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主論文
Anisotropic Structure of Polyethylene Fiber and Its Related Properties

参考論文
Deformation of Sedimented Mats of Polyethylene Single Crystals

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Dielectric Properties of Mixture of Lauryl-Laurate and Hexacosane
Anisotropic Structure of Polyethylene Fiber and Its Related Properties

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Synopsis

Direct observation of internal structure of a bulky polyethylene fiber was made with an electron microscope using selective etching technique by fuming nitric acid. Striations with a period of 230 Å making an angle of 45 deg. with the fiber axis, and wavy striations with a period of 320 Å perpendicular to the fiber axis, are observed on different fractured surfaces perpendicular to each other. Fibrils of 5000 Å in width are also seen. These values are consistent with a model of double-textured fiber studied by x-ray scattering methods. Amorphous tie molecules between lamellae are oriented perpendicularly to the fiber axis, and parallel to the plane of fiber strip. Tie molecules give the anisotropic structure and the broad range of melting of the fibers.
§ 1. Introduction

Polymer chains are usually oriented to the direction of the resultant maximum strain in the severely drawn sample. When the oriented crystalline polymers are annealed, they give meridional or off-meridional reflections (so-called two-point or four-point pattern) in x-ray small angle scattering (XSAS). The mechanism of fiber formation by drawing was proposed by Kobayashi and by Peterlin. Development of chain orientation in sedimented mat of polyethylene (PE) single crystals was studied through structural changes by hot press in our previous work. Morphological investigation of the final fiber texture in severely drawn PE will help us to understand the mechanism of chain orientation more clearly.

Various models of fiber texture have been proposed by x-ray scattering and electron microscope (EM) studies. On the structure corresponding to the four-point XSAS, morphological investigations having been reported until now were made chiefly by XSAS method. The reliability of the models could be increased by direct EM observation of internal structure of fibers with double texture, but no EM observation has been reported yet. Though Gezovichi et al. succeeded in EM observation of a double texture of as-rolled polyoxymethylene and polyethylene oxide, the resultant strain of their specimens appears to be too small to consider the texture to be a model of fiber with double texture.

We investigated the internal structure of bulky linear-PE fiber with double texture by direct EM observation of fractured surfaces. The PE fibers were prepared in the same way as used
by Seto et al.\(^7\sim 9\), and the anisotropic structure was confirmed by XSAS method. Specimens for the EM observation were made by use of the selective etching technique by fuming nitric acid (FNA) developed by Palmer et al.\(^{14}\), Hay et al.\(^{15}\) and Peterlin et al.\(^{16}\). In the course of the morphological studies, tie molecules were found to play an important role in the anisotropic nature of fiber strips. Hence we investigated and discussed qualitative effects of tie molecules on appearance of the anisotropic structure, and on melting behaviour of the fiber strips.

§ 2. Experimental

Unfractionated linear-PE (Sholex-6009) was molded into 1 mm thick sheets. They were once melted and then quenched in ice water. Strips about 10 mm wide were cut out of the sheets and were drawn at room temps. through the stage of necking. Dimensional changes after the neck-drawing were about 8 times in length, 1/5\sim 1/6 time in thickness, and 4/5\sim 5/6 times in width. The neck-drawn parts of strips were then annealed at a temperature between 122\sim 128°C for half a day under zero stress. The sample preparation so far was almost the same as Seto et al. did\(^7\sim 9\).

In the following, the specimens are represented as "fiber strips" or simply "strips". The axes of coordination in the strip are given in Fig. 1, where the draw axis and the normal of the plane of fiber are chosen as the z- and y-axes respectively.

Two kinds of fractured surfaces of the strips, parallel to the y-z and x-z planes, were prepared at liquid nitrogen temp. to observe the anisotropic structure expected from the reports by Seto et al.\(^7\sim 9\). Since the y-z fractured surfaces were observed
to be covered over with microfibrils introduced by the fracture, they were etched by FNA at 80 °C; the density of the nitric acid is 1.50 g.cm\(^{-3}\). Other strips were treated by FNA at 80 °C for \(\frac{1}{2} \sim 8\) h in advance, and then fractured parallel to the y-z plane.

These fractured surfaces were shadowed with Pt-C along the z-axis and stripped with polyacrylic acid. The replicas were backed with carbon and the polyacrylic acid supporter was removed in water. Because heavily treated strips were too fragile to apply the above mentioned replication technique, some other strips were Pt-C shadowed, immediately backed with carbon, and removed by exposure to vapour of boiling xylene.

The magnification of the EM (HU-11, Hitachi Ltd.) was corrected by 1/576 and 1/2000 mm grating replicas previously to the present experiment. X-ray wide and small angle scattering patterns were taken simultaneously with a point collimated camera, with Ni-filtered Cu-K\(_{\alpha}\) radiation, at room temp$\ldots$.

The experimental procedures to investigate effects of tie molecules on the anisotropic structure, and on melting behaviour of the strips will be given in §4.

§ 3. Texture of Fractured Surfaces

The x-ray scattering patterns in Fig. 2 were taken with incident beams along the x-axis on the right. One-third portion of the y-z plane of a fiber strip, which was so fixed that the developing direction of the neck during the cold drawing was...
directed upwards along the patterns. The central one-third part fixed in the same orientation gives four-point reflections with equal intensity in XSAS. (See, for example, Fig. 8a) The remaining left part gives the mirror images of the patterns in Fig. 2 with respect to the x-z plane. The XSAS with beams along the y-axis gives no trace of off-meridional reflections but meridional two-point reflections. These show that our strips have the same type of double texture and of anisotropic structure as was reported by Seto et al.\textsuperscript{7,8,9) Further they display a biased distribution of the double texture in the strip.

Seto's model viewed along the x-axis is schematically shown in Fig. 3 together with structural parameters. The twin-like arrangement of adjacent fibrils corresponds to the four-point XSAS. The texture is assumed to develop along the x-axis to explain the meridional two-point pattern with x-rays along the y-axis.

From such scattering patterns of strips annealed at 126 °C as given in Figs. 2 and 8a, the thickness of layers composed of crystalline and amorphous parts, D, is determined as 220~240 Å, and the tilting angles of the lamellar normal θ and of chains φ referring to the z-axis are about 45 deg. and 5 deg. respectively. The long period L is calculated to be about 310~340 Å by the relation , \( L = D / \cos \theta \), which is deduced from the model.

As to the width of fibrils, W, we assume a lower limit of W to be 350 Å as a criterion for the EM observation, which is calculated from the lower limit of the coherent length for XSAS reflections estimated by Seto et al.\textsuperscript{8})
To check the FNA treatment effects on the texture, wide angle x-ray pattern of non-treated strips was compared with that of the strip treated at 80°C for 40 h. It is seen that, by the treatment, a disorientation of chains is caused a little, and amorphous halo in Fig. 4a disappears in Fig. 4b. Since the halo in Fig. 4a is stronger in meridional directions than in the equatorial direction, chains in the amorphous regions are partly oriented perpendicularly to the z-axis. The treatment gradually decreases the intensity of XSAS peaks, and increases the intensity of diffuse scattering near the direct spot. (Fig. 4c) The former is considered to be caused by the increase in electron density in the amorphous parts due to oxidation, and the latter is ascribed to voids formed by the treatment. The XSAS pattern Fig. 4c is given by the strip treated for 8 h at 80°C, and is obtained by 10 times as long exposure to x-rays in the evacuated camera as one given in Fig. 2. The D and \( \theta \) remain constant within experimental errors by the treatment for up to 8 h at least.

In the following the word "fractured" will be abbreviated for brevity. The y-z surfaces of non-treated strips are more rough than the x-z surfaces of non-treated strips by the EM observations. From Fig. 5a, structure elements of 5000 Å in width (we call them "fibrils" henceforth) are extending along the z-axis, though they are covered over with microfibrils running to various directions. Some fibrils were found to protrude from the shadowed film, since
they were fluttering due to the electron bombardment during EM observation. We cannot see any other characteristic structure in it. In marked contrast to the y-z surfaces, we can see, on the x-z surfaces, striations with periods of $320 \pm 30 \text{ Å}$, which are rather wavy and perpendicular to the z-axis.

Though the y-z surfaces of etched strips are still covered with microfibrils, their x-z surfaces give a clear lamellar structure (Fig. 6); lamellae, stacked in layers along the z-axis, exhibit their accidental branching and various waviness.

FNA treatment previous to the fracture were found to reduce the microfibrils appreciably, and periodic striations are observed on the y-z surfaces after the treatment for 8 h at 80 °C. (Fig. 7) Since the disappearance of microfibrils is considered to be due to scissions of tie molecules which tightly connect contiguous stacks of crystalline layers, tie molecules will not be easily torn up but bring out microfibrils on the fracture, especially on the fracture parallel to the y-z plane. Hence the different roughness of the x-z and y-z surfaces indicates the preferential orientation of tie molecules parallel to the x-axis. This type of orientation has already been suggested by Seto et al..7,9)

From Fig. 7, the normal of the striations with periods of $230 \pm 30 \text{ Å}$ makes an angle of 45 deg. with the z-axis. The fibrillar structure in Fig. 5a is not observed, and a single region with striations inclining in the same direction spreads in an average width of a few ten microns. Of course, this single region might be composed of some fibrils, and the width of fibrils will be more than 5000 Å at least. This satisfies the criterion described above.
Moreover, Seto et al. assumed that adjacent fibrils with rather small width should shift along the z-axis each other to avoid the creation of excess voids due to misfit between them. Our observation means that fibrils are so wide that the importance of the arrangement of fibrils is reduced.

The biased distribution of the double texture seen in Fig. 2 seems to develop on a more macroscopic scale than the width of each fibril. It should be pointed out that the symmetry of the texture is consistent with that of the plastic strain in an as-drawn strip. We can observe the macroscopic local strain by drawing parallel marker lines perpendicular to the z-axis in advance. After the neck-draw central portions are seen to be drawn more severely than both sides by the distortion of the lines. Such strain pattern is very similar to that of flat metal bar test piece. Hence, in spite of the microscopic texture of the initial strip, the plastic strain is a conspicuous feature of a flat bar test piece. This can explain the symmetry of the texture seen in Fig. 2, because the symmetry will not be lost before the perfect melting. The inhomogeneity of the initial strip, i.e., spherulitic texture, tends to produce local variations in the plastic strain, which determine the inclining direction of lamellae in each fibril.

It is safely concluded that the results obtained by EM observations agree with those determined by x-ray scattering methods, qualitatively in the anisotropic nature and quantitatively in the structural parameters.
§ 4. Tie Molecules in Fiber Strips

Tie molecules have been considered to play some roles on the anisotropic structure,\textsuperscript{7,9} and superheating of PE fibers.\textsuperscript{17\textemdash}19 It is almost confirmed that the FNA treatment cuts tie molecules and weakens the tight connection between lamellae and between fibrils. Hence, effects of the treatment, in other words, effects of decrease in tie molecules on the fiber texture were studied through structural change of the strips by re-annealing. Effects on melting behaviour were also examined with DSC (Rigaku-thermoflex, Rigakudenki Ltd.)

4.1 Effects of tie molecules on the anisotropic structure

Strips annealed at 122 °C were treated for various durations including zero hour, and thereafter re-annealed just below their respective melting points. Their melting points are indicated by $T_3$ in Tables II and III in § 4.2. Corresponding structural changes studied by XSAS are given in Fig. 8 and in Table I.

The double texture in non-treated strip is maintained up to at 132.5 °C (Fig. 8a) and complete disorientation of lamellae is observed at 133 °C. Re-annealing at 132.8 °C causes partial disorientation of lamellae, but some of them still hold their initial orientation. (Fig. 8b) On the contrary, the textures of strips once treated for longer than 1 h are changed by the re-annealing. (Figs. 8c \textemdash}8f) It is notable that such changes will not be caused by stored energy which was accumulated in the amorphous in the course of the previous drawing,\textsuperscript{10} since the texture in non-treated strip does show considerable stability.

The structural change can be explained as follows. Seto et al.
suggested that the volume effect discussed by Flory\textsuperscript{20} could account for chain tilting in oriented semi-crystalline polymers. Following Flory's discussion, we can deduce a modified equation, 

\[ \frac{N_a}{N_c} = \frac{A_c}{2 \cdot A_a \cdot \cos \alpha} \]  

(1)

where \(A_c\) and \(A_a\) are the areas of chain cross-section in crystalline and amorphous regions respectively, and the angle \(\alpha\) is the tilting angle of chains mentioned above. Number of crystalline chains crossing unit area of crystal-amorphous interface is represented by \(N_c\), and the number \(N_a\) is that of chains crossing unit area of a plane placed in the amorphous region apart from the interface by a certain distance. Directions of chains crossing the plane are assumed to be completely random. This condition is not satisfied in the present strips, but here we neglect it. The ratio \(\frac{N_a}{N_c}\) is considered to be an upper limit of number fraction of tie molecules, which leave from the interface and cross the plane at least.

Scissions of tie molecules reduce the ratio \(\frac{N_a}{N_c}\), which leads to the decrease in the angle \(\alpha\) following to eq. (1). Thus we can estimate the ratio \(\frac{N_a}{N_c}\) from the observed angles \(\theta\) and \(\phi\). The last column in Table I is the calculated values of the ratio, where we assume the ratio of \(A_c\) to \(A_a\) equal to unity for simplicity. We obtain a rate of scission of tie molecules to be 9\% per h, which is consistent with the fact that it takes 8 h to observe the y-z fractured surfaces free from the microfibrils.
4.2 Tie molecules and melting behaviour of fiber strips

Typical thermograms are given in Fig. 9 together with characteristic temperatures, $T_1 \sim T_4$. The $T_1$, $T_3$ and $T_4$ are the initiation-, peak- and termination-temps. of the melting respectively. The $T_2$ is the temperature at which the tangential of the melting curve crosses the base line of the thermogram at an angle of 45 deg. in the case of a chart speed of 8 mm per min as shown in Fig. 9. The $T_3$ is regarded as the melting point of the fiber strip. The effect of heating rate and that of duration of the FNA treatment are studied separately. The former is summarized in Table II, and the latter is in Table III.

Non-treated strip shows appreciable rise in $T_4$ with a fast heating rate of 20 °C per min as seen in Table II. The melting point $T_3$ increases a little, but not so appreciable as $T_4$. The high-temp. tails of melting curves are decreased considerably by the FNA treatment.

The effect of treatment duration is very interesting. The treatment of only $1/2$ h lowers $T_4$ by 4.5 °C, and further lowering of $T_4$ is only 2 °C by the treatment for 5 h. (Table III) Since the treatment for 8 h enables us to observe the internal structure free from the microfibrils, it is reasonable to assume that scissions of almost all tie molecules need the treatment for 8 h at least. Further we can assume a constant scission speed from the discussion in §. 4.1. The observation leads to the idea that rather small fraction of tie molecules should owes to the high-temp. tails of melting curves, since a fraction of 93% or more of tie molecules still survives after the treatment for $1/2$ h.
Similar ideas to explain the superheating of stirred crystals of PE were given by Kawai et al.,\textsuperscript{17} Keller et al.\textsuperscript{18} and Zachmann.\textsuperscript{19} Zachmann\textsuperscript{19} studied theoretically the entropy of semi-crystalline polymer, and broad melting in such system could be ascribed to the decrease in the entropy of melting due to two factors; the one was that both ends of non-crystalline parts of chains were fixed, and the other was that volume available to those parts of chains were limited by surrounding crystallites. Such considerations can be applicable to the present strips.

By a short treatment, the $T_1$ is increased and the portion of the melting curves below $T_2$ is lowered as seen in Fig. 9. Tie molecules in semi-crystalline system should take rather extended conformations in thermodynamically equilibrium state.\textsuperscript{19} Since such tie molecules begin to melt or relax at lower temperature due to their characteristic entropies, the semi-crystalline polymer with many tie molecules gives a lower temperature $T_1$ than the one without tie molecules.

\textbf{§5. Summary}

An anisotropic structure of a specially oriented PE with double texture is observed directly with EM by use of the selective etching technique by fuming nitric acid. On the $y$-$z$ fractured surfaces, striations with a mean period 230 Å making an angle of 45 deg. with the z-axis are observed in accord with the off-meridional four-point XSAS. On the $x$-$z$ surfaces, rather wavy striations with a mean period of 320 Å are observed to be perpendicular to the z-axis on an average. Fibrils are estimated to be wider than 5000 Å at least.
The different roughness of the $y$-$z$ and $x$-$z$ fractured surfaces of non-treated strips indicates a preferential orientation of tie molecules to the $x$-axis. Results of EM observation agree well with a model of fiber with double texture proposed by Seto et al.

Reannealing experiments show that the FNA treatment influences the texture through change in tilting angle of chains, and this can be caused by the reduction of tie molecules. Scission of even a small fraction of tie molecules can cause appreciable lowering of the termination-temp. of melting $T_4$, and tie molecules are also responsible on the shape of lower-temp. side of the endotherms of the fiber strips.

Acknowledgements

The author would like to thank Professor K. Asai and Mr. H. Miyaji for their valuable discussions. Thanks are also due to Professor H. Okamoto for his continuous encouragement.
References

2) K. K. Kobayashi: in P. H. Geil "Polymer Single Crystals"
9) T. Seto, T. Hara, Y. Tajima and H. Miyaji: in the preprints of
Table I  Changes in the ratio \( N_a / N_c \) of fiber strips with duration of the FNA treatment by re-annealing.

Strips are pre-annealed at 122°C, then treated by FNA at 80°C and at last re-annealed just below respective melting points. See the text and Fig. 8.

<table>
<thead>
<tr>
<th>Strips treated for</th>
<th>( \theta )</th>
<th>( \phi )</th>
<th>( \lambda )</th>
<th>( N_a/N_c )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 h</td>
<td>48 deg.</td>
<td>5.0 deg.</td>
<td>53.0 deg.</td>
<td>0.83</td>
</tr>
<tr>
<td>1/2 h</td>
<td>46</td>
<td>4.5</td>
<td>50.5</td>
<td>0.80</td>
</tr>
<tr>
<td>3/2 h</td>
<td>40</td>
<td>3.0</td>
<td>43.0</td>
<td>0.69</td>
</tr>
<tr>
<td>3 h</td>
<td>29</td>
<td>1.0</td>
<td>30.0</td>
<td>0.58</td>
</tr>
<tr>
<td>6 h</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.5</td>
</tr>
</tbody>
</table>
Table II  Variation of endotherms of fiber strips with heating rate.

Strips are annealed at 122°C.

<table>
<thead>
<tr>
<th>Strips</th>
<th>Heating rate per min</th>
<th>Characteristic Temps.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$T_1$</td>
</tr>
<tr>
<td>Non-treated</td>
<td>20 °C</td>
<td>100.0°C</td>
</tr>
<tr>
<td></td>
<td>10</td>
<td>103.5°C</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>106.0°C</td>
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<tr>
<td></td>
<td>1</td>
<td>124.5°C</td>
</tr>
<tr>
<td>Treated</td>
<td>20 °C</td>
<td>97.0°C</td>
</tr>
<tr>
<td>for 6 h</td>
<td>10</td>
<td>100.0°C</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>106.0°C</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>124.0°C</td>
</tr>
</tbody>
</table>
Table III. Variation of endotherm of fiber strips with duration of the treatment by FNA at 80°C.

All the strips were pre-annealed at 128°C for 12 h.

Heating rate: 5°C per min.

<table>
<thead>
<tr>
<th>Duration of treatment</th>
<th>Characteristic Temps.</th>
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<tbody>
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<td></td>
<td>$T_1$</td>
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<td>0 h</td>
<td>100.0°C</td>
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<td>1/2 h</td>
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<tr>
<td>2 h</td>
<td>106.0</td>
</tr>
<tr>
<td>5 h</td>
<td>105.0</td>
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</tbody>
</table>
Figure captions

Fig. 1: The axes of coordinations in the fiber strip.

Fig. 2: X-ray scattering patterns of (Right) small angle, and of (Left) wide angle, taken with beams along the x-axis on the right one-third portion of the y-z plane of a fiber strip, which was so fixed that the developing direction of the neck was directed upwards along the patterns.

Fig. 3: Seto's model of fiber with double texture,

Twin-like adjacent fibrils, composed of stacks of oppositely inclined lamellar crystal-amorphous parts are arranged side by side consecutively. Hatched areas are crystalline parts and lines in the areas represent chain axes, which make an angle θ with the z-axis. Blanked areas are amorphous parts.

n_L ; The normal of lamellae,  L ; The long period,

D ; The total thickness of a pair of crystalline and amorphous layers,  W ; The width of a fibril.

Fig. 4: X-ray scattering patterns with beams along the x-axis,

Wide angle patterns of strips (a) non-treated, (b) treated by FNA for 40 h at 80°C. (c) Small angle pattern of the strip treated for 8 h at 80°C.

Fig. 5: Electron micrographs of fractured surfaces of non-treated strips; (a) the y-z surface, (b) the x-z surface.
Fig. 6: An electron micrograph of the x-z surface etched for 4 h after the fracture, Cr-shadowed.

Fig. 7: An electron micrograph of the y-z surface, treated for 8 h previously to the fracture.

Fig. 8: Changes in x-ray small angle patterns of treated and non-treated strips with re-annealing.

Strips were pre-annealed at 122 °C, and FNA treatment was done at 80 °C.

Non-treated, (a) Annealed at 132.5 °C, (b) Annealed at 132.8 °C, Treated, (c) For 1/2 h, (d) For 1.5 h, (e) For 3 h, (f) For 6 h, then annealed just below respective melting points.

Fig. 9: Variations of endotherms of strips pre-annealed at 128 °C with duration of FNA treatment, Heating rate; 5 °C per min.
Fig. 1.

Draw Axis

Plane of Strip
Fig. 2

Fig. 3は次のページです。

Fig. 4
Fig. 3
Fig. 5(a)

Fig. 5(b)
Fig. 8
Fig. 9.

- 0h
- ½h

$C_p$ vs. Temperature

$T_1$, $T_2$, $T_3$, $T_4$