Characterization of the ageing of an epoxy resin using high temperature nanoindentation

Bruno Passilly* and Romain Delannoy

Département Matériaux et Structures, ONERA, 29 avenue de la division Leclerc, 92322 Chatillon Cedex, France

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Abstract. This article aims to understand better the mechanical properties and behavior of organic matrix composite materials under elevated temperature conditions. Two specific specimens of cured RTM6 epoxy resin are tested with DMA analysis: one being unaged and the other one aged for 5000 h at 130 °C under ambient air. Anti-plasticization effects seem to occur on aged resin. Series of nano-indentation tests are carried out from the surface to the core of the sample so as to measure gradient properties of the resin, at temperatures up to 150 °C using a high temperature indentation machine prototype. Thermo-oxidation phenomena involve oxidized layer formation during thermal ageing of the epoxy resin which is characterized through measurements of indentation Young’s modulus. After aged treatment, the variation of Young’s modulus of the oxidized layer at the surface of the sample is not clearly affected by the increasing test temperature whereas Young’s modulus of the core of the sample is decreasing significantly with the temperature test as on unaged epoxy resin. Thus, asymptotic growing of the oxidized layer is then confirmed.

Keywords: high temperature / nanoindentation / Young’s modulus / ageing / epoxy resin / durability

1 Introduction

The development of the aerospace industry during the last century led to an unrestrained pursuit of better technologies and overall aircraft performances. As such, weightlessness remains today a major challenge for aeronautic companies.

Metal and alloy structures have progressively been replaced by composite materials; especially carbon/epoxy laminates, as they possess interesting specific mechanical properties. However, a better understanding of their durability is necessary to implement these materials further, especially in severe thermal conditions.

To predict mechanical behavior of polymer matrix composite materials at high temperatures, it is necessary to understand the degradation mechanisms occurring during thermal ageing. In presence of oxygen, oxidation of the polymer can take place forming a layer of oxidized epoxy
polymer at the surface of the material [1–3]. Mechanical properties degradation during thermal ageing have yet to be thoroughly predicted, specifically at higher temperatures.

This study focuses on an experimental investigation of the evolution of surface mechanical properties of an epoxy resin matrix with thermal ageing. The material behavior will be characterized using high temperature nanoindentation to simulate flight conditions in high temperature regions of the aircraft.

2 Samples preparation

Specimens used in the present study are cut from a thermoset RTM6 epoxy resin plate. RTM6 epoxy resin presents a gel temperature of 180 °C for a gel time of 30 min. The plate is prepared according to a specific polymerization cycle, composed of a 2 h long cure at 180 °C, followed by a post-cure of 2 h at 200 °C to obtain additional crosslinking.

Glass transition of RTM6 resin is about 210 °C (data produced by Hexcel Corporation).

The epoxy resin is obtained by mixing pre-polymer chains of tetraglycid methylene dianiline – TGMDA, with two amine hardeners: 4,4'-methylenebis (2-isopropyl-6-diethylaniline) – M-MIPA, and 4,4'-methylenebis (2,6-diethylaniline) – MDEA as shown in Figure 1.

Crosslinking is mainly obtained by polyaddition between cyclic ether functions of the pre-polymer chains with amine functions of the two hardeners.

Isothermal ageing is carried out on the monolithic epoxy samples at 130 °C in an oven under atmospheric air, for a duration up to 5000 h. Samples are then taken out from the chamber after ageing time, and cut to appropriate dimensions to be positioned in the nanoindentation machine prototype. Specimens were polished down to 0.25 μm to get a polished surface to indent on. To avoid any problems of humidity, samples stayed at least 48 h at 70 °C before performing any indentation test series. Humidity of the specimen would influence the measured results of mechanical properties and could create plasticization of the polymer [4]. Characterization of the epoxy specimens are carried out at multiple temperatures, up to 150 °C.

3 Instrumented high temperature nanoindentation

3.1 High temperature nanoindentation apparatus

The high temperature nanoindentation device shown in Figure 2 used for this study is a prototype developed by ONERA [5]. It uses the same functioning principles as commercial nanoindentation devices. The force $P$ applied onto the sample by the indenter, and the penetration depth $h$, are recorded throughout the indentation tests.

Indenter displacement is controlled by a piezoelectric actuator, and measured by a capacitive displacement transducer. The sample is placed onto a load cell measuring the applied load by the indenter. An additional displacement transducer is placed underneath to measure the load cell deformation. Penetration depth is then determined by subtracting the load cell displacement to the indenter displacement.

The indenter and the sample are both heated separately by cylindrical furnaces, thus preventing any thermal gradient and drift upon contact between the indenter and the studied specimen [6]. Temperature is monitored, and studied temperatures can go up to 200 °C.

3.2 Loading and unloading cycle

The same loading protocol is used for every nanoindentation test and a typical indentation curve is shown in Figure 3. First, a loading phase is carried out, during
which the indenter is steadily applying a load onto the sample surface. The loading and unloading rate is about 2.5 mN/s. Both elastic and plastic deformations occur, as the hardness impression formed can attest to it.

Once the maximum load of 100 mN is reached, a creep phase is initiated for 200 s, during which the load is maintained at maximum value. This phase is very important for high temperature nanoindentation tests on viscoelastic materials such as RTM6 epoxy resin.

Finally, an unloading phase is carried out until there is no more contact between the indenter and the material. During unloading, the mechanical behavior is assumed to be only elastic, thus facilitating the calculation of indentation hardness and modulus.

### 3.3 Determination of the mechanical properties

A Vickers diamond indenter was used during this study with Young’s modulus and Poisson coefficient assumed to be respectively of $E_i = 1141$ GPa and $\nu_i = 0.07$. Vickers hardness is defined as the quotient of the maximal applied load $P_{\text{max}}$, expressed in Newton, by the projected indentation area $A_p$, expressed in square millimeters:

$$H_v = \frac{P_{\text{max}}}{A_p}. \tag{1}$$

Macroscopic hardness values are usually determined by measuring dimensions of residual indentations imprinted on the sample surface. In indentation, pile-up and sink-in effects are usually studied to be taken into account or not [7] (Fig. 4).

The method introduced in 1992 by Oliver and Pharr [8] is used to determine indentation hardness $H_{IT}$ and modulus $E_{IT}$ using unloading measurements during indentation tests. As such, pile-up of the material is not taken into account for the further calculations because this effect has not been observed by scanning electron microscope. It would be interesting to use in situ high temperature nanoindentation measurements by SEM in order to describe the behavior of the indented sample during the indentation test [9].

The profile of the contact geometry around the indenter is described in Figure 5. A few quantities are defined: the sink-in displacement of the surface at the contact perimeter, $h_s$, and the effective contact depth of the indenter into the material, $h_c$. Maximum displacement $h_{\text{max}}$ can be then defined by using these quantities:

$$h_{\text{max}} = h_{s_{\text{max}}} + h_{c_{\text{max}}}. \tag{2}$$

It is assumed the pyramidal tip punch behavior can be modeled by a conical indenter with a half-included angle of 70.3°. Models [10,11] show the amount of sink-in displacement around the tip is calculated using the applied loading $P_{\text{max}}$ and the indentation unloading stiffness $S_{IT}$:

$$h_{s_{\text{max}}} = \frac{\varepsilon P_{\text{max}}}{S_{IT}}, \tag{3}$$

where $\varepsilon$ is a geometric constant depending on the indenter used for the study: $\varepsilon = 0.72$ for a conical punch, $\varepsilon = 1$ for a flat punch, and $\varepsilon = 0.75$ for a paraboloid of revolution. The standard value of $\varepsilon = 0.72$ is used for the further calculations.

Using equation (3) to approximate the sink-in displacement of the contact periphery around the Vickers tip, the contact depth at maximum load $h_{c_{\text{max}}}$ can be determined with equation (2):

$$h_{c_{\text{max}}} = h_{\text{max}} - \varepsilon \frac{P_{\text{max}}}{S_{IT}}. \tag{4}$$
Projected contact area $A_{P\text{max}}$ described in Figure 6, at maximum load contact depth can be then determined using standard geometry of Vickers indenter tip, with the half-included angle of the pyramid $\theta = 68^\circ$:

$$A_{P\text{max}} = 4\tan^2\theta \left( h_{\text{max}} + \frac{P_{\text{max}}}{S_{IT}} \right)^2,$$

where $h_{\text{max}}$ is the maximum load contact depth.

Once the contact area is determined, the indentation hardness is then computed using equation (1).

Effective indentation modulus is then acquired using measured unloading stiffness and projected contact area:

$$S_{IT} = \frac{2\beta}{\sqrt{\pi}} E_{\text{eff}} \sqrt{A_{P\text{max}}},$$

where $\beta$ is a correction factor depending on the geometry of the indenter. It plays a major role in measuring precisely indentation modulus and hardness. Although many studies have been carried out to find an accurate value of this factor, $\beta = 1.012$ is used as determined by King [12] for a square-base indenter.

As elastic displacement occurs in both indented material and indenter during indentation, the indenter mechanical behavior has to be taken into account to determine the material indentation modulus $E_{IT}$:

$$\frac{1}{E_{\text{eff}}} = \frac{1 - \nu_i^2}{E_i} + \frac{1 - \nu_e^2}{E_e},$$

4 High temperature mechanical properties of epoxy resin

4.1 DMA results

DMA tests were carried out both on aged and unaged RTM6 epoxy resin samples to characterize global modulus (Fig. 7). The samples have the following dimensions: 45 mm long, 5 mm wide, and a thickness of 3 mm.

Storage modulus of the unaged resin decreases from 3.1 to 2.2 GPa, while storage modulus of the aged resin decreases from 3.1 to 2.5 GPa, when the temperature rises from 30 to 150°C.

DMA tests show that ageing of the epoxy resin slightly increases global mechanical properties of the sample which is due to oxidized layer formed on the surface of the sample. Young’s modulus is higher on aged RTM6 sample, due to anti-plasticization effect which has been studied by different authors [13,14]. Nevertheless, glass transition temperature would be lower for aged sample.

To explain these global results, nanoindentation tests are performed in order to test local properties on aged and non-aged materials from the edge to the center of the sample from 30 up to 150°C.

4.2 Effect of ageing

Indentation tests on aged and non-aged resins show in Figure 8 that Young’s modulus is increasing when the distance to the edge is less than 100 micrometers whereas Young’s modulus is constant on the non-ageing resin.

Polymer network becomes more rigid as oxygen diffuses in the sample. This phenomenon of thermo-oxidation affects the external surface of the sample. Thermo-oxidation effect can also be studied by weight loss measurement under oxidizing atmosphere [15] (Fig. 8).
4.3 Unaged resin characterization

Elevated temperature nanoindentation tests drawn in Figure 9, up to 150 °C are carried out in the center of an unaged sample of cured RTM6 epoxy resin. Indentation modulus decreases from 3.6 to 2.5 GPa when the temperature rises from 30 to 150 °C.

Under glass transition temperature Tg, viscous effect of polymeric material leads to a decrease of the value of indentation modulus. This effect is due to polymer network degradation, with breakage of polymer chains.

4.4 Aged resin characterization

Elevated temperature nanoindentation tests are presented in Figure 10. Tests are then carried out on a RTM6 epoxy resin sample aged for 5000 h at 130 °C to characterize local modulus near the edge of the sample. Ageing of the polymer in presence of oxygen creates a layer of oxidized resin on the first 100 μm underneath the surface and Young’s modulus is decreasing significantly in the thickness of the oxidized layer.

A maximum of 6 GPa is measured near the edge. No significant variation of the modulus in the oxidized layer in the first 100 micrometer thickness is shown when the temperature is increasing. Indentation modulus of the core of the sample, however, decreases to reach 3.0 GPa when the temperature is evolving up to 150°C.

After ageing, the thickness of the oxidized layer is growing asymptotically and oxidation growing is blocked. Evolution of temperature has no effects on Young’s modulus of the oxidized layer because polymer chains do not evolve anymore. In the core of the sample, Young’s modulus is decreasing with temperature increasing as observed on non-aged resin, due to degradation of polymer networks.

5 Conclusion

This study focuses on an experimental investigation of the evolution of gradient mechanical properties of an epoxy resin matrix with thermal ageing and with temperature test. The variation of Young’s modulus of the oxidized layer at the surface of the sample is not clearly affected by the increase in temperature because polymer chains do not evolve anymore, whereas Young’s modulus of the core of the sample is decreasing significantly with the temperature due to the viscous effect created by polymer network degradation which is also observed on unaged resin.

These new results show the opportunity of temperature nanoindentation set up to perform tests under severe conditions to determine gradient mechanical properties with a good accuracy of measurement.

References

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