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Thermal oxidation of three epoxy resins prepared from flexible or rigid prepolymer and hardeners was studied by monitoring epoxy mechanical and physical changes. The physical changes were followed by mass measurements, glass transition temperature using DSC and sub-glass \( T_g \) transition using DMA. It was put in evidence that embrittlement is not directly associated to \( T_g \) or mass loss changes since epoxy network based on isophorone diamine (IPDA) hardeners were shown to undergo mainly a chain scission at the beginning of exposure process whereas epoxy network based on trioxatridecane diamine (TTDA) hardeners exhibits a crosslinking process with a significant mass loss. The only common feature for both epoxy systems to understand embrittlement is the drop of amplitude of \( T_g \) transition with oxidation.

1. Introduction

Epoxy family cover a large diversity of polymer networks differing by the nature of resins and hardeners commercially available. Depending on resin and hardener used, range of \( T_g \) can vary from 60 °C up to 250 °C for instance tris(4-hydroxyphenyl) methane triglycidyl ether (Tactix 742) and diglycidyl ether of bisphenol A (Tactix 123), with an aromatic diamine hardener: 4,4'-diaminodiphenyl sulfone [1,2].

In many applications such as composites for aeronautical components, a peculiar concern is associated to long term ageing effect since oxidation affects strongly the epoxy mechanical properties, especially the failure properties. In this context, epoxy lifetime is often empirically related to mass loss induced by oxidation. However, to predict lifetime with an accurate manner for all epoxy networks, embrittlement mechanisms associated to oxidation have to be deeply studied. Since embrittlement process corresponds to failure properties drop, let us recall some approaches relating these properties with epoxy networks characteristics.

Failure properties, especially strain at break or toughness \( G_{IC} \), are governed by the capacity to promote plastic deformation. Even if the relationships between structure and plastic deformation for epoxy networks in glassy state have been studied since the seventies [3–8], the molecular origin for plastic deformation mechanisms in the epoxy networks still is an open question. The variety of epoxy networks studied in the literature and the difficulties to investigate plastic deformation zone can explain this lack of consensus in literatures. Several approaches have been proposed to understand the molecular aspects of plastic deformation in epoxy networks:

- Plastic deformation is promoted if glass transition is close to mechanical test temperature. In other words, plasticity is associated to cooperative mobility.
- Plastic deformation is linked to molar mass between crosslinks (\( M_c \)). In this approach, plastic deformation obeys the relationships proposed in rubber elasticity theory where deformation at failure is associated to the chain extensibility limit. Indeed, several authors have put in evidence that epoxy network fracture energy is proportional to \( M_c^{1/2} \) [9]. However, an opposite trend has been witnessed by Urbaczewski-Espuche et al. [10].
- Plastic deformation is associated to secondary transition below \( T_g \), which is often called \( T_\beta \) transition. The \( T_\beta \) transition in networks based on DGEBA is attributed to local motions linked to hydroxyether segments [11]. As a result, plastic deformation (cooperative motions) is activated by this local mobility for a given state stress.

In this paper, we propose to follow epoxy networks modifications during oxidation in terms of glass transition and \( T_\beta \) transition and to relate these modifications to embrittlement process. For this purpose, we have chosen three epoxy networks: two systems having a relatively low \( T_g \) (DGEBA/TTDA and DGBU/IPDA, 69 and 60 °C respectively) and a system having a high \( T_g \) (DGEBA/IPDA, 166 °C). Furthermore, it has been witnessed that among these
systems, one system undergoes mainly chain scission (DGEBA/IPDA) whereas the other undergoes mainly crosslinking [12]. The final objective is then to propose a common scenario to explain embrittlement induced by oxidation for epoxy networks.

2. Experimental

2.1. Materials

Two kinds of prepolymerms are used.

- The diglycidyl ether of bisphenol A (DER 332, CAS 1675-54-3—ref 31185 supplied by Sigma Aldrich) has a degree of polymerization n ~ 0 and a number average molecular mass equal to 340 g/mol.

- The diglycidyl ether of 1,4-butanediol named DGBEU (CAS 2425-79-8—ref 220892 by Sigma Aldrich) has an average molecular mass of 202.25 g/mol.

Two kinds of hardeners have been chosen:

- The isophorone diamine here named IPDA (CAS 2855-13-2—ref 118184 by Sigma Aldrich, M = 170.3 g/mol).

- The 4,7,10-trioxa-1,13-tridecanediamine here named TTDA (CAS 4246-51-9—ref 369519 by Sigma Aldrich, M = 220.3 g/mol).

The chemical structure of chemicals is given in Fig. 1. These components were mixed in stoichiometric ratio and fully cured checked by the total disappearance of exothermal signal and by FTIR from the loss of 914 cm⁻¹ peak attributed to epoxide group. Details of films preparation, thermal characterization of the cure and glass transition of final networks obtained can be found elsewhere [13]. Thin films have been processed in heating press. Thermal ageing under atmospheric air was performed in ventilated ovens (calibrated at ± 3 °C) at 110 °C. Films having a thickness lower than 100 µm to avoid diffusion limited oxidation were periodically removed for analysis.

2.2. Characterization

2.2.1. Oxidative products concentrations

To assess oxidative products concentration during exposure, FTIR spectroscopy in transmission mode was performed on free standing films using a Frontier spectrophotometer (PerkinElmer) in the 550 to 4000 cm⁻¹ wavenumber range by averaging 16 scans with a 4 cm⁻¹ resolution. Spectra were interpreted using the Spectrum software (PerkinElmer) in order to determine the absorbance value from which the carbonyl and amide concentrations, the main oxidation products, were calculated according to the method defined elsewhere [13].

2.2.2. Mechanical properties

Changes of mechanical properties during the oxidation of epoxy/amine have been followed by tensile testing on an Instron® 5881 machine. Dumbells samples have been obtained by cutting 100 µm films. The effective zone has a dimension of 30 mm length and 4 mm width for DGEBA/TTDA and 10 mm length and 2 mm width for DGEBA/IPDA and DGBEU/IPDA. The crosshead speed was at 1 mm/min was constant during the test and a 100 N cell was used. Tests have been performed in a controlled environment at 23 °C on five samples.

2.2.3. Mass loss

Weight changes have been monitored during the thermal oxidation of epoxy resin thanks to the Dual Range XS105 scale from Mettler Toledo with an accuracy of ±0.01 mg. Those measurements have been duplicated on two samples with an initial mass of approximately 100 mg.

2.2.4. Glass transition measurement

Differential scanning calorimetry measurements were made with a DSC Q1000 (TA Instruments). Samples with mass ranging between 3 and 5 mg are sealed in aluminum pans were heated from 0 °C to 250 °C at a 10 °C/min ramp under nitrogen flow (50 ml/min). Results were interpreted using TA Analysis software. DSC analyses were done to check the total cure of samples and to measure the value of the glass transition temperature of aged samples. Tg values were measured during the second heating ramp (i.e. after having removed the thermal history of samples). Two samples of the same resin have been analyzed for each exposure time to control measurements reproducibility.

2.2.5. Sub glass mobility

To put in evidence sub glass transition, i.e. β transition, dynamical mechanical analysis (DMA) measurements were performed between −100 and 40 °C at 1 Hz on films samples. For this purpose a DMA Q800 from TA instruments was used in tension mode.

3. Results

3.1. Mechanical properties changes

To put evidence drop of plastic deformation by oxidation, samples were characterized by classical tensile tests up to failure at room temperature. Fig. 2 shows stress-strain curves for all epoxy networks during exposure at 110 °C. The stress-strain curves show a classical shape for epoxies [14–16] from which the strain at break was determined to monitor plastic deformation.

First of all, for DGEBA/IPDA and DGEBA/TTDA networks (Fig. 2a and b), a decrease of strain at break is associated to a modulus increase up to complete embrittlement whereas stress-strain changes for DGBEU/IPDA are clearly more complex.

First, a modulus increase is observed during thermal oxidation in the case of DGEBA/IPDA and DGEBA/TTDA. This phenomenon, so called anti-plasticization phenomenon, has already been observed during the epoxy cured with aromatic amine during oxidation [1] where modulus increase is related to decrease in local mobility associated to β transition.

Strain at break changes clearly shows an embrittlement process in the case of DGEBA/IPDA and DGEBA/TTDA whereas strain at break...
changes for DGEBU/IPDA are surprisingly more complicated. To put in evidence the embrittlement process for each system, strain at break values as a function of exposure time are reported in Fig. 3:

- for DGEBA/IPDA, it decreases continuously. The time at which strain at break is half is initial value is ca 200 h.
- for DGEBA/TTDA, the initially low strain at break value decreases at a low rate. The time at which $\varepsilon_R = \varepsilon_{R0}/2$ is ca 2200 h.
- last, DGEBU/IPDA displays the most surprising trend: $\varepsilon_R$ increases first very strongly and reaches values ca 100%. After a sort of plateau, it decreases to a value close to $\varepsilon_{R0}/2$ after ca 1630 h.

3.2. Mass changes vs embrittlement

In the introduction part, we have mentioned that mass loss is often used as a tracer for embrittlement. In other words, it is proposed that samples become brittle when mass loss exceeds 10%. To check the validity of this assumption, we have followed mass changes during exposure in the same conditions. Mass changes as a function of exposure time are displayed in Fig. 4. In this latter, characteristic time for embrittlement (time for $\varepsilon_R \sim \varepsilon_{R0}/2$) is also reported for each networks by an arrow.

According to Fig. 4, DGEBU/IPDA displays an important mass loss (more than 20% after 1000 h of exposure at 110 °C). This network is by far the most unstable system from the mass loss point of view. Indeed, embrittlement occurs at a mass loss close to 2% and 12% for DGEBA/IPDA and DGEBA/TTDA respectively.

As a result, it is obvious there is no direct correlation between an only one value of mass loss and the embrittlement time for the three systems under investigation. It means that the three systems studied here differ by the consequences on oxidation on the architecture of the network that will be studied below.
3.3. Glass transition changes vs embrittlement

During oxidation, it is expected that epoxy network undergoes macromolecular changes during oxidation since a chain scission process occurs. Since it is challenging to detect the chain scissions in networks, measurement of glass transition temperature is particularly interesting since $T_g$ is related to the concentration in elastically active chains [14,17], which decreases with chain scissions and increases with crosslinkings. $T_g$ changes during exposure and characteristic time for embrittlement (arrow) are reported in Fig. 5, which shows:

- DGEBA/IPDA displays a significant $T_g$ drop revealing a predominant chain scission process.
- DGEBA/TTDA displays a significant $T_g$ increase revealing a predominant crosslinking process.
- DGEBU/IPDA show first a slight $T_g$ drop followed by an increase. These changes can be attributed to chain scission events at low degree of conversion followed by a majority crosslinking process.

Interestingly, it appears the common point of both networks which undergo crosslinking (DGEBA/TTDA and DGEBU/IPDA) is the presence of methylene sequences. A quantitative study and mechanism proposals of all these aspects can be found elsewhere [13].

3.4. Sub-glassy transition

Since two epoxy networks (DGEBA/IPDA and DGEBA/TTDA) remain in glassy state at room temperature even after oxidation process, it seems to us that sub-glass transition (so called $\beta$ transition) has to be investigated for both systems. Tan $\delta$ spectra obtained by DMA are shown in Fig. 6 for several exposure times.

First, a clear $\beta$ transition is put in evidence between $-60$ and $-20 \, ^\circ C$ for both epoxy networks. Secondly, it appears that tan $\delta$ amplitude decreases clearly for exposure times increasing. These spectra modifications could be associated to oxidation DGEBA segments. Indeed hydroxypropylether groups in DGEBA can be easily oxidized since they contain some unstable sites such as CH groups or CH$_2$ groups in alpha position of heteroatoms [18].

4. Discussion

The main objective of this study is to propose mechanisms responsible for embrittlement process during oxidation. In terms of structure-property relationships, different approaches have been proposed in the literature to relate macromolecular architecture and failure properties: gap between $T_g$ and temperature for mechanical testing, molar mass between crosslinks mechanically active ($M_c$) or sub glass transition intensity. To investigate the most relevant parameter, in other words relevant whatever the epoxy
networks and the macromolecular modifications (chain scission or crosslinking) occurring during oxidation, we propose here to check out these three approaches successively.

4.1. Gap between Tg and temperature for mechanical testing

To check out the first approach, strain at break has been plotted as a function of difference between Tg measured and imposed temperature for tensile testing, room temperature (RT) in this study (Fig. 7). First, since initial Tg-TRT value for DGEBU/IPDA system is below 30 °C and decreases below 20 °C during oxidation up to 1000 h time of exposure, the mechanical behavior of this system tends to a rubbery behavior which explains large deformation (see for instance the stress-strain curve at 1006 h in Fig. 2c). At high oxidation level, the Tg-TRT increase leads to return to a glassy state and thus to strain at break values. It is noteworthy that the particular DGEBU/IPDA behavior during oxidation requires low initial Tg-TRT value and a chain scission followed by a crosslinking mechanism.

In the case of DGEBA/TTDA network, even if the gap at the initial state is low (45 °C), this gap is inversely proportional to strain at break. In a same time, strain at break drops with the gap value initially close to 140 °C for the DGEBA/IPDA. As a result, the gap between Tg and RT is not a common feature to explain embrittlement for both systems.

4.2. Molar mass between crosslinks

It has been proposed that properties at failure are proportional to M^1/2 with M average molar mass between mechanically active crosslinks [9]. This relationship has been also experimentally validated for rubbers and is based on the fact that strain at break is related to maximum stretching of chains between crosslinks. The failure then occurs when chain stretching reaches its limit of chain extensibility defined as the square root of the number of crosslinks [19]. One can notice this approach in the case of glassy networks implies that large deformation is related to cooperative motions.

To assess MC values in our case, a possible way is to use DiMarzio’s approach to estimate crosslink density (n) as following:

\[ T_g = \frac{T_{gl}}{1 - (K_{DM}Fn)} \]

With K_{DM} is the DiMarzio’s constant ca 2.91 for epoxy/amine trifunctional networks [20], n is the crosslink density (mol kg^{-1}), T_{gl} is the glass transition of the corresponding « virtual » linear polymer and F is the Flex parameter (kg mol^{-1}) related to the molar mass per flexible bond. Validity and physical meaning for the networks under study have been reviewed in Refs. [20,21].

Molar mass between crosslink (MC) can then be obtained from crosslink density (n) by:

\[ M_C = \frac{2}{fn} \]

Where f is the network functionality equal to 3 in our case.

In Fig. 8, we have reported strain at break as a function of M^1/2. Concerning DGEBA/TTDA networks, a good correlation is obtained. However, DGEBA/IPDA follows an opposite trend: strain at break decreases with MC increasing. As a result, even if this approach could be consistent for DGEBA/TTDA network possibly because its low Tg value promotes cooperative motions, DGEBA/IPDA network shows a counterexample leading to conclude that MC is not the relevant parameter to monitor embrittlement.

4.3. Sub glass transition intensity

According to previously published results [22] evidencing that impact strength is governed by sub glass transition so called β transition, sub glass transition depletion could be associated to strain at break (and related quantities) decrease [23–26]. To check out this scenario, Fig. 9 below shows strain at break as a function of the intensity (area) of the β transition. It is observed that the plastic deformation decreases with the area the β transition. This scenario thus appears valid to describe not only the embrittlement of DGEBA/TTDA but also, and especially, the embrittlement of DGEBA/IPDA. However, the linear characteristic for this relationship between ductility and β transition amplitude needs to be confirmed in the future.

5. Conclusions

By following physical modifications of three epoxy amine networks during oxidation, we have investigated the most relevant physical parameter responsible for embrittlement. Among these epoxy systems, DGEBA/TTDA has a low initial Tg and undergoes mainly crosslinking whereas DGEBA/IPDA shows a relatively high
Tg and undergoes mainly chain scission. The third chosen system (DGEBU/IPDA) has an initial Tg close to room temperature and shows a chain scission process followed by a crosslink process at high oxidation level. Firstly, it has been proven that mass changes cannot be considered as a good parameter to monitor properties at failure. Indeed, DGEBA/IPDA becomes brittle for a mass loss close to 2%, embrittlement of DGEBA/TTDA occurs at 12%.

To put in evidence the relevant parameter according to different approaches proposed in the literature, we have followed Tg changes focusing especially on the gap between Tg and temperature of tensile testing TRT. It appears that only strain at break changes for DGEBU/IPDA can be explained by the Tg-TRT evolution. To investigate DGEBA/TTDA and DGEBA/IPDA embrittlement, two physical parameters have been correlated to strain at break drop. The first one is the molar mass between crosslink MC calculated from Tg by DiMarzio’s theory and the second one is the intensity of peak associated to transition. As a result, transition disappearance during oxidation is the only molecular modification which can be associated to embrittlement for both epoxy systems. This finding shows clearly that embrittlement is not directly associated to chain scission or crosslinking mechanism. Knowing that the transition assigned to hydroxypropylether groups local mobility is responsible for ductility, embrittlement appears to be governed by oxidation of these groups in the DGEBU segment.

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Appendix

We are here interested in discussing on the relative weight loss observed in each kind of epoxy/amine system so as to try to justify them from a structural point of view.

For that purpose, we will first try to quantify the occurrence of a volatile compound from chain scissions. In the aim of simplicity, we will first only consider that only CH2 in x position of nitrogen are reactive (having in mind that the effect of the other CH groups is isopropanol will involve the same kind of effects).

From Table 1, it can thus be estimated that 8 scissions in DGEBA/IPDA are needed to generate 1.66 volatiles. This number may be lower if we consider that the tertiary C-H in x position of nitrogen rather yields to scissions of 3’ and 3” bonds instead of scission of 3 bond (which would give a >N′ radical, that was not observed, to the best of our knowledge, in epoxy oxidation).

In DGEBA/TTDA (Table 2), the generation of volatile compounds is favored for at least two reasons:
- the absence of tertiary CH which favors the yield in scissions of elastically active chains
- the participation of CH2 in x position of oxygen

References


