibres [89], and some evidence for this type of microstructure has also been found in cellulose-II fibres [90]. The work recently reported for silk fibres [89] showed that the Raman band shifts as a function of stress \( (d\Delta v/d\varepsilon_f) \) are proportional to the inverse of the fibre modulus \( (1/E_f) \), i.e. conforming with the uniform strain model (Table 1). This result is in direct contrast to the rigid-rod polymer fibres where there is a proportionality of \( d\Delta v/d\varepsilon_f \) with the modulus \( E_f \) [for instance Ref. [45]], i.e. uniform stress model (Table 1).

### 3.4. Modelling stress-induced Raman band shifts in polymeric fibres

Perhaps, the first attempt to explain a Raman band shift in theoretical terms was made by Mitra et al. [30] who gave an empirical relationship for the effect based on anharmonic potentials, and predicted that the shifts were seen in polydiacetylene single crystal fibres (Fig. 5). Important work has been undertaken by Tashiro [88] upon the prediction of spectroscopic (particularly Raman) band shifts, including work
upon POM (polyoxymethylene) and PE (polyethylene) chains. Using general expressions for the strain energy and potential functions of asymmetric and symmetric vibrations, Tashiro [88] was able to obtain good predictions for the rate of his experimentally determined Raman band shifts. This type of approach was later applied to a PBO chain by Kitagawa et al. [91] and more rigorously for the case of a semicrystalline structure [92].

Now, this approach is adequate in the case where non-bonded interactions do not significantly contribute towards the deformation of the structure. However, little or no work exists on modelling Raman band shifts using molecular force fields. This type of approach uses a normal mode analysis to predict vibrations of minimised structures. Normal mode analyses on static theoretical cellulose structures have been previously performed [93], but not for deformed polymer chains. In recent times Eichhorn and others [90,94] have used a commercial force field, a molecular mechanics approach and normal mode analysis to predict band shifts in cellulose polymorphs (both I and II). Each structure was taken from recently published atomic coordinates [74-76] and minimised within a commercial force field (COMPASSᵀᴹ) under restraint. The normal mode analysis used predicted infrared intensities, and by a process of elimination a number of bands were chosen which best represented the experimentally observed Raman band shifts. Good agreement with the experimental data was obtained, and also with chain stiffness data obtained from X-ray diffraction, a subject to which we will turn to now.

4. X-ray diffraction and crystal deformation

The development of high-performance polymer fibres relies on the ability to maximise the influence of the mechanical properties on the stiff crystals that reinforce them. In a semicrystalline polymer this is never easy, due to misorientation and the effects of the amorphous regions. However, to obtain a theoretical upper limit to the mechanical properties achievable from such fibres, it has been a goal to determine, either by theoretical or experimental means, the inherent stiffness of the crystalline component. So far we have discussed the use of spectroscopic methods, which by their very nature obtain information from both the crystalline and amorphous regions. Therefore, the use of techniques such as X-ray diffraction and modelling has enabled a better picture of crystalline deformation to be obtained.

4.1. Background and theory

One of the earliest studies on the prediction of the stiffness of a polymeric chain was that of Meyer and Lotmar in 1936 [95] on cellulose. This theoretical treatment relied on primitive X-ray diffraction determinations of the structure of native cellulose from ramie fibres [96], and an assumption of a pyranose ring structure for the anhydroglucose units. Nevertheless, values of around 130 GPa were obtained using a classical mechanical approach for the deformation relying on chain stretching and rotation of bonds. Subsequent spectroscopic information allowed Lyons [97] to approach this determination with better precision, but due to an error in his calculation he obtained an overestimated value. This was corrected by Treloar [98] through an alteration in the angle bending stiffness calculations, but his value of 56 GPa was subsequently found to be too low, probably due to a lack of hydrogen bonding in his structure. Treloar also studied other polymers such as polyethylene and nylon [99] and PET [100]. Prompted by Treloar’s efforts, Mann and Roldan-Gonzalez [101] in 1962 measured the crystal modulus of cellulose by deforming Fortisan (cellulose-II) fibre bundles and determined the percentage changes in the crystal strain from shifts in the 040 Bragg reflections. Sakurada et al. simultaneously published a classic paper where they measured, using X-ray diffraction on bundles of fibres held under constant stress, the elastic moduli of crystalline regions of polyethylene, poly(vinyl alcohol), polypropylene, polyoxymethylene and cellulose [102] by assuming a uniform stress response of the structure to applied deformation (Table 2). Examples of the crystal deformation of polyoxymethylene and cellulose are reported in Fig. 10 [102]. Subsequently,
people were then able, from macromechanical tensile measurements and more detailed diffraction studies, to suggest chain-folded structures for polymer fibres [103]. Such folded structures and a full review of the development of morphological studies of this kind are somewhat beyond the scope of this article. However, it is worth noting that this type of structure was refuted strongly for cellulose [104], with some of the arguments against such morphologies being based on the work of Sakurada et al. [102]. This despite initial experimental evidence being in favour of a chain-folded structure [105]. However, a chain-folded structure could be considered to be a fully uniform strain type morphology, and it is clear that this is not always the case, with a possible hybrid model best explaining mechanical properties [90].

4.2. Crystal deformation in synthetic polymeric fibres

During the mid-to-late 1970s people began to develop oriented fibres, of which some, such as aromatic polyesters, were investigated in terms of their crystal response to deformation using X-ray diffraction [106]. With the advent of high-performance fibres, the so-called “rigid-rod” polymers, such as Kevlar [107], new studies on their properties and potential as reinforcements in composites, amongst other applications, were published. The first such study on the crystal structure of Kevlar or poly(p-phenylene terephthalamide) (PpPTA) fibres was reported by Northolt and van Aarts in 1973 [108], although this has recently been refined using neutron diffraction [109]. Subsequently, Northolt went on to elaborate in more detail the mechanical response of highly oriented PpPTA fibres relating crystal modulus and the orientation parameter of misoriented crystals in a formal manner [110] by equation

$$\frac{1}{E_t} = \frac{1}{E_c} + \frac{<\sin^2 \Theta>}{2g}$$

(10)

where $E_t$ is the fibre modulus, $E_c$ is the crystal modulus, $<\sin^2 \Theta>$ is the crystal orientation and $g$ is the shear modulus. Northolt [110] plotted the reciprocal of the dynamic fibre modulus against changes in orientation (determined by X-ray diffraction) for a number of deformed PpPTA fibres to show that this relationship was followed (Fig. 11). Subsequent deformation cycles also showed that the uniform stress model was consistent for these types of fibres.

Another notable study on these fibres was a measurement of the crystal moduli of Kevlar 49, 29, HM-50, Ekonol, Vectran and PEEK (poly(ether ether ketone)) fibres parallel to their axes [111]. For the first time the relationship between fibre properties and crystal modulus was reported [111].

Perhaps the most notable addition to the family of rigid-rod fibres has been the development of the poly(p-phenylene benzobisoxazole) (PBO) types; commercialised as Zylon™ by Toyobo, Japan. The structure was first synthesised by Wolfe and others as part of the US Air Force research programme into high-performance rigid-rod fibres [112]. One of the first publications to appear on the structure and morphology of PBO fibres was by Krause et al. [113]. They used wide-angle X-ray analysis and high-resolution electron microscopy to study the morphology, and reported a modulus of 370 GPa for the fibres compared to 132 GPa for PpPTA. A few years later Martin and Thomas used X-ray diffraction techniques to obtain unit cell dimensions and HRTEM (high-resolution transmission electron microscopy) and SAED (selected area electron diffraction) to obtain lateral spacings between molecules [114]. They were also able to image and characterise defects in the fibres and show that they were edge dislocations. More detailed analyses of the packing disorder, using X-ray imaging and computer simulation, were subsequently reported by Tashiro et al. [115], and in a similar manner to Northolt’s treatment of PpPTA fibres [110], a crystal modulus was estimated to be 370 GPa. This value was later found to be similar to an experimental value (372 GPa) determined by X-ray diffraction of three different forms of PBO fibre bundles [116].

However, the most recent development in the analysis of the crystal moduli of rigid-rod polymeric fibres using X-ray diffraction has been the development of microfocus beamlines, one of which has been at the ESRF (European Synchrotron Radiation Facility) in Grenoble [117]. This development has enabled, for the first time, the determination of crystal moduli of single filaments of high-performance fibres, the first of which to be reported was PpPTA [118]. Single fibres are perceived to be better for such studies, than bundles say for instance, because for the latter, average measurements have to be taken over a larger number of fibres. It is also hard to apply high uniform strains to bundles of fibres, and this could be quite critical if one were attempting to differentiate between uniform stress and strain situations. Following this, a series of papers were published by Davies and co-workers [119–124] using this beamline to study in detail the crystalline deformation and texture analysis of three forms of PBO fibre; namely AS (as-spun), HM (high modulus) and HM+ (an advanced form of HM). The first of these papers [119] showed...
that the crystal moduli of the AS and HM+ fibres were very similar (on an average $\sim 450 \text{ GPa}$) and the HM sample showed the highest overall value of this parameter. Raman spectroscopy was also used to follow the molecular deformation of the fibres and it was shown that a direct proportionality between band shift rate with strain and fibre modulus exists (Table 1), in agreement with a uniform stress model (see Fig. 12). A later paper [120] compared the crystal moduli of PBO fibres to that of PBT and PIPD (initially commercialised as M5 fibres by Magellan and subsequently in partnership with DuPont Advanced Fibre Systems). Similar crystal moduli for PBO and PIPD were obtained ($\sim 440 \text{ GPa}$) but PBT was found to have a much lower value of this parameter (350 GPa). This difference was attributed to the non-linearity of the PBT monomers in the unit cell, and demonstrated the superior properties of PBO, and the benefit of post heat treatment during processing.

The use of a micron-sized beam allows the crystal strain and skin-core orientation of fibres to be mapped spatially across their diameter, and this has been done in some detail using the facility at the ESRF. In one particular study on PBO fibres [121] it was shown that the distribution of crystal strain was uniform across the HM form of the filaments. This study also reported the onset of reorientation across an AS fibre, and showed that this form of deformation mechanism was more significant compared to what was seen for HM and HM+ fibres. This suggested that the uniquely high initial crystal orientation of HM and HM+ fibres contributed to their superior mechanical properties compared to AS fibres. Subsequent experiments also looked at cyclic studies on AS, HM and HM+ fibres and in addition to previous studies, the transverse changes in unit cell parameters [122]. The unit cell angle was found to increase, and the unit cell volume change was found to be invariant for all fibre types. For HM+ fibres the changes in $c$-spacing were comparatively larger than for HM fibres, hence the lower crystal modulus. These changes were however due, for HM+ fibres, to volume change invariance compensated by changes in the $a$ and $b$ cell dimensions. This result confirmed why differences were seen between the fibre types [123]. A major advance in thinking regarding radial fibrillar texturing (RFT) and fibre homogeneity was then reported [123]. This study showed that by using the FWHM (full-width-half-maximum) of equatorial peaks, and an assumption of the $a$-axes oriented radially about the fibre axis, one could show that the degree or distribution of RFT was directly related to fibre modulus. Heat treatment was shown to reduce this distribution. By plotting the FWHM distribution against fibre modulus it was shown that a closer estimation ($\sim 470 \text{ GPa}$) of crystal modulus was obtained then by the Northolt approach [110,115].

Finally, a recent study [124] on heat treated PBO fibres (HM and HM+) has shown that variations (at a spatial resolution of 0.1 μm) in the intensity of layer lines indicate a non-uniform morphology, a factor which contributes to the fact that the fibre modulus does not reach the magnitude of the crystal modulus.

4.3. Crystal deformation in natural fibres

The approaches adopted for highly oriented and crystalline polymer fibres are sufficient to determine the local mechanical properties. However, in the case of semicrystalline and misoriented polymers, such as cellulose, silk, chitin and others, a difficulty exists in the interpretation of results based on seemingly well-intentioned assumptions. In brief, the assumptions of uniform stress and theories of local deformation mechanisms are more difficult to apply when multicomponent and semicrystalline materials are present. In addition to this it must be noted that at low applied stresses, crystal strains will normally increase linearly with fibre stress. However, at higher strains other effects, such as the morphology of the fibres and hydrogen bonding breakdown may influence the form of the deformation. In addition to this, natural polymer fibres tend to be more susceptible to radiation damage in the X-ray beam than synthetic polymers, so particular care must be taken when investigating the materials using a microfocus X-ray beam from a synchrotron source.

The first studies on the crystalline deformation of cellulose using X-ray diffraction were published simultaneously in 1962 [101,102] although only one of these studies was on a native sample [101]. Both studies gave a value of around 130 GPa for the cellulose crystal. These initial studies were followed by an extensive study by Nishino et al. [125] on cellulose polymorphs I, II, IIIa, IIIb and IV giving values of 138, 88, 87, 58 and 75 GPa for their respective elastic moduli along the chain direction. A fundamental assumption of this study was that the stress in the crystalline regions was the same as that applied to the samples. This assumption was shown at a later stage to be valid for a range of synthetically produced polyethylenes produced by different methods, where, irrespective of the sample type a value of 235 GPa was obtained [126]. However, this type of assumption was questioned by Becker et al. [127], since experimental values ranging from 17 to 29 GPa were obtained for silk fibres, which were in stark contrast to theoretical determinations.
As already discussed, this assumption of uniform stress may be invalid for some polymeric fibres, and certainly cellulosic fibres seem to have more of a series-parallel type structure than a series-type configuration [90]. Nakamae et al. [128] attributed their small experimentally determined value of 23 GPa for the crystal modulus of Bombyx mori silk to a 5% shrinkage of the polypeptide chains from a fully extended planar zig-zag conformation, but again one could conceive that in a non-uniform stress situation, the crystal stress is likely to be underestimated, if it is just assumed to be the applied value. Indeed, a recent study by de Oca and Ward [129] has showed for PGA (poly-glycolic acid) crystals that a parallel-series structure might be the best way to represent the deformation of the material, since the theoretical and experimental values of stiffness were so different. Other studies on similar materials to cellulose, such as chitin and chitosan [130] have also shown that seemingly small values of modulus are obtained by these methods, when the uniform stress assumption is applied. Nishino et al. [130] found that α-chitin and chitosan had crystal moduli of 41 GPa and 65 GPa, respectively. These are much lower than values obtained for cellulose, a result that seems strange considering that chitin and cellulose are structurally very similar. It could be that the side chain functionality of chitin changes the conformation of the molecule in chitin, thus reducing its modulus. Nishino et al. [130] carried out identical experiments on chitin when saturated with water and found that the same crystal modulus of 41 GPa was obtained. This result was used to show that the uniform stress model must be valid for these fibres, but again it was assumed prior to the determination of the crystal modulus. It would therefore be interesting to see what value of modulus could be obtained from a non-uniform stress model, such as that proposed by Takayanagi et al. [11].

Other biopolymers have been deformed using the methods developed by Sakurada et al. [102], such as collagen [131]. In this study they found that the chain moduli of a number of samples from various sources ranged from 3–9 GPa. To arrive at these values, Sasaki and Odajima [131] used a quasi-hexagonal packing of chains in the collagen fibrils, a structure that had been postulated from the results of a previous study [132]. An approach like this may be the appropriate method for deriving the true stress–strain curve of cellulosics, and other polymers such as silk, also taking into account the location and proportion of paracrystalline material. Wide-angle X-ray diffraction studies on silk have been performed by Grubb and Jelinski [133], who showed that the crystalline component of the fibres only comprises 12% of the overall structure, with the amorphous component taking up both oriented and misoriented conformations. This may represent the best evidence we have that there are indeed two components of amorphous material in biopolymmeric fibres, but more work is required to confirm the hypothesis that the stress is not uniform in such structures. Recently, the shapes of protein segments in collagen-I have been reported, using a small-angle X-ray diffraction technique to derive their three-dimensional topography [134]. Such an approach could be extended to other biopolymers to better understand the relationship between structure and mechanics.

The deformation of cellulose, as revealed by X-ray diffraction, has been used quite extensively to study the structure–property relationships of wood. Wood itself is a biocomposite of fibres (cellulose) and matrix materials (lignin, hemicellulose). The fibres contain a structural feature within their cell walls called the microfibril angle; the angle to the fibre axis at which the cellulose microfibrils (which are of nanoscale proportions in terms of width) are oriented. In this respect, wood can be considered to be a natural nanocomposite. The longitudinal tensile mechanical properties of wood, such as modulus, have long been known to depend intrinsically on the magnitude of the microfibril angle [135]. This angle can be measured using a small-angle X-ray scattering (SAXS) technique [see for example Ref. [136]]. Distinctive streaks corresponding to the microfibrils in the fibre cell walls can be used to accurately measure this parameter. Although SAXS does not determine the detailed properties of the crystal, it is a useful technique for structurally regular semicrystalline natural fibres such as those that comprise wood. A complete review of the work of Fratzl and co-workers is beyond the scope of this article, but they were the first to identify directly that changes in the microfibril angle of wood related directly to mechanical properties using combined in situ deformation and SAXS [137]. They have also shown that the macroscopic loss in tensile strength and increase in energy absorption are best compromised with a microfibril angle of 27° [138], and that the cell walls of wood were recovered by a so-called “velcro” mechanism after irreversible deformation [139]. The change in the microfibril angle (μ) with applied tensile strain (ε) was shown to follow a simple linear relationship (Fig. 13). A model for the deformation of the wood was also derived which relied on uniform strain and a critical shear stress between the matrix and fibres. This has certain synergies with misoriented reinforcing phases within

![Fig. 13. The change in microfibril angle (μ) of a wet wood foil as a function of tensile strain (ε) as revealed by WAXD (wide-angle X-ray diffraction). Insets show the undeformed and deformed XRD (X-ray diffraction) patterns and the approximate value of the microfibril angle as derived from the integrated intensity around the (002) circle. A linear relationship between microfibril angle (μ(ε) = μ(0) – cot(μ(0))ε) was used to fit the data [139], where μ(0) is the initial microfibril angle of the specimen (data replotted from Ref. [139]).](image-url)
polymeric fibres that have a uniform strain type structure, and hence complementary approaches to their deformation could be applied.

Other notable contributions to the understanding of deformation of wood fibres, but using wide-angle X-ray diffraction (WAXD), has been from the recent water absorption—desorption studies of Abe and Yamamoto [140]. They found that the peak positions of the (200) and (004) reflections shifted during desorption, the form of which indicated a lateral expansion of the crystal, indicating that there must be some mechanical interaction between the amorphous matrix constituents of the materials and the fibres. Lateral changes in the crystal dimensions compared to the longitudinal, again using shifts in the 200 and 004 reflections of the WAXD patterns of wood fibres, have recently been shown to indicate local negative Poisson’s ratios [141]. Negative values of Poisson’s ratio for cellulose fibres have been reported before for cellulose-II [142]. In addition to these effects, it has been shown that the microfibril angle, as derived by SAXS, can decrease after the yield point in the macroscopic deformation of wood fibres [143], a result which has also been seen independently in wet samples of the compressed wood of another species [144].

5. Molecular and fibrous nanocomposites

Discussions will now turn briefly to the deformation micromechanics of fibrous nanocomposites. A full understanding of the local deformation mechanisms in such materials has not yet been achieved within the scientific community, and hence techniques for developing this area are likely to receive significant interest in the near future. Nevertheless some attempts have been made both to determine the interfacial phenomena and to define the laws that govern their properties, for instance, the nanoscale scaling of flaw violations [145].

The work that will be reviewed in this section relates well to the micromechanics and deformation of fibres already discussed. It is now possible to produce nanostructured fibres using electrospinning by the addition of hierarchical pores [146], carbon nanotubes (CNTs) [147] or montmorillonite [148]. The use of CNTs is reminiscent of the ideas first put forward by Takayanagi and his collaborators in 1980; that of a molecular composite [149]. This type of system, where a molecular hard component reinforces a soft polymeric phase, was realised by Takayanagi and others a few years later [150], but only in the form of a film of nylon with phase separated and dispersed chains of PpPTA. The principle of such molecular composites (or as has become customary of late, nanocomposites), as outlined by Takayanagi et al. [149], is that the aspect ratio of the rod-like molecules (in this case PpPTA [150]) should have a high enough aspect ratio, with the covalent bond between reinforcement and the “matrix” material in an ideal state.

It is entirely conceivable, based on the concepts of Takayanagi et al. [149,150] that polymer fibres could be structured in terms of their local orientation to generate the desired mechanical properties by the addition of a second phase. However, one hurdle to overcome is the development of a satisfactory interface, and one way to characterise the efficacy of that property is to use Raman spectroscopy and the other is adequate dispersion, or as stated by Takayanagi et al. [150], phase separation. This degree of dispersion can also be ascertained using Raman spectroscopy, and this will now be discussed.

The concept of a molecular composite, containing diacylenes, was described explicitly by Lando et al. [151], although much had been done prior to that to characterise their deformation. As already mentioned, Raman band shifts indicating the molecular deformation of polydiacetylene single crystal fibres was the first report of its kind [30]. This was, however, followed a few years later by Galiotis et al. [152] who showed that it was possible to follow the interfacial adhesion between two lap-jointed polydiacetylene fibres made using an epoxy resin adhesive. They found that strong well-defined Raman spectra could be obtained from the fibres and that the strains, as determined by a shift in the polydiacetylene Raman peak, were different in each fibre and within the lap-jointed region [152]. This was the first study to show that it was possible to follow the interface between a fibre and a resin using Raman spectroscopy, but a later paper showed that a shear-lag type stress-transfer occurred over the ends of a fully embedded fibre [153] (Fig. 14). The full details of the number of systems that have been subsequently studied by this method are beyond the scope of this review, and are covered in detail elsewhere [154–156]. It is worth noting that the Takayanagi model [11] has been shown to readily apply to urethane—diacetylene segmented block co-polymers [156]. Block co-polymers are rapidly becoming useful for nanostructuring materials, the progress of which has been reported elsewhere [157]. However, the principle of using Raman spectroscopy to map local deformation in such systems relies on the fact that one can use the local molecular deformation of a fibre or phase, as revealed by the shift in the position of a Raman peak, as a sensor of the local stress state within a composite, provided that the resin used is transparent to the laser or does not interfere spectroscopically with the fibre or phase.

![Fig. 14. Axial strain of single poly-1,6-di-(N-carbazoyl)-2,4-hexadiene (pDCH) fibre as a function of position along the fibre in a model composite at different applied matrix strains (εm) (data replotted from Ref. [153]).](image-url)
After this initial work on a lap-jointed composite [152] other papers were published that showed that local deformation could be followed [158–160] in various forms of molecular composites. To the authors knowledge no other papers have been published which follow the local deformation of two-phase nanocomposites using Raman spectroscopy, other than for CNT systems, which are not within the scope of this article, but recently covered in detail by Wagner and Zhao [161]. In recent times, however, the use of Raman spectroscopy to follow the deformation of natural based nanocomposites has been reported. Before discussing these developments, the production and characterisation of polysaccharide-based (e.g. cellulose, chitin and starch) nanocomposites will be reviewed.

The first report on the mechanical properties of a cellulose-based nanocomposite was published in 1995 by Favier et al. [162]. This paper showed that it was possible to isolate rod-like cellulose whiskers, from an edible grade tunicate Microcosmus sulcatus, and disperse them in a resin (latex) to make nanocomposites with good reinforcement capability. A great deal of work has been done to generate a number of cellulose nanocomposites by Dufresne et al. [163,164] and Oskman et al. [165] using a variety of sources for the base reinforcement material. Another large development in the field has been the use of bacterial cellulose (from sources such as Acetobacter xylinum) for the manufacture of transparent and high-performance composite materials [166,167]. These fibres are produced in culture and so are attractive from an environmental perspective. Since they also have nanometre transverse dimensions they make very attractive materials, not only for composites, but also as support for the growth of cells for tissue engineering applications [168–170]. Here the definitions of “high-performance” are best replaced with “functional”, in that the fibres need to reinforce the generated biomaterial, and eventually a composite material is formed.

Other major developments in the use of natural fibres for composite materials have been the production of all-cellulose composites. The concept was first suggested by Nishino and co-workers [171]. The principle of these materials is to dissolve other cellulose fibres and use that as a matrix which is then reinforced by the intact material. Quite remarkable properties have been obtained for these materials, wherein two-fold increases in the modulus compared to epoxy-cellulose have been observed [172]. All-cellulose nanocomposites have also been produced [173] which also show enhanced properties compared to a composite of two chemically distinct phases. In a similar vein, a group led by Teeri has also developed a method of grafting cellulose fibres to other polymeric materials using XETs (xyloglucan endotransglycosylases) [174]. These are enzymes that under certain conditions can generate glucan chains that can then be grafted to materials such as PMMA (poly-methylmethacrylate). In all these systems the concept of an “interfaceless composite”, or at least a non-conventional interface, has to be introduced. Little or no information about the nature of such interfaces exists, although some work has been done on following the relative changes in orientation of dissolved cellulose and a reinforcing fibre using X-ray diffraction from a synchrotron source [175]. These data are shown in Fig. 15 [175] and clear differences between the matrix and reinforcement can be seen, with a linear increase in orientation for both. Clearly great advances have been made in developing these materials, where cellulose is grafted with proteins and phases generated by dissolution and enzymatic catalysis of similar materials, but much needs to be done to better understand the “interfaces” that are forming, and how they deform, whether by uniform stress or strain.

6. Conclusions

The development of high-performance fibres for a variety of applications has invigorated research into the understanding of molecular and crystalline orientations, local deformation and morphology. This article has shown both spectroscopy (in particular Raman spectroscopy) and X-ray diffraction (particularly using synchrotron sources) have been critical techniques for our understanding of these concepts. Early developments in these areas showed that it is possible to predict properties of semicrystalline polymers on the basis of stiff crystals embedded in an amorphous matrix, such as in a nanocomposite. However, it is the morphology of this composite...
structure that will ultimately determine the mechanical properties. Moreover the location of the amorphous material in relation to the crystals affects whether the structure responds under uniform stress or strain. The early determinations of crystal modulus assumed uniform stress for all polymers. It is clear that a distinction between natural biopolymer-based fibres and those produced by synthetic routes can be made, wherein the former generally appear to have uniform strain type structures and the latter uniform stress. It has been shown that general relationships can be made between physical properties, such as Raman band shift and crystal strain and applied stress and strain, that give a clear signature of whether the structure is a uniform stress or strain type. It is therefore clear that only a combination of these techniques can definitively discriminate such structures. Some materials, such as cellulose, have been shown to have more of a hybrid structure, whereas it is clear that measurements upon silks may have been misinterpreted, with most experimental evidence pointing towards a uniform strain type microstructure.

The effect of crystal orientation has also been reviewed as this plays a critical role in the mechanical properties of high-performance fibres. It has only recently become clear, due to the advances in the spatial resolution of X-ray sources (such as synchrotron), that the skin—core orientation of fibres may play a critical role in mechanical properties. Raman spectroscopy really only looks at an average of crystal and amorphous structure, wherein the former generally appear to have uniform strain type structures and the latter uniform stress. It has been shown that general relationships can be made between physical properties, such as Raman band shift and crystal strain and applied stress and strain, that give a clear signature of whether the structure is a uniform stress or strain type. It is therefore clear that only a combination of these techniques can definitively discriminate such structures. Some materials, such as cellulose, have been shown to have more of a hybrid structure, whereas it is clear that measurements upon silks may have been misinterpreted, with most experimental evidence pointing towards a uniform strain type microstructure.

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Professor Young’s main research interest is the relationships between structure and properties in polymers and composites, publishing over 250 papers and a number of books. His research has been supported by the EPSRC, industry and a number of other organisations. He was elected as a Fellow of the Royal Academy of Engineering in 2006.

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