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Mechanism of strain-induced crystallization in filled and unfilled natural rubber vulcanizates

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Structure evolution during deformation of unfilled natural rubber (NR) vulcanizate and filled ones with carbon black or calcium carbonate was investigated by the synchrotron x-ray diffraction. The crystallization onset strain, $\alpha_0$, was found to decrease by the inclusion of the filler. However, corrected $\alpha_0$ values into the effective strain ratio of deformable rubber portion were almost constant between filled and unfilled samples. Accordingly, our model of strain-induced crystallization of unfilled NR vulcanizates, assuming that melting temperature is independent of network-chain length ($n$), was applied to the filled samples. The discrepancy between classical theories and experimental results was thought to come from the distribution of $n$. By the inclusion of filler, the lateral crystallite size was decreased but the orientational fluctuation increased. The lattice of the strain-induced crystallites changed almost linearly with the nominal stress. In addition, the degree of lattice deformation decreased with the filler content, especially in the carbon black-filled system. All these experimental results are consistent with the proposed model. © 2005 American Institute of Physics. [DOI: 10.1063/1.1900927]

I. INTRODUCTION

An elastomer is a polymer material which has the ability to resume its shape after large deformation. The long polymer chains of an elastomer are cross-linked with each other and account for the flexible nature of the material. Among various types of elastomers, natural rubber (NR) is the most typical one and an indispensable material for many industrial and household applications.1 The versatility of NR is mainly due to its outstanding tensile properties and the good crack growth resistance, which originates in its ability to crystallize upon elongation. Thus the strain-induced crystallization in cross-linked polymer system is a significant subject from both academic and industrial viewpoints. Even though the strain-induced crystallization of NR was recognized long ago,2,3 the crystallization mechanism in cross-linked NR had not been much understood in spite of the extensive studies on this subject.2,4-17

There is a characteristic feature in the wide-angle x-ray diffraction (WAXD) patterns of highly strained NR, which is the coexistence of highly oriented crystalline reflections and near isotropic amorphous halo. This feature has been described in the early work by Katz in 1925.4 Mitchell reported the same feature almost six decades later, and investigated the crystallinity and the degree of molecular orientation using the conventional WAXD technique.4,5 One may understand the peculiarity of this feature when the WAXD patterns of other semicrystalline linear polymers, e.g., poly(ethylene terephthalate)18 (PET) or poly(ethylene naphthalene-2,6-dicarboxylate) (PEN),18,19 during the uniaxial deformation were compared with the case of NR. In the case of PET or PEN, as the strain ratio increases, the amorphous halo is localized onto the equator and subsequently crystalline reflections appear, indicating that the overall molecular orientation occurs first and the crystallization follows. Despite the peculiarity, because the isotropic halo from the strained NR could be expected as a result of relaxation during long exposure using a conventional x-ray source, not much attention has been paid. In the subsequent synchrotron WAXD studies, this feature of NR was more clearly recognized by time-resolved WAXD measurements.14,20,21 Since synchrotron x-ray allowed the in situ examination of structural evolution during deformation, the detection of a persistent isotropic amorphous halo in the highly strained NR cannot be related to the relaxation of the stretched chains but to an intrinsic feature of the NR samples.

In the recent quantitative studies by some of the authors of the present paper, the feature was attributed to the coexistence of stretched chains and random coiled chains.4,20,21 A model mechanism has been presented for the strain-induced crystallization of unfilled NR on the basis of the features found through a close inspection of the WAXD patterns, considering the polidisperse distribution of the network-chain length which results in the inhomogeneous deformation of the molecular network of NR vulcanizates.22,23 This model will be extended to the filled system.

In this paper, we discuss the structure evolution and its mechanical response in NR samples containing nanosized particulate fillers. Since the filled NR (e.g., with carbon 0021-8979/2005/97(10)/103529/9/$22.50 © 2005 American Institute of Physics

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TABLE I. Formulations of the NR compounds in parts by weight per hundred parts of rubbers.

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>NR-0</th>
<th>NR-20H</th>
<th>NR-40H</th>
<th>NR-40CA</th>
<th>NR-60CA</th>
</tr>
</thead>
<tbody>
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<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Stearic acid</td>
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<td>2.0</td>
<td>2.0</td>
<td>2.0</td>
<td>2.0</td>
</tr>
<tr>
<td>Active ZnO</td>
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<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>CBS(^c)</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Sulfur</td>
<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
<td>1.5</td>
</tr>
<tr>
<td>Carbon black</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium carbonate</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Curing time(^d)</td>
<td>14</td>
<td>14</td>
<td>14</td>
<td>15</td>
<td>15</td>
</tr>
<tr>
<td>Volume fraction of filler, (\phi)</td>
<td>0.097</td>
<td>0.177</td>
<td>0.135</td>
<td>0.135</td>
<td>0.190</td>
</tr>
</tbody>
</table>

\(^a\) X-cyclohexyl-2-benzothiazole sulphenamide.
\(^b\) Curing temperature 140 °C.

II. EXPERIMENT

The grade of NR used in this study was ribbed smoked sheet (RSS) No. 1 from Malaysia. Carbon black (HS-HAF, Diablack N339) with an average particle size of 26–30 nm and ultrafine calcium carbonate (CaCO₃, ACTIFORD700) with an average particle size of 20 nm were used as fillers. The formulations for the preparation of varying rubber compounds are shown in Table I, along with the volume fraction, \(\phi\), of the filler. Each of the fillers was dried in an oven at 120 °C for 2 h. Subsequently, all of the ingredients were mixed on a two-roll mill at room temperature. The cure time was determined based on unfilled samples.\(^d\)\(^2\) The results of the tensile stress-strain measurements are shown in Fig. 1. For the filled samples, the stress upturn occurred at a smaller \(\alpha\) value, compared to the unfilled sample. This effect was more significant for the samples with carbon black than those with calcium carbonate, according to the comparison of, e.g., NR-40H and NR-60CA having closer \(\phi\) values. This is in accord with the general trend reported previously.\(^d\)\(^2\) Although calcium carbonate of nanometer size (ACTIFORD700) was used, its effect on the tensile behaviors of NR was less than that of a conventional carbon black [high abrasion furnace (HAF)].

A tabletop stretching machine from Instron, Inc., modified to allow the symmetric deformation of the sample, was used to illuminate the same sample position during stretching with the focused x-ray.\(^d\)\(^2\)\(^2\)\(^4\)\(^8\) The incident beam impinged on the sample perpendicular to the stretching direction. Each specimen was continuously elongated up to the predetermined strain ratio and then retracted to the original length, not stopping during the exposure. The strain ratio, \(\alpha\), was determined from the distance between the clamps during the deformation. Here, \(\alpha=\ell/\ell_0\) in which \(\ell_0\) is the initial length of the sample and \(\ell\) is that of the elongated one. The initial distance between the clamps was 25 mm and the deformation rate was 10 mm/min. All the measurements were carried out at room temperature (ca. 25 °C).

III. RESULTS

A. Tensile properties

The results of the tensile stress-strain measurements are shown in Fig. 1. For the filled samples, the stress upturn occurred at a smaller \(\alpha\) value, compared to the unfilled sample. This effect was more significant for the samples with carbon black than those with calcium carbonate, according to the comparison of, e.g., NR-40H and NR-60CA having closer \(\phi\) values. This is in accord with the general trend reported previously.\(^d\)\(^2\)\(^2\)\(^4\)\(^8\) Although calcium carbonate of nanometer size (ACTIFORD700) was used, its effect on the tensile behaviors of NR was less than that of a conventional carbon black [high abrasion furnace (HAF)].

![FIG. 1. Stress-strain curves.](image-url)
B. WAXD patterns of unfilled and filled rubber samples

Figure 2 shows the examples of the WAXD patterns at $\alpha=5$. Only the highly oriented crystal reflections in WAXD are observed in NR. On the other hand, the ring-shaped (unoriented) reflections of calcium carbonate (calcite), which can be indexed as 012 (weak inner one) and 104 (strong outer one) reflections of the trigonal crystal ($a=0.4989\ \text{nm},\ c=1.7062\ \text{nm},\ \text{space group No. }167:\ R_{3c}^7$),\textsuperscript{32} are seen in the WAXD pattern of NR-40CA. As confirmed by the ring-shaped reflections, the filler particles did not have a preferred orientation in the rubber samples upon elongation.

If some sites on the filler particle were acting as nucleating points, randomly oriented crystallites of NR would have appeared, resulting in ring-shaped reflections of NR crystallites. Such ring-shaped reflections of NR have not been found in this study or in the previous studies.\textsuperscript{22,23} The absence of such randomly oriented reflection of NR crystallites strongly suggests that both carbon black and calcium carbonate in this study are not acting by themselves as nucleating agents.

C. Development of strain-induced crystallites

The development of crystallinity was evaluated by using the integrated intensity of 200 and 120 reflections ($I_{eq}$), obtained by a conventional curve-fitting method, from the equatorial intensity distribution that included $\pm 25^\circ$ from the equator. Figures 3(a)–3(e) show the development of $I_{eq}$ with $\alpha$. During the stretching process, the $I_{eq}$ value of all the samples increased monotonically. In addition, during the retracting process, in accord with our previous studies,\textsuperscript{22,23} the $I_{eq}$ value of the unfilled sample decreased monotonically [Fig. 3(a)]. In the case of the filled samples, the $I_{eq}$ value increased when the retracting process started, and then decreased monotonically [Figs. 3(b)–3(e)]. This increase would not be due to the increase of the sample thickness (i.e., the scattering mass, synchronizing the contraction of the sample) because the change in the sample thickness was not so large when the sample was highly stretched; and furthermore, such an increase in intensity has not been observed for unfilled...
The crystallite size was estimated by using the Scherrer equation,

\[ L_{200} = \frac{K \cdot \lambda}{\beta \cos \theta}, \]

where \( L_{200} \) is the crystallite size in the direction perpendicular to the (200) plane, \( \lambda \) is the wavelength, \( \beta \) is the half-width of the 200 reflection in the radial direction, \( \theta \) is the Bragg angle (half of the scattering angle), \( K \) is a constant, and the value of 0.89 was used in this study.\(^{33}\) The detailed procedure for the estimation of \( L_{200} \) has been described in our previous paper.\(^{22}\) Figure 4 shows the variation of \( L_{200} \) during the stretching and retracting processes. As shown in the previous paper,\(^{22}\) the \( L_{200} \) values of all the samples decreased during the stretching process, whereas the values increased gradually during the retracting process. On the whole, the \( L_{200} \) values of the filled samples were smaller than those of the unfilled one.

D. Crystallite size

The crystallite size was estimated by using the Scherrer equation,

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where \( L_{200} \) is the crystallite size in the direction perpendicular to the (200) plane, \( \lambda \) is the wavelength, \( \beta \) is the half-width of the 200 reflection in the radial direction, \( \theta \) is the Bragg angle (half of the scattering angle), \( K \) is a constant, and the value of 0.89 was used in this study.\(^{33}\) The detailed procedure for the estimation of \( L_{200} \) has been described in our previous paper.\(^{22}\) Figure 4 shows the variation of \( L_{200} \) during the stretching and retracting processes. As shown in the previous paper,\(^{22}\) the \( L_{200} \) values of all the samples decreased during the stretching process, whereas the values increased gradually during the retracting process. On the whole, the \( L_{200} \) values of the filled samples were smaller than those of the unfilled one.

E. Orientational fluctuation of crystallites

The degree of crystallite orientation was evaluated based on the azimuthal intensity distribution of the 200 reflection. The azimuthal half-width, \( \beta_{az} \), of the intensity distribution was estimated as described in our previous paper.\(^{22}\) The smaller value of \( \beta_{az} \) indicates the smaller fluctuations in orientation. Figure 5 shows the variation of \( \beta_{az} \) during the stretching and the retracting processes. For each sample, the orientational fluctuation decreased with \( \alpha \) during stretching, and then increased gradually during retracting. It was noted that the \( \beta_{az} \) values of all the filled samples were larger than the unfilled ones.

F. Lattice deformation of strain-induced crystallites

In order to evaluate the deformation of the crystal lattice by the stress upon stretching and retracting processes, the lattice constants were estimated from the WAXD patterns during deformation by using the least-square regression method. The unit cell was assumed to be a rectangular parallelepiped, according to the structure analysis by Nyburg.\(^{34}\) The estimated values of the lattice constants and the volume...
of the unit cell are plotted in Fig. 6 against the stress values obtained in the dynamic experiment. We note that both the stretching and retracting processes are included in this figure. The contraction of the unit cell in the $a$ and $b$ directions (perpendicular to the stretching direction), and the elongation along the $c$ direction (parallel to the stretching direction) with the increase in the nominal stress were recognized in both the unfilled and filled samples. The differences of the lattice constants among the samples at zero stress, which were estimated by the extrapolation of the plots, may be due to the slight differences of the experimental configurations and corresponding errors. Unfortunately, measurements for NR-0 and NR-40H were performed at different experimental periods. The differences of the absolute values of the lattice constants are, therefore, not discussed in this paper.

IV. DISCUSSIONS

A. The model of strain-induced crystallization for unfilled NR

In our previous papers, the following model has been presented for the strain-induced crystallization of unfilled NR.\textsuperscript{22,23}

When the stretching of a NR vulcanizate is started, the network chains in the sample are divided into highly stretched chains and coiled chains, as a result of inhomogeneous deformation due to inhomogeneous topology of chains. (In the field of polymer physics, the term “coiled” is customary used to express the tangled chain conformation in general.\textsuperscript{35}) The coiled chains result in the nearly isotropic amorphous halo in the WAXD pattern.\textsuperscript{20,21,30} Upon further elongation, the melting temperature ($T_m$) of the sample is increased due to elongation of the coiled chains.\textsuperscript{3,4,7,8,11,16,22,23} When $T_m$ exceeds the ambient temperature, the system enters a supercooled state because polymer molecules cannot crystallize otherwise under sufficient supercooling.\textsuperscript{36} Subsequently, the oriented crystallites will be formed onto the stretched chains. The molecular chains that are consumed for the crystal growth (in the same way as the shish-kebab structure\textsuperscript{37–41}) would be supplied by the surrounding coiled chains.

As a result of crystallization, the molecular chains are partly extended in the stretching direction, and the rest, namely, the amorphous portion, is conversely relaxed. Thus the entropy of the amorphous portion will be increased after crystallization,\textsuperscript{4,15,22,30,42} and as a result, further crystallization would be hindered without additional elongation of the sample. The persistence of almost isotropic amorphous halo in the WAXD pattern of highly strained NR samples\textsuperscript{20,21,30} can be explained in this way.

An important postulate assumed in the previous studies is that $T_m$ of the strain-induced crystallites is almost independent of network-chain density, $\nu$. This postulate arises from the result that the $\alpha^0$ values for unfilled NR vulcanizates were almost independent of $\nu$.\textsuperscript{22,23} If $T_m$ were a function of both $\alpha$ and $\nu$, $\alpha^0$ (where $T_m$ minus supercooling equals to ambient temperature) would depend on $\nu$. However, not only in our previous studies\textsuperscript{22,23} but also in another synchrotron study,\textsuperscript{43} $\alpha^0$ was found to be almost independent of $\nu$. Even when the temperature was changed by the ambient condition, $\alpha^0$ was still independent of $\nu$. The latter result was implicitly reported on cross-linked NR;\textsuperscript{7} by plotting the data of $\alpha^0$ against crystallization temperature (in Table IV of Ref. 7), we can obtain curves for three samples with different $\nu$, which are almost superposed with each other as shown in Fig. 7. At least, in the case of “temperature-induced” crystallites, $T_m$ is independent of $\nu$ according to the dilatometric study\textsuperscript{44} of a series of NR samples with different $\nu$ values, though the growth rate of the temperature-induced crystals\textsuperscript{42} was suppressed by increasing $\nu$.\textsuperscript{5} In this way, this postulate is based on sufficient experimental evidences as stated above.

On the other hand, classical theories of strain-induced crystallization for homogeneous network have predicted that $T_m$ depends on the network-chain length, that is, on $\nu$.\textsuperscript{4,11,45} However, there is not enough evidence to prove that $T_m$ depends on $\nu$ for the strain-induced crystallites of vulcanized NR. For example, in Ref. 7, the results shown in Fig. 7 were analyzed by using the Flory’s equation.\textsuperscript{4}
\[
\frac{1}{T_i} = \frac{1}{\tau^0_i} - \frac{R}{\Delta H} \psi(\alpha, n),
\]

where

\[
\psi(\alpha, n) = \left( \frac{6}{\pi n} \right)^{\frac{3}{2}} \frac{\alpha^2}{2} - \frac{1}{\alpha} - \alpha - \frac{n}{n}
\]

with \(T_i\) being the incipient crystallization temperature (namely, the ambient temperature where crystallization starts at strain ratio \(\alpha\)), \(\tau^0_i\) a constant, \(R\) the gas constant, \(\Delta H\) the melting enthalpy, and \(n\) the number of statistical units per network chain. The value \(n\) is not equal to the number of repeating unit per chain, \(N\). The ratio \(N/n\) is a measure of rigidity of the chain. The data that give the plots in Fig. 7 for the samples with different \(N\) have led to different values of \(\Delta H\) and \(N/n\) for each sample. In this way, the dependence of \(T_m\) on \(\nu\) has been believed unquestioningly by regarding other constants as fitting parameters. However, it is unreasonable to attribute different values of \(\Delta H\) and \(N/n\) for the series of NR samples prepared in the similar manner.

The discrepancy between the experimental results and the theoretical prediction may arise from the inhomogeneous nature of the chemically cross-linked NR networks. That is, the premise of the classical theories for the homogeneous network may not be valid in the real NR network system. The coexistence of highly oriented crystalline reflections and nearly isotropic amorphous halo in the WAXD pattern (Fig. 2) suggests the coexistence of highly stretched network chains and coiled chains, as stated before. This would be due to a polydisperse but finite distribution of network-chain length in the real system. A theoretical treatment for such realistic systems, considering the distribution of network-chain length, has been presented by Dietrich et al. Though their model for strain-induced crystallization was different from ours and the theory failed to predict the experimental trends of the crystallization onset strain, the coexistence of highly stretched chains and coiled ones was successfully predicted in their paper.

The hypothesis of the coexistence of two types of network chains has led us to the above-discussed model to satisfy all the experimental evidences consistently. Hereafter, we discuss the results for filled samples from this viewpoint.

### B. Onset of crystallization

As shown in Table II, the strain-induced crystallization of the filled samples starts at a smaller strain ratio than the unfilled sample. This shift of \(\alpha^0\) by the inclusion of carbon black filler has already been reported by Gehman and Field. To explain these results, we have to take account that the filler particles do not deform and only the rubber portion can be deformed. Because of the existence of the filler particles, the effective strain ratio of the rubber portion is larger than the nominal (macroscopic) one. In the simplest model, where no deformation of filler particles and uniform deformation of the rubber portion are assumed, the relationship between the filler volume fraction \(\phi\) and the effective strain ratio \(\alpha_i(\phi)\) can be formulated as

\[
\alpha_i(\phi) = (\alpha - \phi)/(1 - \phi).
\]

Of course, the real value of the effective strain ratio depends on the network structure and the aggregation of filler particles, but we tentatively adopt this equation as the starting point. The \(\alpha^0\) values are converted using Eq. (3) and the values, \(\alpha_i(\phi)\), are shown in Table II. The \(\alpha_i(\phi)\) values are closer to each other compared to the \(\alpha^0\) values.

Two types of correction for the usage of Eq. (3) are considered to explain the relatively large deviation of \(\alpha_i(\phi)\) for samples with carbon black (NR-20H and NR-40H) than those with calcium carbonate (NR-40CA and NR-60CA). In the case of carbon black-filled NR, formation of “bound rubber” layer which is confined (both physically and chemically) around the filler particle is widely known, and the tendency of calcium carbonate is known to be less enhanced. Equation (3) assumes that the whole portion of rubber deforms uniformly while the filler particles do not deform. If we further assume that the bound rubber layer does not deform upon elongation as well as the filler particles, the smaller \(\alpha_i(\phi)\) values in Table II for the carbon black-filled samples may be explained by the underestimation of the volume fraction of undeformable components. In the reverse way, we calculated the thickness of the bound rubber layer from \(\alpha^0\) to be ca. 3–4 nm. (The calculation procedure is described in the Appendix.) The calculated thickness of the bound rubber is within the range of reported values.

The other type of correction considers the aggregate structure formed by the elementary particles of carbon black. The aggregate of carbon black particles traps a part of matrix rubber in the internal void space, and accordingly, the effective volume of the aggregate is larger than the net volume of carbon black particles. The trapped rubber in the carbon black aggregate is called “occluded rubber” and also contributes to the undeformable component. The effective volume fraction, \(V\), of carbon black including the occluded rubber can be estimated by using dibutylphthalate absorption (DBPA) values in \(\text{cm}^3/100 \text{ g}\) as

\[
V/\phi = 0.5 \{ 1 + [1 + 0.02139(\text{DBPA})]/1.46 \}. \tag{4}
\]

In the case of the carbon black used in this study (HS-HAF N339), DBPA is about 120 \(\text{cm}^3/100 \text{ g}\) (according to ASTM D-1765) and the corresponding value of \(V/\phi\) is calculated to be 1.72 by using Eq. (4). The corrected \(\alpha^0\) values in Table II were converted to \(\alpha_i(\phi)\) using Eq. (3).
based on Eq. (3) by using \( V \) instead of \( \phi \) [namely, \( \alpha_0^s(V) \)] for NR-20H and NR-40H are shown in Table II along with the values of \( V \). The deviations of \( \alpha_0^s(V) \) values for the carbon black-filled samples from the \( \alpha_0^s \) value of the unfilled sample are comparable with those of \( \alpha_0^s(\phi) \) for the calcium carbonate-filled samples. Slight fluctuations of these values may be partly due to the oversimplification of the model assumed for the derivation of Eq. (3).

The bound rubber and occluded rubber coexist in the real system, i.e., the undeforume rubber component would be composed of both of them. However, their relative contributions to the real effective strain are presently very difficult to estimate. Therefore, we have estimated each of the effects separately, as described above. The more detailed quantitative analysis is in the scope of future works.

Because \( \alpha_0^s \) for the unfilled sample, \( \alpha_0^s(\phi) \) for the calcium carbonate-filled samples, and \( \alpha_0^s(V) \) for the carbon black-filled ones distribute in the small range, we may be able to assume them to be constant in the range of experimental error for the samples used in this study. Under this assumption, the model for the onset of strain-induced crystallization in our previous papers\(^{22,23} \) may be applicable also for the filled NR. That is, \( T_m \) of NR would be determined as a function of the effective strain of the deformed rubber portion. Apparent increase in \( \nu \) by mixing with filler would not affect \( T_m \). Strain-induced crystallization will start when \( T_m \) of NR is increased with the increase in the effective strain and the system enters a supercooled state.\(^{11,22,23} \) Here, we further assumed that the fillers do not suppress the melting temperature. This is because filler particles are too large to be incorporated in the crystal lattice of NR,\(^{34,55} \) and accordingly, they will not affect the melting temperature.

### C. Growth of crystallites

In the previous paper, on the basis of the observations that the crystalline reflections of NR were highly oriented even when they started to appear, we proposed that the strain-induced crystals are nucleated from the stressed network chains. This assumption was consistent with the result that the values of \( L_{\text{200}} \) were smaller for the samples with the larger \( \nu \); the more densely crystallization starts, the smaller crystallite size results due to the limited availability of coiled network chains. Note that the resulted crystallites can be either chain-folded lamellae or sheaflike (fringed micelle) crystals,\(^{25} \) depending on the longitudinal crystallite size, the available length of network chains, strain ratio, and so on. Such a morphological change with the strain ratio was suggested by Gent.\(^{5} \) We note that the reason for the smaller crystallite size in the samples with the larger \( \nu \) should not be due to the suppressed rate of crystallization.\(^{24} \) Strain-induced crystallization is a fast process that has a time constant of ca. 100 ms,\(^{10} \) and accordingly, the observed size of the crystallites must reflect the size near the completion of the growth. Furthermore, suppression of crystallinity due to the higher \( \nu \) has not been observed in our previous experiments.\(^{22,53} \)

The smaller crystallite size in the filled samples (Fig. 4) can be explained in the similar way. According to transmission electron microscopy (TEM) of stretched NR samples filled with carbon black,\(^{56-58} \) the filler particles (or the aggregates of filler particles) are connected by dark bands running in the stretching direction. The strain-induced crystallites may be formed in the bands, considering the origin of diffraction contrast in bright-field TEM images.\(^{59} \) A strain analysis based on the finite element method\(^{60} \) (FEM) also leads to the same interpretation. The result of FEM indicates that adjacent filler particles are connected by the region of high strain in deformed rubber. Therefore, there should be much stretched chains between the filler particles. These stretched chains will lead to dense occurrence of crystallites. On the basis of the TEM and FEM studies, the smaller crystallite size of the filled samples may be attributable to the dense crystal formation on the stretched chains between the filler particles.

If the crystallites are formed from preciously stretched chains which are bundled with each other, it is very difficult to explain why the smaller crystallite size results from the more densely existing stretched chains. The reason why the stretched chains do not gather together to form crystallites must be due to their very low mobility. Being tightly stretched would be incompatible with the mobility for the network chains. In order to attain the mobility to form the crystal lattice, the network chains must be relaxed to some extent. Given this perspective, the observed increase in crystallinity [Figs. 3(b)–3(e)] at the beginning of the retracting process can be explained by the crystallization incorporating the slightly relaxed chains that attained the mobility from the stretched immobile state. The director of the stretched chains can be fluctuated to connect the filler particles. This would result in larger orientational fluctuations for the filled samples (Fig. 5).

### D. Stress field around the strain-induced crystallites

Based on the linear relationship between the nominal stress and the lattice constants, we have proposed that the transfer of the stress to the crystallites is mediated by frames of surrounding chains.\(^{22} \) We can observe the similar linearity between the nominal stress and the lattice constants for the filled samples in Fig. 6. Therefore, it is conceivable that the similar mechanical model of the stress transfer for the unfilled sample can be applied to the filled ones. It is seen that the absolute values of the gradients in Fig. 6 decrease with the increase in filler content. This trend is apparent for the carbon black-filled samples, and is slightly recognized for the calcium carbonate-filled ones. The smaller degree of lattice deformation in the filled systems indicates that the smaller stress is applied to the crystallites due to the presence of filler particles.\(^{23} \)

The mechanism on how the smaller stress is applied to the NR crystallites by the inclusion of filler particles can be explained with the schematic illustration in Fig. 8 which is drawn on the basis of the discussion in Ref. 22. (We note that there are coiled chains surrounding the frames and crystallites in Fig. 8, though the coiled chains are not drawn.) When the system is stretched in the vertical direction, the frame will compress the crystallites in the horizontal direction. If an elementary filler particle (average size: 20–30 nm) occu-
The comparison of IR with NR suggests the importance of the strain-induced crystallites as filler.

The reinforcing effect of heterogeneous filler, especially of carbon black, is much straightforward compared to that of strain-induced crystals. The larger reinforcing effect of carbon black than calcium carbonate (e.g., in Fig. 1) has been explained by the formation of bound rubber layer and aggregate structure of carbon black particles. Though differences between carbon black and calcium carbonate were detected in the \( a_s^0(\phi) \) values in Table II and in the change in the slopes of the plots in Fig. 6, detailed discussion about the tensile properties, effect of filler, and crystallization requires additional experimental evidences. These points are the subjects in our future works.

V. CONCLUSION

The synchrotron WAXD and the simultaneous stress-strain measurements were performed for the filled and unfilled NR vulcanizates. The following results were obtained.

1. The \( a_s^0 \) value was shifted to the lower value by the inclusion of fillers. If we consider the effective strain ratio of the deformable rubber portion in the filled samples, the strain-induced crystallization may start at almost the same strain ratio for the unfilled sample in this study.

2. By the inclusion of fillers, the lateral crystallite size, \( L_{200} \), became smaller, while the degree of orientational fluctuation, \( \beta_{av} \), became larger.

3. The lattice of the strain-induced crystallites was deformed almost linearly with the nominal stress. The degree of deformation was smaller for the samples filled with the larger amount of carbon black. The samples filled with calcium carbonate showed the similar but weaker trend.

On the basis of these results, we extended the model for the strain-induced crystallization of unfilled NR described in the previous papers, to explain the crystallization in the filled NR. The important assumption for this model is that \( T_m \) is independent of \( \nu \), which may be valid for the filled NR vulcanizates because of conclusion (1). Another assumption is that the strain-induced crystals grow consuming the surrounding coiled chains, which was supported by the observation of the small crystallite size for the filled samples.

The lattice deformation of the filled and unfilled samples could be explained by considering the pantograph-like frame structure and by assuming that the crystallites tend to be compressed by the frames in the direction perpendicular to the stretching direction.

### TABLE III. Values estimated for the calculation of the thickness of the bound rubber layer.

<table>
<thead>
<tr>
<th>Sample code</th>
<th>Onset strain of crystallization, ( \alpha_c )</th>
<th>( \phi_0 )</th>
<th>( \phi_{av} )</th>
<th>( (\phi_{av}/\phi_0)^{1/3} )</th>
<th>Thickness of the bound rubber layer (nm) assuming the average particle size</th>
</tr>
</thead>
<tbody>
<tr>
<td>NR-20H</td>
<td>2.4</td>
<td>0.10</td>
<td>0.20</td>
<td>1.3</td>
<td>3.5</td>
</tr>
<tr>
<td>NR-40H</td>
<td>2.3</td>
<td>0.18</td>
<td>0.34</td>
<td>1.3</td>
<td>3.2</td>
</tr>
</tbody>
</table>

FIG. 8. Mechanical model to explain the deformation of crystallites for the filled sample. The filled squares and shaded circles represent NR crystallites and filler particles, respectively.
In this way, the model presented in our previous papers could consistently explain the strain-induced crystallization of filled NR and the mechanical response of the crystallites.

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APPENDIX

The thickness of the bound rubber layer was calculated as follows. First, we assumed that the properly corrected on-set strain of crystallization (namely, $\alpha_0^*$) was constant among the samples used in this study, i.e., $\alpha_0^* = 3.0$ for all the samples [substituted to the left side of Eq. (3)]. Next, the effective volume fraction of undeformable component containing the bound rubber, $\phi_{\text{eff}}$, was calculated by solving Eq. (3) for the measured (uncorrected) $\alpha_0$ substituted to the right side. The $\phi_{\text{eff}}$ values calculated in this way were compared with the net volume fraction of filler, $\phi$, indicated in Table I. The fraction $\phi_{\text{eff}}/\phi$ is the volume expansion ratio of the undeformable component due to the formation of the bound rubber layer. The linear-expansion ratio of the undeformable component, namely, the expansion ratio of particle diameter, $d$, due to the formation of the bound rubber layer is $(\phi_{\text{eff}}/\phi)^{1/3}$. The thickness of the bound rubber layer is calculated from the average particle size, $D$, of the carbon black and the linear-expansion ratio to be $D[(\phi_{\text{eff}}/\phi)^{1/3} - 1]/2$. The values estimated for the calculation of the thickness of the bound rubber layer are indicated in Table III.

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