Time-Dependent Behavior of a Graphite/Thermoplastic Composite and the Effects of Stress and Physical Aging

Thomas S. Gates¹ and Mark Feldman²

ABSTRACT: Experimental studies were performed to determine the effects of stress and physical aging on the matrix dominated time-dependent properties of IM7/8320 composite. Isothermal tensile creep/aging test techniques developed for polymers were adapted for testing of the composite material. Time-dependent transverse ($S_T$) and shear ($S_S$) compliance's for an orthotropic plate were found from short-term creep compliance measurements at constant, sub-$T_g$ temperatures. These compliance terms were shown to be affected by physical aging. Aging time shift factors and shift rates were found to be a function of temperature and applied stress.

KEYWORDS: physical aging, composite materials, viscoelasticity, creep compliance, superposition

Tensile Creep Compliance

The experimental study centered on using momentary creep compliance tests to determine the aging behavior of the principal compliance terms for a laminated composite as a function of stress and temperature. Using a three-parameter model of creep compliance, aging shift factors and shift rates were investigated for three temperatures below the $T_g$. Isothermal, elevated temperature creep tests provided the short-term creep compliance data. To establish stress levels for the aging tests, short-term, isothermal creep tests were run at increasingly higher stresses. In addition, creep/recovery tests provided insights into the effects of short versus long recovery periods and the applicability of Boltzman’s superposition principle [5]. Other tests examined the effects of stress on creep compliance while aging to equilibrium.

Analytical Models

The analytical modeling provided a means of describing changes due to physical aging in creep compliance of a laminated plate.
For the laminates in this study, the usual axes' definitions for an orthotropic plate were used, for example, $1 = $ along the fiber, $2 = $ transverse to the fiber, $x = $ axial loading direction, $y = $ transverse to loading direction, and the angle between the 1,2 axes and the $x - y$ axes was measured clockwise.

The general case of a thin orthotropic plate was considered. For a state of plane stress parallel to the $x = y$ plane in an orthotropic solid, the constitutive relation can be given by

$$
\begin{bmatrix}
\varepsilon_{11} \\
\varepsilon_{22} \\
\gamma_{12}
\end{bmatrix} =
\begin{bmatrix}
S_{11} & S_{12} & 0 \\
S_{21} & S_{22} & 0 \\
0 & 0 & S_{66}
\end{bmatrix}
\begin{bmatrix}
\sigma_{11} \\
\sigma_{22} \\
\sigma_{12}
\end{bmatrix}
$$

where $\varepsilon$ and $\gamma$ are the strains, $S_{ij}$ is the compliance, and $\sigma$ is the stress.

This set of equations implies that there are four independent constants needed to characterize the material. These constants are the longitudinal and transverse compliance, $S_{11}$ and $S_{22}$, respectively, Poisson's ratio $v_{12}$, and the shear compliance $S_{66}$. A primary assumption was that physical aging occurred only in the polymer-matrix constituent. On the basis of tests performed by Sullivan [2] and Hastie and Morris [3], an assumption was that $S_{22}$ and $S_{66}$ were the two compliance terms to be affected by physical aging.

Creep tests provided the means for assessing the time-dependent compliance changes in both the $S_{22}$ and $S_{66}$ terms. These creep tests were analyzed using linear viscoelastic theory and standard sub-Tg creep/aging data reduction procedures. The term linear viscoelastic behavior implies that the principles of proportionality and superposition were both met. These conditions were checked experimentally and the results are given in following sections. The authors recognize that the polymer-matrix material may exhibit nonlinearity, particularly at elevated temperatures. However, it was felt that the simplicity of linear viscoelasticity made it desirable enough to investigate the use of it as a initial modeling scheme.

For a short-term (less than 10 h) creep test, the $log$ compliance versus $log$ test time data for both $S_{22}$ and $S_{66}$ tests will appear as shown in the example provided by Fig. 1. A three-parameter expression was chosen to model the test data

$$
S_{ij}(t) = S_{ij}^0 e^{(t / \tau) \beta}
$$

where $S_{ij}^0$ is the time-dependent creep compliance term from Eq 1, $S_{ij}$ is the initial creep compliance, $t$ is time, $\tau$ is the characteristic retardation time, and $\beta$ is the shape parameter. The fitting algorithm was based upon the Levenberg-Marquardt method [6]. Equation 2 has been used successfully by Struik [1] to model a wide range of polymers. The contribution of each parameter to the shape of the curve is well understood. For the continuous fiber-reinforced composite, the individual terms can be written

$$
S_{22}(t) = S_{22}^0 e^{(t / \tau_2) \beta_2}
$$

The short-term creep compliance curves, as shown in Fig. 1, had a horizontal or time shift between curves defined by $log \ a$, and the vertical or initial compliance shift between curves defined as $log \ b$. Figure 1 schematically illustrates these shifts. It was assumed that both shifts were due to physical aging effects.

Once the time shifts ($log \ a$) were measured for a given set of compliance tests, the aging shift rate ($\mu$) was calculated using

$$
\mu = \frac{d \ log(a_\tau)}{d \ log(\tau)}
$$

where $\tau_\text{is}$ the specific aging time for a given test. Struik [1] indicates that for sub-$T_g$ temperatures, the shift rates would be constant over the relatively short aging time of a test sequence and would have a value close to unity. By raising the test temperatures sufficiently close to $T_g$, Struik [1] and Lee and McKenna [7], found the shift rate of an aging test decreased significantly below unity. This was attributed to the material approaching an equilibrium state. The time required to reach this equilibrium state is dependent on the test temperature. Lee and McKenna [7] have stated that this equilibrium time was not dependent upon the state of stress. An empirical formula for estimating this equilibrium time ($t_\infty$) for polymers was given by Struik [1] to be

$$
t_\infty = 100 e^{787 (T_g - T)}
$$

where $T_g$ is the glass transition of the test material, $T$ is the test temperature in degrees C, and $t_\infty$ is in seconds.

Test Material and Specimen Configuration

The composite material used in this study was a continuous carbon-fiber-reinforced amorphous thermoplastic fabricated by AMOCO and designated IM7/8320 (formally IM7/RadelX). Static lamina properties measured isothermally and given previously [8] are repeated for reference purposes in Table 1. The $T_g$, as measured by a differentiating scanning calorimeter (DSC), was 221.3°C. The static material property tests were run using a rectangular specimen geometry of approximately 24 by 2.5 cm with a 12- or 8-ply thickness and a ply thickness of approximately .0135 cm. A set of thin end tabs were added to each end of the specimen. For the creep tests, the transverse ($S_{22}$) and shear ($S_{66}$) compliance data came from [90]$_1$, and [± 45]$_b$ specimens, respectively.

<table>
<thead>
<tr>
<th>Temperature, °C</th>
<th>$E_1$, MPa</th>
<th>$E_2$, MPa</th>
<th>$G_{12}$, MPa</th>
<th>$v_{12}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>150</td>
<td>152.9</td>
<td>7.3</td>
<td>4.4</td>
<td>0.3</td>
</tr>
<tr>
<td>175</td>
<td>153.9</td>
<td>7.2</td>
<td>3.4</td>
<td>0.3</td>
</tr>
<tr>
<td>200</td>
<td>147.0</td>
<td>5.5</td>
<td>2.6</td>
<td>0.3</td>
</tr>
</tbody>
</table>

FIG. 1—Schematic of typical creep compliance data from a sequenced aging test.
**Test Procedures and Equipment**

Before testing, all specimens were dried for at least 15 h at 110°C in a convection oven. After drying, the specimens were stored in a disector until the start of the testing. Any thermal exposure prior to the start of testing did not affect the data since the specimen was rejuvenated immediately prior to testing (see following sections). After each test sequence, the specimens were inspected for matrix cracks along their edges with an optical microscope.

Three test temperatures selected for study were 195, 200, and 212°C. The two lower temperatures provided data near the upper use temperature of the material. The highest test temperature was selected to ensure that the aging to a point near equilibrium would occur within a reasonable amount of time. For IM7/8320, which has a Tg of 221.3°C, the aging time to equilibrium based upon Eq 5 would vary from 35.2 h for the upper test temperature of 212°C to $17.3 \times 10^6$ h for the lower test temperature of 195°C.

All of the creep tests were performed in convection ovens equipped with digital controllers. Temperature monitoring and feedback thermocouples were placed near the test section. Thermal apparent strain was corrected for by using the compensating gage technique [9]. Load was applied through a dead-weight cantilever arm and reacted at a point outside the test chamber. Mechanical wedge grips held the specimen during the loaded or creep segments. Stress was calculated based upon the applied load and the specimen cross section measured before testing. During the unloaded or recovery segments, the lower grip was released using a remote cable and pulley system. This mechanism allowed the test chamber to remain closed during the entire test sequence.

Strain in the gage section was measured with high-temperature foil strain gages applied in the center of the specimen. The gage type was Micro-Measurement WK-00-250BG-350 with the Mbond-600-gage adhesive. Proper selection of gage type and adhesive gave coefficient of thermal expansion match and stability at elevated temperatures.

For the [90]s specimens, the average measurement of two back-to-back gages, aligned longitudinally, were used to compute $S_{22}$. For the $[\pm 45]$s specimens, the average measurement in each direction of four gages, two back-to-back aligned longitudinally and two back-to-back aligned transversely, were used to compute $S_{66}$. The compliance terms for these specimens are as follows

$$S_{22}(t) = \frac{\epsilon_2(t)}{\sigma_2}$$

$$S_{66}(t) = \frac{2\epsilon_6(t)[1 - \epsilon_6(t)/\epsilon_6(t)]}{\sigma_6}$$

(6)

Each gage formed a quarter-bridge circuit. Thermal strain compensation was accomplished during data reduction. Commercially available instrumentation provided bridge completion, excitation, and signal conditioning. A personal computer equipped with a 12-bit A/D board converted and stored the data.

**Creep Compliance and Physical Aging** — A procedure for measuring creep compliance, as described in Struik [1], was used for all tests. This procedure consisted of a sequence of isothermal creep and recovery tests. Each creep period was short in comparison with the previous aging time. During the aging process, the short-term creep tests required a constant load. After each creep segment, the specimen was unloaded and allowed to recover until the start of the next creep test. Figure 2 is a schematic of this procedure.

Table 2 provides the temperatures and applied axial stress for each test sequence.

As part of the test procedure, immediately before the start of any physical aging test sequence, the specimen was held at 235°C (approximately 15 degrees above it’s $T_g$) for 15 min and then rapidly quenched with compressed air to the test temperature. This excursion above the $T_g$ rejuvenates the specimen and erases any prior physical aging history. After reaching the test temperature during the quench, the aging time started.

The ratio between the prior aging time and each creep test time was at least 10:1 for the sequenced tests. Using this ratio and the sequencing procedure, up to seven creep tests were conducted during an aging time of approximately 120 h. One point of concern for such sequenced tests is the effects of repeated loading on the aging process. To check out such loading effects, sequenced creep tests were run for 27 h aging (three loading segments). The 27-h sequenced compliance data was then compared to data from a specimen with the same 27-h aging time but without the intermediate creep segments.

During the creep recovery segments of each aging test sequence, the specimen had no load on it. The recovery times were long enough to ensure creep compliance relaxation was complete by the start of the next creep test. Table 2 provides the temperatures and applied axial stress for each test sequence.

<table>
<thead>
<tr>
<th>Test Temperature</th>
<th>$\sigma_2$, MPa</th>
<th>$\tau$, s</th>
<th>$\beta$, $\mu$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$S_{22}$</td>
<td>195</td>
<td>3.04</td>
<td>0.128</td>
</tr>
<tr>
<td>$S_{195}$</td>
<td>195</td>
<td>3.72</td>
<td>0.126</td>
</tr>
<tr>
<td>$S_{22}$</td>
<td>195</td>
<td>4.39</td>
<td>0.127</td>
</tr>
<tr>
<td>$S_{195}$</td>
<td>195</td>
<td>5.07</td>
<td>0.129</td>
</tr>
<tr>
<td>$S_{22}$</td>
<td>200</td>
<td>2.32</td>
<td>0.138</td>
</tr>
<tr>
<td>$S_{200}$</td>
<td>200</td>
<td>2.99</td>
<td>0.134</td>
</tr>
<tr>
<td>$S_{22}$</td>
<td>200</td>
<td>3.65</td>
<td>0.133</td>
</tr>
<tr>
<td>$S_{66}$</td>
<td>195</td>
<td>5.18</td>
<td>0.239</td>
</tr>
<tr>
<td>$S_{195}$</td>
<td>195</td>
<td>5.61</td>
<td>0.237</td>
</tr>
<tr>
<td>$S_{200}$</td>
<td>200</td>
<td>3.49</td>
<td>0.244</td>
</tr>
<tr>
<td>$S_{200}$</td>
<td>200</td>
<td>4.48</td>
<td>0.269</td>
</tr>
<tr>
<td>$S_{200}$</td>
<td>200</td>
<td>5.47</td>
<td>0.257</td>
</tr>
</tbody>
</table>

**FIG. 2—Schematic of creep/recovery sequence for an aging test.**
enough to allow for nearly complete strain recovery. However, to account for any remaining strain, the strain measured in the creep segment was corrected by subtracting the extrapolated prior recovery curve from the creep curve, as shown in Fig. 2. To further investigate the effects of short versus long recovery times, two sets of sequenced creep/recovery tests were run with identical aging times but different creep/recovery times. Figure 3 is a schematic illustrating these two test schedules.

Thermoreversibility of Physical Aging—Due to the supposed thermoreversibility of the physical aging process, a single specimen could be used for several aging tests as long as the rejuvenation occurred before each test sequence. This reversibility was verified by measuring creep compliance from a single specimen loaded several times with a rejuvenation sequence before each loading.

Determination of Applied Stresses—For linear viscoelastic behavior, it was assumed that superposition and proportionality would hold. Given an initial state of stress $\sigma'$ applied for a time $t_0$ and an additional stress $\sigma''$ applied at a time $t$, superposition implies that

$$
e[\sigma'(t) + \sigma''(t - t_0)] = \epsilon[\sigma'(t)] + \epsilon[\sigma''(t - t_0)]$$  \(7\)

while proportionality states that for an applied stress $\sigma$, the strain in a material at any other stress state is found using

$$\epsilon[c\sigma(t)] = ce[\sigma(t)] \quad \text{where} \quad c = \text{constant}$$  \(8\)

Creep and creep/recovery data from several loadings at different stress levels provided data for checking superposition. Proportionality was checked by plotting isothermal, isochronous stiffness versus applied stress for a specimen that was repeatedly rejuvenated, quenched, and loaded at increasingly higher stress levels. The supposed transition from linear to nonlinear behavior should be evident by a slope change in the data.

Results: Tensile Creep Compliance

Results from the experiments illustrate the effects of stress and physical aging on the elevated temperature creep compliance of IM7/8320. Superposition, proportionality, sequencing effects, load duration effects, stress overload effects, aging to equilibrium, aging shift factors, and aging shift rates were measured and quantified.

Superposition—For linear viscoelastic behavior, superposition should hold. For the creep test, superposition states that the recovery strain can be found using Eqs 2 and 7 such that

$$\epsilon(t^*) = \sigma S_o \{e^{\alpha \beta} - e^{\alpha \beta - t^*} \}$$  \(9\)

where $\sigma$ is the constant stress, $t_0$ is the time of load removal, $t^*$ is the time during the recovery segment, and $S_o$, $\alpha$, and $\beta$ are the constants from the measured creep compliance curve. Equation 9 was used to predict the recovery of a $[90]_t$ transverse specimen loaded at sequentially higher load levels. The specimen was rejuvenated before each test and aged for exactly 30 min before loading. Figure 4 provides the predicted strain/time curves along with the measured data.

Comparison of test and prediction for the recovery segments indicates that only the recovery test data at the lowest load level was accurately predicted by superposition. However, the predicted strain rate during recovery corresponded well with the measured value for all load levels tested.

Proportionality—One of the clearest ways to check if proportionality holds for creep tests is to plot the isochronous stiffness versus applied stress levels at a constant temperature. The stiffness values were calculated by inverting the measured creep compliance data for any given time during the loaded segment of the test. Therefore, it should be noted that the stiffness values were not true stiffness as calculated from stress-relaxation tests. Unproportional behavior will be evident by deviations of the data from horizontal lines as stress increases. Figures 5 through 8 give examples of such data.

Figures 5 and 6 indicate little or no unproportional behavior as a function of applied stress, whereas Figs. 7 and 8 show a transition to unproportional behavior. Comparing Figs. 5 through 7 shows that these deviations from linearity became more apparent as the test temperature increased. The stress levels chosen for the aging experiments are indicated on these figures. Using the isochronous stiffness data from Figs. 5 through 8, the aging test stress levels were chosen to correspond to the range where little or no deviation from proportionality was observed.

Sequencing Effects—To ensure that the sequencing procedures of the momentary creep tests worked, checks were made using different specimens and test temperatures. For the sequencing to work properly, any creep compliance data from the sequenced
creep/aging experiment should coincide with an unsequenced creep compliance measurement taken at the same aging time. Figure 9 shows an example of this behavior for a [±45]$_{3s}$ specimen loaded at 1, 3, 9, and 27 h of aging time versus the same specimen loaded only at 27 h aging. Both tests included a rejuvenation period before the start of aging.

Examination of the compliance data in Fig. 9 indicates that the sequenced tests produced equivalent 27 h compliance data compared to the unsequenced test. This demonstrates that the short loading periods of the sequenced tests did not measurably effect the data, thereby ensuring that the sequenced test procedure gave a true measure of physical aging.

Load Duration Effects—The momentary creep compliance measured during the aging tests was essentially a "snapshot" of the aging process. This required that the duration of a creep segment be kept much lower than the cumulative aging time, thereby ensuring minimal influence on the aging process moreover allowing for a sufficient time for strain recovery to occur. The data from all the strain recovery segments of the sequenced creep compliance measurements indicated that near complete recovery took place. As suggested by Struik [1], the unrecovered strain was subtracted from the total strain during the following creep segment. The unrecovered strain was computed using a linear extrapolation of the steady-state portion of the creep recovery curve. Figure 2 schematically illustrates this procedure.

To explore the effects of load duration on creep compliance, two different test schedules were followed. Figure 3 gives the times for the creep and recovery segments of these two test schedules. The test sequence with the short creep segments was designated Schedule 1 and the test sequence with the long creep segments was designated Schedule 2. Figure 10 shows the measured effects on creep compliance due to loading according to these schedules.

Figure 10 clearly illustrates that the extended creep times used in Schedule 2 produce different creep compliance curves than the data from Schedule 1. The differences are due to the additional creep occurring during the extended load time. If the curves from both schedules are fit with Eq 2, different parameters will result. However, the effects of aging (log a shifts) are apparent using either test schedule. For material systems unlike IM7/8320 that would not exhibit such rapid strain recovery, the effects of extended load periods would become even more pronounced.

Stress Overload Effects—As outlined by Struik [1], a momentary stress overload during a normal sequenced creep compliance test

\[ \log S(t) \]

\[ (1 / \text{GPa}) \]

\[ \text{IM7/8320 [±45]}_{3s} \]

200°C

Sequenced aging at: 1 hr.
3 hr.
9 hr.
27 hr.

Aged for 27 hrs.

FIG. 9—Comparison of sequenced versus unsequenced creep compliance test data.
for aging may upset the aging process in the material. He states that if this overload is sufficiently high, the material may apparently deage. To explore this idea, a sequenced creep compliance test was run on a [90]_12 transverse specimen at 200°C. Figure 11 shows the load/time history imposed on this specimen. For the first three segments of the sequenced test, the load was kept at a constant. The aging time for these segments was 1, 3, and 9 h. The forth segment occurred at 24 h and consisted of an overload that was approximately 1.7 times the previous load. The initial load was then used for the subsequent segments. To capture any immediate or transient effects on the aging process, the segments after the overload started at 1, 3, 9, and 26 h after the overload or 26, 28, 34, and 51 h total aging time, respectively.

Figure 12a gives the resulting creep compliance curves. The 1-, 3-, 9-, and 24-h aging time segments collapsed to form one master curve referenced to the 1-h curve. The 26-, 28-, 34-, and 51-h aging time segments collapsed to form a master curve referenced to the 26-h curve. Apparently these curves are distinct in both shape and location.

Figure 12b gives the resulting shift factors. All the factors are first plotted using the 1-h curve as a reference for all the data. As shown in the Fig. 12b, a slight disruption of the data occurs after the overload. However, deviations from linearity of the shift rate µ are not significant. Fitting all the data gave a shift rate of .90. If just the post-overload shift factors are plotted referenced to the 26-h test, the shift rate decreased to .30.

**Aging to Equilibrium**—An experimental study on the ability to age the material to a state near equilibrium was performed using S_{22} specimens tested at 212°C and four stress levels. Figures 13 and 14 give the shift factors for these tests. Shift factors used in Fig. 13 are from using only the horizontal time shift to create the master curves. Figure 14 shows results from the same tests but with the time shifts computed after a vertical compliance shift. A bilinear least squares fit to the data was computed for all four data sets and is shown in Figs. 13 and 14. The break in these bilinear fits was made at the 3-h aging point. Additional data points before and after this 3-h point would further define these break points. The 3-h data point for the 385 psi (2654.5 kPa) data set was not included in the curve fit due to excessive scatter.
A comparison of Figs. 13 and 14 show that inclusion of the vertical shift in the collapse of the data to form the master curve may decrease the scatter of the aging factors. However, the general trends in the data do not change by including this vertical shift.

Aging Shift Factors—Given a set of momentary creep compliance curves, as shown in Fig. 1, the time shift factors ($\log a$) for each curve were measured by referencing all curves to the first (1-h) creep compliance curve. To facilitate computation of these shift factors, each curve was fit with the three-parameter model shown in Eq 2. The smooth curves generated from these fits were shifted along the time axis until they coincided with the reference curve and therefore created a momentary master curve. The master curve data was also fit with Eq 2. Table 2 gives the three parameters from the analysis of the master curve data for the 195 and 200°C sequenced tests. Figures 15 through 18 provide the master curves for $S_{22}$ and $S_{66}$.

For many of the tests, a vertical or compliance shift would facilitate the collapse of the data. However, the time shifts due to aging were of primary importance in this study; therefore, all the shift factor and shift rate curves shown in the results were found using the time ($\log a$) shifts only, unless otherwise noted.

Figures 19 through 22 present the time shifts versus aging time. In these figures, a linear least squares fit to each data set is also shown. Each linear fit represents a different applied stress level. Figures 19 and 20 are the $S_{22}$ data at 195 and 200°C, respectively. Figures 21 and 22 are the $S_{66}$ data at 195 and 200°C, respectively. All the data presented in these figures have a data point at the origin.

Aging Shift Rates—The aging shift rates for all tests were computed by using linear least squares fits to the aging shift factors. Tables 2 and 3 give the computed shift rates. Figure 23 is a plot of these shift rates as a function of applied axial stress. The $S_{66}$ data at 195 and 200°C collapsed and was fit with one curve. The $S_{22}$ data sets for each temperature were separate and fit with their own curves. The shift rates shown in Fig. 23 for the $S_{22}$ specimen at 212°C are values from the secondary slope of the bilinear fit.
These rates were computed using shift factors found from the aging time shift to the data.

Figure 24 uses the same shift rate data as in Fig. 23, but shows it as a function of ply stress for the $S_{22}$ and $S_{66}$ specimens. The transverse stress $\sigma_{22}$ in the 90° ply of the $S_{22}$ specimen is equal to the applied axial stress. The shear stress $\sigma_{12}$ in a 45° ply of the $S_{66}$ specimen has a magnitude equal to one half of the applied axial stress. The 195°C, $S_{66}$ specimen tested at 960 psi (6619 kPa)

<table>
<thead>
<tr>
<th>$\sigma_{12}$, psi</th>
<th>$\mu^a$</th>
<th>$\mu^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>200</td>
<td>0.132</td>
<td>0.239</td>
</tr>
<tr>
<td>256</td>
<td>0.114</td>
<td>0.147</td>
</tr>
<tr>
<td>350</td>
<td>0.091</td>
<td>0.147</td>
</tr>
<tr>
<td>385</td>
<td>0.065</td>
<td>0.147</td>
</tr>
</tbody>
</table>

$^a \mu^a$ = horizontal shift rate using horizontal and vertical shift.

$^b \mu$ = horizontal shift rate using horizontal shift only.
Shift Rate

FIG. 23—Measured aging shift rates for all tests versus the applied stress.

Shift Rate

FIG. 24—Measured aging shift rates for all tests versus the applied stress.

was the only specimen in which damage (transverse matrix cracks) was detected during post-test inspection. This data point was not used in the curve fit of Fig. 23 or 24.

Discussion and Summary: Tensile Creep Compliance

Linear viscoelasticity assumes that both proportionality and superposition hold. Data from Figs. 4 through 8 reveals the difficulty in checking these concepts for elevated temperature PMCs. The inability of superposition to accurately predict strain recovery for all stress levels indicates some stress-level dependency. However, the proportionality data suggests that there should be a range of stress levels available for testing in the linear viscoelastic range. For temperatures sufficiently close to the $T_g$, PMCs may exhibit nonlinear behavior in the matrix constituent. The lack of a sharp transition from linear to nonlinear behavior would suggest that if test stress levels are low, linear viscoelasticity may provide a first level approximation when developing test and analysis procedures.

The sequenced creep/recovery procedures seem to work well for characterizing physical aging in PMCs. It appears necessary to keep the ratio between aging time and test time to a value of ten or greater to ensure that loading has a minimum influence on aging and that the material has sufficient time to undergo strain recovery.

The tests on the $S_{22}$ specimen at 212°C indicated that if the temperature was sufficiently close to the $T_g$, a change in the aging shift rate may occur. Figures 13 and 14 indicate that these breaks in shift rate, or time-to-equilibrium, are relatively independent of applied stress level. This result has important implications for long-term testing. The aging towards equilibrium behavior for the IM7/8320 material was similar to the results achieved by Lee and McKenna [7] for epoxy glasses.

Momentary master curves do not reveal a clear trend when examined as a function of stress level. Examining Figs. 15 through 18 show that, in general, the master curves will shift to the right along the time axis as stress level increases. However, the data does not appear to lend itself well to a time-stress superposition type of analysis. The plots of shift factor versus aging time given in Figs. 19 through 22 verify the trend in shift factors as a function of applied stress.

Plotting shift rate $\mu$ as a function of applied axial stress, as in Fig. 23, indicates that $\mu$ for the two creep compliance terms have different aging shift rates. The $S_{22}$ term shows a higher dependency on applied axial stress. In addition, the $S_{22}$ data shows some temperature dependency when compared to the $S_{66}$ data. This difference in temperature dependency between the two compliance terms is probably due to the low creep strains of the $S_{22}$ data and the inability of the instrumentation to accurately resolve strains less than 10 $\mu e$. The $S_{66}$ data gave less scatter at both test temperatures.

Struik [7] has shown that the shift rate $\mu$ of rigid PVC loaded under two separate loads, tensile and shear, will collapse when plotted against applied stress. This stress was assumed to be large and to cause nonlinear creep. He further states that the collapse of the $\mu$/stress data appears to be independent of test temperature. Plotting the shift rates from Table 2 as a function of ply stress, as in Fig. 24, indicated the data for the $S_{22}$ and $S_{66}$ specimens may collapse into a single data set where the temperature and load-type dependency are less apparent.

Struik [7] has interpreted this stress-level dependency in polymers as an indication that large stresses produce an erasure or deaging of the material. Lee and McKenna [7] have offered an alternative interpretation of the decrease in shift rate as a function of an increase in the applied stress. They state that “an increase in the amplitude of the stress applied in the physical aging experiment results in a decrease in the shift rate $\mu$, simply because the changes in structure accompanying volume recovery affect the non-linear response differently at large stresses than at small ones.” They further state that this does not imply that the structure of the polymer is changed by the application of mechanical loads. At this point, the authors feel that additional data on the composite material is needed to fully validate either of the theories presented.

Acknowledgment

The authors would like to acknowledge the expert technical laboratory assistance of Ms. Lois White and Mr. Reggie Rose.

References


