Early Cure Behavior of a Liquid Dicyanate Ester Resin

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ABSTRACT: The early cure behavior of 4,4-dicyanato 1,1-diphenolethane resin with and without incorporating Cr(acac)_3, Co(acac)_3, and Cu(acac)_2, respectively, as catalysts was investigated by gel permeation chromatography. The curing intermediates were separated by the column elution method and characterized by Fourier transform infrared, ^1H, and ^13C NMR spectroscopies. The results indicated that the formed dimer in the early cure stage is a straight chain containing a primary amino group. The formed triazine ring in the trimer has a strong catalytic effect on the remaining cyanate groups so that the reactivity of the trimers was significantly increased. The reactivities of the curing intermediates decreased with molecular size until 7-mer was reached. The initial monomer consumption is described by second-order-rate kinetics. In the presence of metal acetylacetonates, the curing reactions may be accelerated, but they did not change the reaction path and preceding sequence of reactivities. © 2001 John Wiley & Sons, Inc. J Polym Sci Part A: Polym Chem 39: 3085–3092, 2001

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INTRODUCTION

Cyanate ester resins are thermosetting materials, featured with a low dielectric constant, high glass-transition temperature, and good mechanical and processing properties. It has generally been agreed that cyclotrimerization reactions of cyanate groups that form a highly crosslinked polytriazine network are dominant curing reactions. The initiation of the reactions requires the presence of a small amount of aryl phenol impurities or adventitious water. The reactions can also be catalyzed by transition-metal complexes. However, there are discrepancies in the literature about the formation of dimeric species prior to triazine ring formation.

Fyfe et al. suggested that, based on their study on the mechanism of the curing reactions of bisphenol A dicyanate resin, no NMR evidence for the formation of dimeric or other intermediate species was observed prior to the formation of the triazine ring. However, Grenier-Loustalot and colleagues detected a dimeric species during the cure of cumyl phenyl cyanate, a monofunctional resin, via high-performance liquid chromatography and liquid-state ^13C NMR. They suggested the formation of cyclic structures in the dimers. Gupta and Macosko have also detected dimers during the cure of bisphenol A dicyanate resin with gel permeation chromatography (GPC). However, they disagreed with Grenier-Loustalot and colleagues over the formation of cyclic structures in the dimers because the two-membered rings with bond angles of 90° are highly strained and improbable to form. Gupta and Macosko suggest that the structure of the dimer is most likely a straight chain, although no further evidence was provided.

In this study, we used GPC to investigate the early cure behavior of 4,4-dicyanato 1,1-diphenolethane (DCDPE), a liquid dicyanate resin, with
and without metal acetylacetonates as catalysts. The dimers were separated by the column elution method and characterized by Fourier transform infrared (FTIR), $^1$H, and $^3$C NMR spectroscopies. The results indicate that the formed dimer is a stable straight chain containing a primary amino group, whereas the trimer is unstable and more reactive. The reactivities of the curing intermediates then decreased with molecular size until 7-mer was reached. In the presence of a catalyst, the curing reactions were accelerated, but they did not change the reaction path and preceding sequence of reactivities. Second-order-rate kinetics was also provided to describe the mechanism of initial monomer consumption.

**EXPERIMENTAL**

**Materials**

DCDPE (Ciba Geigy L10) was used as received and mixed with 0.05 phr of chromium acetylacetonate [Cr(acac)$_3$, Hayashi Japan], cobalt acetylacetonate [Co(acac)$_3$, Nakarai Japan], and copper acetylacetonate [Cu(acac)$_2$, Aldrich], respectively, at room temperature by stirring until the latter was completely dissolved. Then, about 200 mg of DCDPE samples with and without metal acetylacetonates were loaded into glass vials individually and sealed. As such, several vials were prepared for each resin formulation. The weight of the resins was accurately measured to 0.01 mg. The glass vials with the samples were then immediately heated to 110 °C in an oven and maintained at that temperature. At a designated time interval (see Fig. 1), one vial for each resin formulation was withdrawn from the oven and immediately loaded with a premeasured volume of ice-cooled tetrahydrofuran (THF) solvent to prepare 0.02 g/mL solution for the test. For separation of the reaction intermediates, the prepared solutions were vacuum-dried at room temperature, and then the dried products were redisolved in dichloromethane before passing through

![Figure 1. GPC spectra of (a) DCDPE resin and those incorporating 0.05 phr of (b) Cr(acac)$_3$, (c) Co(acac)$_3$, and (d) Cu(acac)$_2$, respectively, after curing at 110 °C for various indicated times.](image-url)
the separation column packed with silica gel (Merck Silica Gel 60). The effluents were repeatedly collected from the column for 3 h. After draining all the solution in the separation column, the residue remaining in the column was flushed off with THF solvent and collected.

Characterization

GPC measurements with THF as the eluent (1 mL/min) were carried out at 40 °C using a Testhigh series III pump and a Testhigh refractive-index detector. One PhenolGEL 550A and two PhenolGEL MXL columns in series were used for separation of the reaction intermediates. Their molecular weight was estimated by referring to the polystyrene standards with a narrow molecular weight distribution. FTIR spectra were recorded on a Jasco 300E FTIR spectrometer with the specimens coated on a KBr plate. $^1$H and $^{13}$C NMR spectroscopies of the reaction intermediates were carried out with a Bruker Model AM-500WB NMR spectrometer. The samples were vacuum-dried at room temperature first and then redissolved in deuterated chloroform.

RESULTS AND DISCUSSION

Reaction Intermediates

Figure 1 shows the GPC spectra of DCDPE resins with and without 0.05 phr of Cr(acac)$_3$, Co(acac)$_3$, and Cu(acac)$_2$, respectively, after reacting at 110 °C at various times. The neat resin contained a small amount of dimers and multimers. When the reactions proceeded at 110 °C for 25 min, the amount of dimers and multimers increased for all four samples. The neat resin and that incorporating Cr(acac)$_3$ have a similar reactivity. They both gelled after curing for 750 min. The resin incorporating Co(acac)$_3$ gelled after 425 min, and that incorporating Cu(acac)$_2$ gelled after 115 min. Although the catalytic effect of metal acetylacetonates increases in the sequence of Cu(acac)$_2$ > Co(acac)$_3$ > Cr(acac)$_3$, the GPC spectra of all the resins under the same extent of cure are basically similar.

When the DCDPE resins incorporating 0.05 phr of Cr(acac)$_3$ were reacted for 25 min, their dimers were separated from the cured resins by the column elution method in an elution time period of 110–160 min, as shown in Figure 2. In fact, the dimers were eluted along with a small amount of monomers and barely detectable multimers. Their FTIR spectrum as compared with that of the DCDPE monomer (see Fig. 3) revealed that the effluent contained free primary amine groups (absorption peaks at 3500 and 3350 cm$^{-1}$ contributed by stretching of $\text{O-NH}_2$ and at 1373 cm$^{-1}$ by $\text{C-O-N}$ stretching$^{12}$) and carbonyl groups (at 1720 cm$^{-1}$). The $^1$H NMR spectrum shown in Figure 4 indicates that the effluent had two kinds of free primary amine groups. The major resonance peak at 5.18 ppm was contributed by the dimers whose chemical structure is outlined in Structure 1. On the other hand, the minor peak at 6.72 ppm was mainly contributed by the carbamate groups formed by the reaction of monomers with adventitious water, whose structure is demonstrated in Structure 2.
The splitting of the minor peak resulted from the presence of aromatic protons close to the phenol groups of unconverted monomers. We have checked this small contribution from the $^1$H-$^{13}$C chemical shift correlated two-dimensional NMR spectrum (data not shown). Grenier-Loustalot et al.\(^9\) also reported that the aromatic protons close to the phenol groups of unconverted bisphenol A diacyanate monomers have a chemical shift of 6.6 ppm. The chemical structure of the previously shown dimers was further supported by $^{13}$CNMR spectroscopy (see Fig. 5). It is probable that the dimers were formed by the reaction of a carbamate group of the preceding compound with an isocyanate group of the monomer. They are rather stable after formation and can stay with the DCDPE monomers at room temperature without proceeding with further detectable reactions for at least 1 week.

**Reactivity of Reaction Intermediates before Gelation**

Only part of the reaction intermediates of the DCDPE resins was flown out from the separation column after applying the column elution method with dichloromethane as the mobile phase. After 200 min of elution, almost no reaction intermediate was detected in the effluent from the separation column. After that, the residue remaining in the column was flushed off with THF solvent, and its GPC spectrum exhibited a clear trimer peak at an elution time of 22.2 min, as shown in Figure 6, that was not observed in the previous GPC spectra of the resin and its effluents. Apparently, trimer is less soluble in dichloromethane than in THF. By comparing the sequence of peaks appearing at different elution times among the GPC spectra of resin, its effluent, and the residue remaining in the separation column, we were able to identify peaks from monomer to 7-mer (see Fig. 6). Assuming that the area of the monomer peak of resin before curing is 1, the number percentage of monomers to 7-mers at various curing times can estimated from Figure 1, and the results of the DCDPE resins containing 0.05 phr of Cr(acac)$_3$, Co(acac)$_3$, and Cu(acac)$_2$, respectively, are illustrated in Figure 7. As seen in the figure, the amount of monomers was gradually decreased with time for all three samples. Because of the similarity between the data of the neat resin and that containing Cr(acac)$_3$, the former is not shown. The dimers, having produced about 7% during resin storage, slightly increased and then barely decreased its number percentage before gelling for most of the samples except that con-
taining Cu(acac)$_2$. The detectable decrease of dimers for the latter was due to the higher extent of reaction caused by the greater catalytic effect of Cu(acac)$_2$. The change in number percentage of the trimer, tetramer, and 5-mer almost followed the trend of the dimers. However, 6-mer and 7-mer slightly increased their amount. Nevertheless, the amount of the formed trimers for all four resin systems was the least compared with other multimers. In other words, their reactivity was the highest. As soon as the trimers were produced, they reacted to form tetramers. If the reactivity of the tetramer is higher than the trimer, then its amount would be lower than that of the latter. Additionally, the reactivity of 5-mers was lower than that of the tetramer. The reactivity decreased with molecular size until 7-mer was reached. Thus, the amount of multimers detected by GPC reflects their reactivities. Catalytic effects of Cu(acac)$_2$ and Co(acac)$_3$ on DCDPE resin only accelerated their reactions and did not change the preceding sequence of reactivities.

The cyclotrimerization reactions of cyanate groups that form triazine groups were dominant reactions in the initial cure of DCDPE resins as indicated by the $^{13}$C NMR spectra in Figure 8. The appearance of the resonance peak at 173 ppm contributed by the carbons in triazine rings justified that the trimers and other multimers were a polytriazine. The highest reactivity of trimers as compared with other multimers indicates that the formed triazine ring has a strong catalytic effect on the remaining isocyanate groups. This catalytic effect has long been considered as the catalytic effect of triazine products on the overall curing reaction of cyanate resins.$^4$ We also believe this effect rendered the gel conversion at the gel point deviated from the theoretical prediction of 0.5$^{11,13}$ because the formed triazine rings, which accelerate the reactions of neighbor cyanate groups, might facilitate the formation of intramolecular cycles.

According to the aforementioned experimental results, the initial cure path of DCDPE resins can be described by the following eqs.

$$\text{Ar-OCN} + \text{H}_2\text{O} \rightarrow \text{Ar-C} \rightarrow \text{NH}_2$$

$$\text{Ar-C} \rightarrow \text{N} \rightarrow \text{ArCN} + \text{Ar-OCN} \rightarrow \text{Ar-C} \rightarrow \text{N} \rightarrow \text{OA} \rightarrow \text{H}_2\text{O}$$

$$\text{Ar-C} \rightarrow \text{N} \rightarrow \text{OA} \rightarrow \text{H}_2\text{O}$$

$$\text{Tri-OCN} + \text{H}_2\text{O} \rightarrow \text{Tri-C} \rightarrow \text{NH}_2$$

$$\text{Tri-C} \rightarrow \text{NH}_2 + \text{Ar-OCN} \rightarrow \text{Tri-C} \rightarrow \text{N} \rightarrow \text{OA} \rightarrow \text{H}_2\text{O}$$

$$\text{Tri-C} \rightarrow \text{N} \rightarrow \text{OA} \rightarrow \text{H}_2\text{O}$$

$$\text{Tri-C} \rightarrow \text{N} \rightarrow \text{OA} \rightarrow \text{H}_2\text{O}$$

$$\text{Tri-C} \rightarrow \text{N} \rightarrow \text{OA} \rightarrow \text{H}_2\text{O}$$

**Figure 6.** GPC spectra of (–) DCDPE resin incorporating 0.05 phr Cr(acac)$_3$ after curing at 110 °C for 25 min, (—) its effluent from separation column in an elution time period of 50–53 min, and (⋯⋯) the residue that was unable to be removed from the separation column with dichloromethane, flushed off with THF.
By following the suggested method, the equilibrium constant \( K_1 \) for reaction 1 is defined as

\[
K_1 = \frac{k_1}{k_{-1}} = \frac{I_1}{CW}
\]  

(1)

where \( I_1 \), \( C \), and \( W \) are the concentrations of imidocarbonate, monomers, and water, respectively. Because the concentrations of dimers, trimers, tetramers, and so forth were almost constant in the initial curing stage (see Fig. 7), the following quasi-steady-state conditions were assumed:

\[
\frac{dI_2}{dt} = k_3 I_1 C - k_5 I_2 C = 0
\]  

(2)

\[
\frac{dT}{dt} = k_3 I_2 C - k_4 TW = 0
\]  

(3)

\[
\frac{dI_3}{dt} = k_4 TW - k_5 I_3 C = 0
\]  

(4)

Figure 7. Plot of number percentage of (■) monomers, (●) dimers, (▲) trimers, (▼) tetramers, (♦) 5-mers, (+) 6-mers, and (×) 7-mers versus curing time at 110 °C for DCDPE resins containing 0.05 phr of (a) Cr(acac)$_3$, (b) Co(acac)$_3$, and (c) Cu(acac)$_2$.
\[
\frac{dI_4}{dt} = k_5 I_4 C - k_4 I_4 C = 0
\]  
(5)

\[
\frac{dI_n}{dt} = k_{3n-1}I_{n-1} C - k_{3n+1}I_n C = 0
\]  
(6)

where \(I_n\) is the concentration of n-mers, and \(k_{3n-1/2}\) and \(k_{3n+1/2}\) are the reaction constants at reaction steps of whole numbers of \(3n - 1/2\) and \(3n + 1/2\), respectively. Thus, the rate of monomer consumption, \(\frac{dC}{dt}\), can be estimated by

\[
\frac{dC}{dt} = k_2 I_2 C + k_3 I_3 C + k_4 I_4 C + \cdots + k_{3n+1/2}I_n C
\]  
(7)

\[
= nK_1k_2WC^2
\]  
(8)

From the mass balance, the initial concentration of water, \(W_o\), is expressed as follows

\[
W_o = W + I_1 + I_2 + \cdots + I_n
\]  
(9)

\[
= W \left(1 + K_1 C + \frac{k_2}{k_3} K_1 C + \cdots + \frac{k_2}{k_{3n+1/2}} K_1 C\right)
\]  
(10)

\[
= W(1 + p)
\]  
(11)

where

\[
p = K_1 C + \frac{k_2}{k_3} K_1 C + \cdots + \frac{k_2}{k_{3n+1/2}} K_1 C
\]  
(12)

Assuming that \(p \ll 1\), \(W \approx W_o\). Thus

\[
-\frac{dC}{dt} = kC^2
\]  
(13)

where \(k = nK_1k_2W_o\). By integrating eq 3, we obtain

\[
\frac{C_o}{C} - 1 = kC_0t
\]  
(14)

where \(C_o\) is the initial concentration of monomer. Incorporating the experimental data of Figures 1 and 4, we plotted \(\frac{C_o}{C} - 1\) versus \(t\) for all resin systems, as shown in Figure 9, and found that the data are well fit with eq 14 in the initial cure stage but deviated after curing for 250 min. Using the first-order least-squares method to fit the data before 250 min, we can obtain the slope, \(kC_o\). \(C_o\) was 4.29 M (mol/L), estimated from the density of DCDPE resin at 110 °C. Thus, the rate constants, \(k\), for monomer consumption of neat resin and those containing Cr(acac)_3, Co(acac)_3, and Cu(acac)_2 were estimated as \(2.10 \times 10^{-4}\), \(2.14 \times 10^{-4}\), \(6.62 \times 10^{-4}\), and \(2.29 \times 10^{-3}\) M\(^{-1}\) min\(^{-1}\), respectively. Most of the experimental data for monomer consumption after 200 min of curing were more than those predicted by second-order-rate kinetics, possibly because the latter did not consider the reactions between two individual growing chains, which become significant when the cure reaches the gel point.

**CONCLUSIONS**

This study has clearly identified the curing intermediates of DCDPE cyanate resins from dimer to 7-mer. The dimer is a stable species, but the trimer is the most reactive. The reactivity decreased with molecular size until 7-mer was reached. Although metal acetylacetonates can catalyze the curing reaction, they do not change the preceding sequence of reactivities. The monomer consumption in the initial cure stage can be described by second-order-rate kinetics.
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REFERENCES AND NOTES