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Dipolar Relaxation in a Second-Order Nonlinear Optical Interpenetrating Polymer Network

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**Abstract**
The nature of the relaxation process of the poled order in an interpenetrating polymer network (IPN) system is found to be fundamentally different from that of a guest/host system. The IPN (T_g = 176 °C) consists of a nonlinear optical (NLO) active epoxy-based polymer network and an NLO active phenoxy-silicon polymer network. The decay behavior of the second-order nonlinearity of this IPN was investigated in the range from 110 to 170 °C. The stability of this IPN is superior to those of classic guest/host polymers as indicated by longer relaxation times at temperatures from 110 to 130 °C. The relaxation process of the IPN system follows Arrhenius type behavior at temperatures ranging from 140 to 170 °C. The IPN system provides a new approach to processing for stabilization of the second-order nonlinear optical properties.
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Dipolar Relaxation in a Second-Order Nonlinear Optical Interpenetrating Polymer Network

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Abstract

The nature of the relaxation process of the poled order in an interpenetrating polymer network (IPN) system is found to be fundamentally different from that of a guest/host system. The IPN (Tg= 176 °C) consists of a nonlinear optical (NLO) active epoxy-based polymer network and an NLO active phenoxy-silicon polymer network. The decay behavior of the second-order nonlinearity of this IPN was investigated in the range from 110 to 170 °C. The stability of this IPN is superior to those of classic guest/host polymers as indicated by longer relaxation times at temperatures from 110 to 130 °C. The relaxation process of the IPN system follows Arrhenius-type behavior at temperatures ranging from 140 to 170 °C. The IPN system provides a new approach to processing for stabilization of the second-order nonlinear optical properties.
Second-order nonlinear optical (NLO) materials are of much interest for applications such as electro-optic modulation and frequency conversion.\textsuperscript{1-3} Polymeric second-order NLO materials have been studied extensively in recent years\textsuperscript{4-9} due to their low dielectric constant, large nonlinearity, and ease of processability compared to inorganic NLO materials. In these polymeric systems, the noncentrosymmetric alignment of NLO chromophores achieved by the poling technique\textsuperscript{3} is responsible for the NLO properties. However, the decay of the poled order as the material is subjected to high temperatures is still an outstanding problem. In order to use these materials in practical applications, the long term NLO stability at temperatures up to 100 °C with spikes to higher temperatures is required during device fabrication and operation. Therefore, an understanding of this decay process is the key to the development of stable polymeric NLO materials.

The relaxation of aligned NLO chromophores is often characterized by monitoring the decay of second harmonic (SH) generation as a function of time at different temperatures.\textsuperscript{10,11} In a typical guest/host system where chromophores are dispersed in an amorphous polymer, Stähelin et al. were able to fit the temporal relaxation to a Kohlrausch-Williams-Watts (KWW) equation establishing that the decay of the dipole alignment is
explained by a single relaxation phenomenon. A fit of the relaxation
times to the Williams-Landel-Ferry (WLF) equation pointed out that
relaxation of the second-order NLO properties is mainly related to the
glass transition temperature (Tg) of the media. Thus high Tg polymers are
sought for stability at elevated temperatures. In other reports, increased crosslinking has been used as a vehicle to enhance the NLO
stability. Crosslinking networks reduce the mobility of the polymer chains
which further prevent the aligned NLO chromophores from relaxing to a
random orientation. Improved stability of the nonlinear response was
established in a system with higher crosslinking density. However,
polymers which possess high Tg as well as high crosslinking density still
exhibit slow decay of the NLO properties at elevated temperatures.
Although qualitatively the relaxation behavior of crosslinked systems at
elevated temperatures seems similar to guest/host systems, a detail
experimental comparison of the relaxation behavior of the two systems has
not been reported.

The important issues we would like to address in this report are the
role of crosslinking in the stabilization of the NLO properties and the
differences in the relaxation behaviors between the crosslinked network
and a classic guest/host system. An interpenetrating polymer network
(IPN) containing aligned NLO chromophores is selected to test these issues.
An IPN is a structure in which two or more networks are physically
combined. The IPN is known to be able to remarkably suppress the
t creep and flow phenomena in polymers. The motion of each type of
polymer in the IPN is reduced by the entanglements between the networks.

The IPN system consists of an NLO active epoxy-based polymer
network and an NLO active phenoxy-silicon polymer network. The
epoxy-based NLO network is prepared from the epoxy prepolymer (BPAZO) based on the diglycidyl ether of bisphenol A and 4-(4'-nitrophenylazo)aniline functionalized with crosslinkable acryloyl groups. The second NLO network of a phenoxy-silicon polymer is based on an alkoxyilane dye (ASD) of (3-glycidoxypropyl)trimethoxysilane and 4(4'-nitrophenylazo)aniline, and the multifunctional phenoxy molecule 1,1,1-tris (4-hydroxyphenyl)ethane (THPE). The chemical structures of BPAZO, ASD, and THPE are shown in Figure 1. The IPN is formed simultaneously upon heating where the epoxy-based polymer network is formed through double reaction and the phenoxy-silicon polymer network is formed through sol-gel reaction.

(Insert Figure 1)

The Tg of the IPN was determined to be 176 °C by differential scanning calorimetry. The corona poling technique was employed to align the NLO chromophores. Poling and curing were performed simultaneously at 200 °C for 1 h. The second-order NLO properties of the poled IPN samples were measured by second harmonic generation from 1.064 μm laser radiation. The relaxation behavior of the second-order NLO properties was studied by monitoring the decay of the SH intensity as a function of time at different temperatures (from 110 °C to 170 °C) 24 h after poling and curing to avoid any space charge effect. The NLO coefficient, d33, of the poled IPN sample was measured to be 33 pm/V and 5.5 pm/V at an incident fundamental wavelength of 1.064 and 1.542 μm respectively.
Figure 2 shows the decay of the $d_{33}$ coefficient as a function of time at different temperatures for the IPN samples. The time $t=0$ was defined as the time when the samples were brought to the desired temperatures for the relaxation study. The IPN system shows no measurable decay of SH intensity after being heated at 110 °C for 168 h. An attempt was made to fit the experimental data at higher temperatures to the KWW equation which is often used to characterize the orientational relaxation in polymers.\(^\text{18}\) The KWW equation can be written as\(^\text{11}\)

$$d_{33}(t)/d_{33}(0) = e^{-(t/\tau)\beta} \quad 0<\beta<1$$

where $\beta$ determines the width of the distribution of relaxation times about some central value. The characteristic relaxation time $\tau$ is the time required for the system to decay to $1/e$ of its initial value. The equation fits the decay curves very well only for temperatures in the vicinity of $T_g$ (range from 140 to 170 °C) shown as solid lines in Figure 2 with $\beta$ values falling in a range of 0.3 to 0.4 and the $\tau$ values in a range of $10^4$ to $10^6$ sec. The data from experiments at 120 and 130 °C do not fit the KWW equation and attempted fits result in large deviations from the data. The extremely slow decay of $d_{33}$ at temperatures lower than 130 °C corresponds to very long relaxation times.

(Insert Figure 2)

Figure 3 shows a plot of log $\tau$ as a function of $1/(T_g-T)$ using the equation shown below:

$$\tau = \tau_0 e^{[B/(T_g-T)]}$$
The dashed line represents the data for the guest/host type samples reported by Stähelin et al.\textsuperscript{11} An arrow indicates the temperature at 110 °C, at which no decay of the SH coefficient was observed after a period of 168 h. The error bar for the data point at 130 °C indicates the large deviations in $\tau$ values at lower temperatures. The results show that the relaxation behavior of the IPN is significantly different from those of the guest/host type polymers. The stability of the d33 coefficient for the IPN samples at low temperatures as well as temperatures close to Tg is substantially superior to the guest/host type samples. An attempt to improve the correlation between $\tau$ and Tg, especially when the temperature is close to Tg, was made by plotting $\log \tau$ vs. $1/(Tg + 50 - T)$ as suggested by Stähelin et al.\textsuperscript{11} and Walsh et al.\textsuperscript{19} The results were not significantly improved and the nature of departure was similar to that shown in Figure 3.

\textbf{(Insert Figure 3)}

In contrast to the guest/host systems, the relaxation times $\tau$ obtained from the KWW equation at temperatures near Tg for our IPN samples follow the Arrhenius type behavior (Figure 4). The Arrhenius equation can be written as:\textsuperscript{20}

$$\tau = Ae^{(Ea/RT)}$$

where $Ea$ is the activation energy for the relaxation process. The activation energy calculated from the plot for the temperature range of 140 to 170 °C in Figure 4 is 46 kcal/mole. This is a relatively small value compared to the activation energy that is normally observed for the $\alpha$ relaxation of rigid polymers.\textsuperscript{21,22} It implies that the dipolar relaxation in this temperature range follows Arrhenius behavior similar to side-group
relaxations in amorphous polymers and is not associated with the main chain motion. As the temperature decreases to 110 °C, there is a significant increase in the relaxation time at some critical temperature where the relaxation phenomenon may no longer follow the Arrhenius behavior. It can be inferred from this result that there are at least two different relaxation mechanisms dominating the SH decay process for the IPN samples.

(Insert Figure 4)

We have demonstrated that the dipolar relaxation process of the IPN system is essentially different from that of a typical guest/host system. At temperatures lower than 130 °C the relaxation times significantly increase and deviate from the master line for guest/host systems. In addition, scaling the Tg to Tg + 50 °C does not reduce the data points to the master curve (Figure 3). The decay of nonlinearity of the IPN system at temperatures close to Tg is inherently a slower process than the guest/host system. This property allows for high temperature spikes, which maybe encountered during device fabrication. The extensive crosslinking and chain entanglements in the IPN are responsible for the enhancement of the stability. The combination of high Tg polymers and the IPN architecture is believed to further improve the temporal stability. This area of research is currently being investigated in our laboratory.

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REFERENCES


List of Figures

FIG. 1 The chemical structures of BPAZO, ASD, and THPE.

FIG. 2 Temporal behavior of the second harmonic coefficient of the IPN at different temperatures.

FIG. 3 Relaxation time $\tau$ as a function of $1/(T_g - T)$ for the IPN samples at 130, 140, 150, 160, and 170 °C. The dash line is the master curve from ref. 11 that is expected to fit guest/host system.

FIG. 4 An Arrhenius plot for the relaxation of the IPN at 130, 140, 150, 160, and 170 °C. An arrow showing 110 °C indicates the temperature at which no measurable decay of the $d_{33}$ could be observed.
(a) BPAZO

(b) ASD

(c) THPE

Fig 1