Multiaxial deformations of end-linked poly(dimethylsiloxane) networks. 4. Further assessment of the slip-link model for chain-entanglement effect on rubber elasticity.

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Multiaxial deformations of end-linked poly(dimethylsiloxane) networks.  
4. Further assessment of the slip-link model for chain-entanglement effect on rubber elasticity

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The Edwards–Vilgis slip-link model for the chain-entanglement effect on rubber elasticity is critically assessed on the basis of quasiequilibrium biaxial stress–strain data of end-linked polydimethylsiloxane (PDMS) networks with different entanglement densities. The PDMS networks with different entanglement densities were prepared by end-linking end-reactive long precursor PDMS in solutions with different solvent contents. The slip-link model, in which trapped entanglement is modeled by fictitious mobile slip-link attaching two entangled chains, satisfactorily describes the biaxial data over the entire range of deformation for all the networks examined. The model-specific parameters, i.e., slippage of slip-link (\(\eta\)) and inextensibility of network (\(\alpha\)), were employed as adjustable parameters in data-fitting. The fitted values of \(\eta\) and \(\alpha\) vary reasonably with the degree of dilution at network preparation, i.e., entanglement density. With an increase in dilution, i.e., decrease in entanglement density, \(\eta\) increases, whereas \(\alpha\) decreases. In addition, the fitted values of \(\eta\) and \(\alpha\) are in good agreement with the estimates from another molecular approach independent of mechanical testings: \(\eta = M_e/M_c\), where \(M_e\) and \(M_c\) are the molecular masses between neighboring entanglements and between adjacent cross-links, respectively; \(\alpha = n_i^{-1/2}\), where \(n_i\) is the number of Kuhn segments between adjacent elastically effective junctions including cross-links and trapped entanglements. The satisfactory data-fit with the model parameters of physically reasonable magnitudes supports the validity of the slip-link model for entanglement effects on rubber elasticity. © 2003 American Institute of Physics. [DOI: 10.1063/1.1555636]

I. INTRODUCTION

High elasticity of cross-linked rubbers is dominantly entropic in origin, so that the main aim of molecular theory is to consider how the number of configurations available to the constituent polymers changes under deformation. On the basis of entropic elasticity of a single Gaussian chain, the classical rubber elasticity theories derived the elastic free energy (\(F\)) for a network composed of “phantom chains” which may pass freely through its neighbors,1–5

\[
\frac{F}{RT} = \frac{1}{2} C \left( \sum_{i=1}^{3} \lambda_i^2 - 3 \right),
\]

where \(R\) is the gas constant, \(T\) is the absolute temperature, \(\lambda_i\) (\(i = 1, 2, 3\)) is the principal ratio in the \(i\)th coordinate direction, and \(C\) is a constant related to network structure such as the numbers of elastic chain and cross-link. The system comprised of phantom chains is essentially like an ideal gas with.

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mation varying independently each of two principal strains (general biaxial strains) is one of the most rigorous experimental assessments to identify the models which account for the entanglement effect most successfully and correctly, because general biaxial strains achieve the whole accessible pure homogeneous deformations for an incompressible material. The arguments relying on only simple uniaxial deformation are insufficient to distinguish unambiguously the difference between the models, because uniaxial deformation is only a particular one among all the accessible deformations. Among the five theoretical models tested, the predictive capability of the slip-link model was obviously superior to those of the other models, and the slip-link model satisfactorily described the biaxial data using the fitted parameters with the physically reasonable magnitudes.

To assess further strictly a molecular entanglement theory, it is very important to survey the predictive capability for the biaxial data of rubbery networks with different entanglement densities. Breereton et al. analyzed the uniaxial elongation data of cross-linked polyethylenes with different cross-linking densities on the basis of the slip-link model, but their samples had a nonhomogeneous network structure due to cross-linking in the semicrystalline phase. The complicated structure of the samples as well as the analysis using only the uniaxial data made their assessment ambiguous. An appropriate way to control the entanglement density in the network is to end-link the precursor polymers in solution and vary the polymer concentration. Entanglement density in the end-linked network is significantly influenced by an overlapping degree of precursor chains before end-linking. In addition, the end-linking system enables us to estimate the parameters of the network structure such as the number of network chains and cross-links independently of mechanical testing, on the basis of a nonlinear polymerization theory with the data of the soluble fractions. This is advantageous relative to conventional randomly-crosslinked systems whose structural parameters are obscure.

In the present study, we assess further rigorously the slip-link model on the basis of the biaxial data of the end-linked PDMS networks with four different entanglement densities which are prepared from the melt and the solutions with three different precursor concentrations. In addition, we investigate how each of the best-fit model parameters depends on the degree of dilution at network preparation, i.e., entanglement density. The biaxial data of these four PDMS networks were originally employed in our separate study, where the form of $F$ was phenomenologically estimated using the two invariants of the deformation tensor as independent variables. This approach is entirely phenomenological, and the main objective is drastically different from that in the present study.

II. THE SLIP-LINK MODEL

The elastic free energy of the slip-link model consists of the two contributions originating from chemical cross-links and trapped entanglements. The expression of the former contribution due to chain connectivity is based on the Gaussian form which was derived by the classical rubber elasticity theory. The latter contribution due to trapped entanglements are modeled by a number $N_s$ of fictitious mobile slip-links which attach two entangled chains. The measure of slippage of slip-link, compared to that of a cross-link, is given by a parameter $\eta$. In the lower limiting case where the slippage becomes zero ($\eta=0$), the slip-link acts in the same way as the cross-link and fully contributes to the network modulus. In the upper limiting case where the slippage becomes infinitely large ($\eta=\infty$), the constraints by entanglements are extremely weak and the slip-link does not contribute to the network elasticity.

Edwards and Vilgis introduced the effect of finite network extensibility to the original slip-link model on the basis of the primitive path concept. The chain between entanglements is fully stretched long before the single chain between cross-links is taut when the network is deformed. As a result, the network extensibility is much lower than the extensibility of a single network chain between cross-links. The Edwards–Vilgis slip-link model has the singularity in the chain entropy at the ultimate elongation ratio $\lambda_{\text{max}}$ which is given by $\lambda_{\text{max}}=n_j^{1/2}$, where $n_j$ is the number of the Kuhn segments between adjacent elastically effective junctions. The limited network extensibility effect is introduced by the parameter $\alpha$ which is equal to $\lambda_{\text{max}}^{-1}$. The full expression of the free energy of the Edwards–Vilgis slip-link model is given by

$$\frac{F}{RT} = \frac{1}{2} N_s \left[ (1-a) \sum \lambda_i^2 \right] + \ln \left( 1-a^2 \sum \lambda_i^2 \right) \]$$

$$+ \frac{1}{2} \sum \lambda_i^2 \ln \left( 1 + \eta \lambda_i^2 \right) \]$$

$$+ \ln \left( 1 + \eta \lambda_j^2 \right) \]$$

where $\Sigma$ denotes the summation for $i$ from 1 to 3, and $\lambda_i$ is the principal ratio in the $i$th direction. The elastic contribution of network connectivity, $N_c$, is related to the number densities of the network chain ($n$) and cross-link ($\mu$),

$$N_c = (n - h \mu),$$

where $h$ is an empirical parameter varying from zero (affine limit) to unity (phantom limit) which depends on the degree of thermal fluctuation of the cross-link. In the case of $\alpha = 0$ (i.e., infinite extensibility), Eq. (2) coincides with the free energy of the original slip-link model. Furthermore, $\eta = \alpha = 0$ yields the classical free energy expressed by Eq. (1).

Theoretical stress–strain relation for a network biaxially stretched in the 1- and 2-directions ($\sigma_3 = 0$) is obtained using the following Treloar relations:

$$\sigma_i = \frac{2}{V \lambda_i^2} \left[ k_i \left( \frac{\partial F}{\partial \lambda_i^2} \right) \right]_{T,V} (i=1,2),$$

where $V$ is the volume of the network, and $\sigma_i$ is the nominal stress (the force per unit area of undeformed state) in the $i$th direction. The principal ratio $\lambda_3$ is automatically determined by the relation $\lambda_3 = \left( 1 + \lambda_1 \lambda_2 \right)^{-1}$ due to incompressibility of the rubbery network. The expression of $\partial F/\partial \lambda_i^2$ is given by
entanglement is treated as the localized additional cross-link. Four end-linked PDMS networks with different entanglement densities were prepared from melt and the solutions of the precursor concentrations 70, 50, and 30 wt. %, respectively. The details of the sample preparation are referred to our previous paper. After the end-linking reaction, the resulting networks were used for mechanical measurements without removing the diluent. The weight fraction of the unreacted reactants \( w_{\text{sol}} \) was evaluated by extraction in toluene. Volume fraction of the precursor chain incorporated into the network \( \phi_0 \) was calculated by subtracting \( w_{\text{sol}} \) from the initial precursor concentration.

The molecular parameters characterizing the network structure, i.e., number densities of elastic network chains \( (\nu) \) and cross-links \( (\mu) \), the number-average molecular mass of elastic chains between cross-links (excluding any attached dangling chains) \( (M_{n,el}) \), were estimated using a nonlinear polymerization model for end-linking with the \( w_{\text{sol}} \) data. The estimated values of these parameters for each sample are summarized in Table I.

As can be seen in Table I, the \( M_{n,el} \) values for the different networks are comparable, which indicates that the degree of dilution at network preparation alters the entanglement density of the resulting network without increasing significantly the network defects such as dangling chains and inelastic loops: If the degree of dilution increases the network defects, the resulting \( M_{n,el} \) values will not be constant against \( \phi_0 \). The larger values of \( M_{n,el} \) relative to \( M_{n,p} \) is due to the nonstochiometric ratio \((r \neq 1)\) as well as the finite values of \( w_{\text{sol}} \). Thus a main difference in network structure between the networks with different \( \phi_0 \) is that of entanglement density.

### III. CHARACTERISTICS OF THE END-LINKED PDMS NETWORKS

#### A. Uni- and biaxial elongation

The quasiequilibrium stress–strain data of uniaxial and biaxial deformations were obtained at 40 °C. The quasiequilibrium stress where the time effect was sufficiently eliminated was measured at each strain by the stress relaxation method. The details of the methods and instruments are described elsewhere.

#### B. Structural parameters

The end-reactive linear precursor PDMS used is long enough to form entanglement couplings before end-linking: The number-average molecular mass of the precursor \((M_{n,p} = 4.66 \times 10^4 \text{ g/mol})\) is much larger than the critical molecular mass to form entanglement couplings for PDMS \((M = 1.66 \times 10^4 \text{ g/mol})\). The molar ratio of functional groups in the precursor to those in the tetrafunctional silane (cross-linker) \((r)\) was 1.3 for all the samples. Oligomethylsiloxane with \( M = 3000 \text{ g/mol} \) was employed as a nonvolatile diluent.

Four end-linked PDMS networks with different entanglement densities were prepared from melt and the solutions of the precursor concentrations 70, 50, and 30 wt. %, respectively. The details of the sample preparation are referred to in our previous paper. After the end-linking reaction, the resulting networks were used for mechanical measurements without removing the diluent. The weight fraction of the unreacted reactants \( w_{\text{sol}} \) was evaluated by extraction in toluene. Volume fraction of the precursor chain incorporated into the network \( \phi_0 \) was calculated by subtracting \( w_{\text{sol}} \) from the initial precursor concentration.

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### IV. RESULTS AND DISCUSSIONS

Four parameters, \( h, N_s, \eta, \alpha \), are employed as adjustable parameters in the data-fitting procedure. The best-fit condition was determined so that the error \( \Delta \) would be minimum,

\[
\Delta = \frac{1}{2n} \sum_{i=1}^{n} \sum_{m=1}^{n} \left( \sigma_{i,\exp}^{m} - \sigma_{i,\text{th}}^{m} \right)^2 \right)^{1/2},
\]

where \( n \) is the number of experimental data points, \( \sigma_{\exp} \) and \( \sigma_{\text{th}} \) are experimental and theoretical stresses, respectively. The fitting procedure yields a unique parameter set which can reproduce the most successfully the experimental data. In the minimization of \( \Delta \), each parameter was independently varied in the physically admissible range. For \( N_s \) without the upper limit, the theoretical value of \( G_0 \) was used as a cutoff point, and the parameter set was selected so that the value of \( \Delta \) was the smallest.
condition: When the difference between the theoretical and experimental values of $G_0$ exceeds 10%, the corresponding value of $N_s$ is discarded before fitting to the stress–strain data.

Figures 1–4 display the comparison of the biaxial stress–elongation data with the best-fit theoretical predictions for the end-linked PDMS networks prepared from melt, 70, 50, and 30 wt. %, respectively. The uniaxial data shown by triangular symbols are also included in the figures. The solid lines represent the theoretical stress $\sigma_i$ at constant $\lambda_1$ as a function of $\lambda_2$. The best-fitted parameters are $N_s/N_c = 8.98$, $\eta = 0.105$, $\alpha = 0.178$, $h = 0$.

Fig. 1. Comparison of the theory with the quasiequilibrium biaxial stress–elongation data (Ref. 21) of the end-linked PDMS network of $f_0 = 0.914$. $\lambda_1$ and $\lambda_2$ are the larger and smaller principal ratios in biaxial stretching, respectively, and $\sigma_1$ (a) and $\sigma_2$ (b) are the nominal stresses in the corresponding directions. The triangular and rectangular symbols stand for the data for the uniaxial ($\lambda_2 = \lambda_1^{1/2}$ and $\sigma_2 = 0$) and equibiaxial stretching ($\lambda_1 = \lambda_2$), respectively. The solid lines represent the theoretical stress $\sigma_i$ ($i = 1,2$) at identical $\lambda_1$ as a function of $\lambda_2$. The best-fitted parameters are $N_s/N_c = 8.98$, $\eta = 0.105$, $\alpha = 0.178$, $h = 0$.

Fig. 2. Comparison of the theory with the quasi-equilibrium biaxial stress–elongation data (Ref. 21) of the end-linked PDMS network of $f_0 = 0.625$. $\lambda_1$ and $\lambda_2$ are the larger and smaller principal ratios in biaxial stretching, respectively, and $\sigma_1$ (a) and $\sigma_2$ (b) are the nominal stresses in the corresponding directions. The triangular and rectangular symbols stand for the data for the uniaxial ($\lambda_2 = \lambda_1^{1/2}$ and $\sigma_2 = 0$) and equibiaxial stretching ($\lambda_1 = \lambda_2$), respectively. The solid lines represent the theoretical stress $\sigma_i$ ($i = 1,2$) at identical $\lambda_1$ as a function of $\lambda_2$. The best-fitted parameters are $N_s/N_c = 7.68$, $\eta = 0.120$, $\alpha = 0.160$, $h = 0$.

points, although the error bars are not shown in the figures in order to avoid overlapping. The good agreements are also confirmed by the small $\Delta$ values: The differences between the data and the theoretical predictions are less than 8%. The theoretical values of $G_0$ calculated from Eq. (6) with the best-fit parameters, shown in Table II are also in good accordance with the experimental values listed in Table I. Thus the predictions of the slip-link model successfully fit the biaxial data as well as $G_0$ for the end-linked PDMS networks with different entanglement densities. We discuss below how each of the best-fit parameters depends on network concentration.

The best-fit value of $h$ was zero for all the networks. The condition $h = 0$ corresponds to full suppression of thermal fluctuation of cross-link (affine limit). Patel et al. reported $h = 0$ for the melt-end-linked PDMS networks irrespective of entanglement density on the basis of the dependence of $G_0$ on $\nu$, which accords with the result obtained here. According to the principle of end-linking, the molecular mass between
adjacent cross-links is unaltered by dilution at preparation. Consequently, $N_e$ is expected to vary linearly with $\phi_0$, when the degree of thermal fluctuation of cross-links ($h$) is not altered by dilution. In Fig. 5, the best-fit values of $N_e$ appear to depend linearly on $\phi_0$, which agrees with the expectation.

Figure 5 also shows the double logarithmic plots of $N_e$ vs $\phi_0$. The least square method yields $N_e \sim \phi_0^{2.4}$. Entanglement coupling (slip-link) originates from the binary interaction, which suggests that the $\phi_0$ dependence of $N_e$ obeys a power law with an exponent of two or about 2.3. The experimental exponent is slightly larger but close to the expected values. The ratio $N_e/N_c$ represents the elastic contribution of entanglements relative to that of cross-links. As can be seen in Table II, $N_e/N_c$ decreases with decreasing in $\phi_0$ as a result of the reduction in entanglement density. The large values of $N_e/N_c$ for the networks of $\phi_0 \geq 0.625$ indicate that the networks are entanglement-dominated. Actually, the magnitude of $G_0$ of the melt-ended network is reasonably explained by the quasiequilibrium modulus of uncrosslinked entangled PDMS melt $G_0^0 = 2 \times 10^5 \text{ Pa}$ (Ref. 30) together with the consideration that the unreacted reactants act as diluent in network preparation, $G_0^0(0.92) \sim 1.6 \times 10^5 \text{ Pa}$.

Figure 6 illustrates the best-fit values of $\eta$ as a function of $\phi_0$. As $\phi_0$ decreases, i.e., entanglement density decreases, $\eta$ becomes larger. This tendency accords with the intuitive expectation from the physical meaning of $\eta$: $\eta$ is a measure of slippage for slip-link. An increase in the distance between neighboring entanglements increases the freedom of the motion of slip-link. The quantitative relation of $\eta$ with network structure is unknown at present, but Vilgis et al. proposed the following rough estimate:

$$
\eta \approx M_c/M_e,
$$

where $M_e$ and $M_c$ are the molecular weights between neighboring entanglements and cross-links, respectively. We em-
ploy the following two methods for the estimation of $M_c/M_e$. As is evident from the definitions, $M_c/M_e$ is equivalent to $N_c/N_s$.

$$\eta = M_c/M_e = N_c/N_s. \tag{9}$$

Table II tabulates the $\eta$ values calculated from Eq. (9) with the best-fit values of $N_c$ and $N_s$. In the other approach, $M_c/M_e$ was evaluated using $M_c = M_{e,melt} \phi_0^{-1}$ (Ref. 28) and $M_c = M_{n,el}$ as

$$\eta = M_c/M_e = M_{e,melt} \phi_0^{-1}/M_{n,el}. \tag{10}$$

The $\eta$ values calculated by Eq. (10) and $M_{e,melt} = 10,000$ g/mol for PDMS (Ref. 30) are listed in Table II. The values estimated from Eqs. (9) and (10) are slightly different but comparable. The agreement in $\eta$ between the best-fit values and the estimates from Eq. (9) or (10) is more or less tolerable in view of the accessible range of $\eta (0 < \eta \ll \infty)$ as well as the roughness included in the estimation.

The $\phi_0$ dependence of the best-fit values of $\alpha$ is shown in Fig. 7. The $\alpha$ values decrease with decreasing in $\phi_0$, i.e., entanglement density. This trend is physically reasonable because $\alpha$ is a measure of inextensibility ($\alpha = 1/\lambda_{\text{max}}$): A decrease in the number of entanglements increases the length of the network strand between neighboring entanglements, which increases the extensibility of network. The primitive path concept\(^7\) derives $\lambda_{\text{max}} = N_j^{1/2}$, where $N_j$ is the number of the Kuhn segments between topologically adjacent cross-links or entanglements where cross-link and entanglement are not distinguished. The expression for $\alpha$ is obtained by using the relation between $N_j$ and $G_0$.

$$\alpha \approx \left( \frac{1}{N_j} \right)^{1/2} = \left( \frac{G_0 m p}{c R T} \right)^{1/2}, \tag{11}$$

where $m$ is the molecular mass of a repeating unit, $p$ is the number of the repeating unit per one Kuhn segment, and $c$ is the network concentration. For PDMS, $m = 74$ g/unit and $p = 8.5$.\(^3\) The values of $\alpha$ calculated from Eq. (11), shown in Table I, are fairly close to the best-fit values. The accordance between the best-fit values and the estimates from Eq. (11) is considered satisfactory in view of the accessible range of $\alpha (0 < \alpha \ll \infty)$.

Thus the slip-link model satisfactorily describes the biaxial stress–strain data of the end-linked PDMS networks with different entanglement densities. Each of the best-fit model-specific parameters ($N_c, N_s, \eta, \alpha$) depends on net-

### Table II. Best-fit parameter values and estimated values from molecular approaches.

<table>
<thead>
<tr>
<th>$\phi_0$</th>
<th>$\Delta^* %$</th>
<th>$h$</th>
<th>$N_c$ (mol m$^{-3}$)</th>
<th>$N_s$ (mol m$^{-3}$)</th>
<th>$\eta$</th>
<th>$\eta_{\text{est}}^b$</th>
<th>$\eta_{\text{est}}^c$</th>
<th>$\alpha$</th>
<th>$\alpha_{\text{est}}^d$</th>
<th>$G_{0,\text{cal}}/ \mathrm{kPa}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.914</td>
<td>7.84</td>
<td>0</td>
<td>5.88</td>
<td>52.8</td>
<td>8.98</td>
<td>0.105</td>
<td>0.111</td>
<td>0.141</td>
<td>0.178</td>
<td>0.199</td>
</tr>
<tr>
<td>0.625</td>
<td>3.58</td>
<td>0</td>
<td>3.22</td>
<td>24.7</td>
<td>7.68</td>
<td>0.120</td>
<td>0.130</td>
<td>0.194</td>
<td>0.160</td>
<td>0.161</td>
</tr>
<tr>
<td>0.463</td>
<td>4.10</td>
<td>0</td>
<td>3.02</td>
<td>7.91</td>
<td>2.62</td>
<td>0.169</td>
<td>0.293</td>
<td>0.134</td>
<td>0.116</td>
<td>24.9</td>
</tr>
<tr>
<td>0.283</td>
<td>6.75</td>
<td>0</td>
<td>2.12</td>
<td>3.33</td>
<td>1.57</td>
<td>0.799</td>
<td>0.637</td>
<td>0.510</td>
<td>0.125</td>
<td>0.0880</td>
</tr>
</tbody>
</table>

\(a\) Defined by Eq. (7).
\(b\) Estimated from Eq. (9).
\(c\) Estimated from Eq. (10).
\(d\) Estimated from Eq. (11).
\(e\) Calculated from Eq. (6) with the best-fit parameter values.
\(f\) The values of the best-fit parameters for the sample are slightly different from those published previously (Ref. 11) due to the difference in the best-fit procedures (Ref. 26).
work concentration (or entanglement density) in a physically reasonable way. In particular, the magnitudes of the best-fit values of \( \eta \) and \( \alpha \) satisfactorily agree with the estimates from another molecular approaches independent of mechanical testing. These results suggest that the slip-link model successfully accounts for the chain-entanglement effects on rubber elasticity.

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27. In the previous paper (Ref. 11) the root of mean square (rms) \((1/2n) \times (\sum_i (\sigma_i - \sigma_{r,exp})^2)^{1/2}\) was minimized for data-fitting. Due to the difference in the rms and \( \Delta \) defined by Eq. (7), the best-fit parameter values for the network of \( \phi_0 = 0.625 \) are slightly different from those reported previously (Ref. 11). The rms tends to weight \( \sigma_1 \) relative to \( \sigma_2 \) due to the fact of \( \sigma_1 \approx \sigma_2 \) under any biaxial strains. This tendency is eliminated in \( \Delta \) where the difference \( |\sigma_i - \sigma_{r,exp}| \) is reduced by the stress \( \sigma_{r,exp} \) (i = 1, 2).