A Comparative Study of Dispersion Techniques for Nanocomposite Made with Nanoclays and an Unsaturated Polyester Resin

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Received 12 April 2011; Revised 19 July 2011; Accepted 23 August 2011

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Over the last few years, polymer/clay nanocomposites have been an area of intensive research due to their capacity to improve the properties of the polymer resin. These nanocharged polymers exhibit a complex rheological behavior due to their dispersed structure in the matrix. Thus, to gain fundamental understanding of nanocomposite dispersion, characterization of their internal structure and their rheological behavior is crucial. Such understanding is also key to determine the manufacturing conditions to produce these nanomaterials by liquid composite molding (LCM) process. This paper investigates the mix of nanoclays particles in an unsaturated polyester resin using three different dispersion techniques: manual mixing, sonication, and high shear mixing (HSM). This paper shows that the mixing method has a significant effect on the sample morphology. Rheology, scanning electron microscopy (SEM), and differential scanning calorimetry (DSC) characterization techniques were used to analyze the blends morphology and evaluate the nanoclays stacks/polymer matrix interaction. Several phenomena, such as shear thinning and premature polymer gelification, were notably observed.

1. Introduction

Recent advances in the composite materials field are related to the addition of nanoparticles such as carbon nanotubes or nanoclays to improve thermal, mechanical, or electrical properties. Nanoparticle additives, like nanoclays, are widely used in various industries such as cable coatings, adhesives, inks, pharmaceuticals and automobiles [1, 2]. One of the most common nanoclay forms is MMT layered silicate with a particle thickness of 1 nm and 70 to 100 nm crosswise silica platelets [3]. The choice of montmorillonite nanoparticles in previous researches is mainly due to the fact that they are commonly available in nature and inexpensive. A minimal content of such additives between 1 to 6% wt can improve the properties of the polymer matrix by increasing flexural modulus by up to 31% and lowering the coefficient of linear thermal expansion by 66% [1, 4, 5]. However, the incorporation of nanoparticles into the liquid matrix is still a challenge, because it requires proper dispersion and exfoliation. Nanoclays are widely used in thermoplastic matrices, but only few studies report to their addition in polyester thermoset resins. This explains the lack of results on the thermal and mechanical properties and especially on the rheology of the mix.

Rheology is a widely used evaluation method for detecting the presence of interconnected structures. This technique seems to be relevant for the study of the dispersion state which defines the nanostructure of the mixture between the conventional, the intercalated, or exfoliated nanocomposite as illustrated in Figure 1. However, in practice, the final mixture would probably be a combination of these three morphologies, where the best-case scenario is the fully exfoliated structure [1, 6]. Due to the high aspect ratio of the platelets, a small percentage of nanoclay particles properly dispersed in the matrix can generate a very large surface...
area for polymer/filler interactions [1, 6]. Many factors can influence the dispersion and exfoliation of the nanoparticles in the polymer. The final properties of the nanostructure will mainly depend on the choice of the mixing technique and the resulted degree of exfoliation of the nanoclay platelets [7, 8]. Techniques such as in situ polymerization, solution mixing, or sonication are widely used to disperse nanoparticles in a liquid. The latter technique in particular seems to be relatively effective to obtain an exfoliated structure [1, 9, 10]. The changes of morphology of the mix are associated to the dispersion of the nanoparticles in the liquid matrix. When using a Newtonian polymer, the morphology change is detectable with rheology analyzes by the apparition of a shear-thinning behaviour [1, 7, 11, 12]. This non-Newtonian behaviour can be attributed to various factors such as the change in the nanoparticles volume fraction, shape, and size or size distribution [6, 12]. This decrease of viscosity is due to the reorientation of the layered silicate (MMT) in the direction of flow in response to the external applied shear [8, 13, 14]. The degree of the shear-thinning can then be used as an indicator of the exfoliation state of the nanoclays inside the polymer matrix; a steeper slope can be associated to an exfoliated mixture [7, 8]. When the shear stress is released after testing, a restructuring to the original disorganized structure of the nanoclay is initiated. The viscosity of the blend gradually go back to its original steady-state value [13].

The presence of layered silicates in nonaqueous polymers changes the viscoelastic behavior of the unfilled matrix from liquid-like \((G'\omega^2)\) to solid-like \((G'\omega^0)\) because of the formation of a three-dimensional percolating network of exfoliated or intercalated stacks [15]. This gel-like behavior is a direct consequence of the highly anisotropic nature of the nanoclays which prevents their free rotation and the dissipation of stress [16]. This superstructure formation will then be a liquid. The latter technique in particular seems to be effective to obtain an exfoliated structure [1, 9, 10]. The changes of morphology of the mix are associated to the dispersion of the nanoparticles in the liquid matrix. When using a Newtonian polymer, the morphology change is detectable with rheology analyzes by the apparition of a shear-thinning behaviour [1, 7, 11, 12]. This non-Newtonian behaviour can be attributed to various factors such as the change in the nanoparticles volume fraction, shape, and size or size distribution [6, 12]. This decrease of viscosity is due to the reorientation of the layered silicate (MMT) in the direction of flow in response to the external applied shear [8, 13, 14]. The degree of the shear-thinning can then be used as an indicator of the exfoliation state of the nanoclays inside the polymer matrix; a steeper slope can be associated to an exfoliated mixture [7, 8]. When the shear stress is released after testing, a restructuring to the original disorganized structure of the nanoclay is initiated. The viscosity of the blend gradually go back to its original steady-state value [13].

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The present study focuses on the mixing techniques and the understanding of the dispersion of nanoclays in unsaturated polyester resin. The main challenge is to achieve exfoliation of large stacks of clay nano-platelets into single layers keeping in mind the manufacturing process limitations. A previous study [31] already identifies the main process parameters to consider for proper injection of a nanoclay reinforced resin and impregnation of natural fibers by LCM. This work has also shown improved mechanical and flammability properties of nanoclay reinforced UP. Three dispersion methods were investigated on this study: manual mixing, sonication, and high shear mixing will be investigated. The dispersion state of nanoparticles in the
matrix will be evaluated using both rheology and SEM analyses. The catalytic effect of the nanoclays will also be investigated by gel time measurements and cure kinetics study using rheology and M-DSC techniques. The main objective of this research is to identify the most efficient dispersion technique and its impact on rheological and cure kinetics on nanocharged UP matrix keeping in consideration future manufacturing possibilities.

2. Experimental

2.1. Material. In this work, an unsaturated polyester petroleum-based resin (UP) R937-DPE24 from AOC was used, which has an average viscosity of 0.1804 Pa.s at 23°C. The resin was prepromoted with cobalt ethylenediyolate and initiated using 1.5 phr of methyl ethyl ketone peroxide (MEKP 925) from Norox. The resin was reinforced using 3% wt of Cloisite 30B nanoclay from Southern Clay Products. These nanoparticles are organically modified layered magnesium aluminum silicates and their properties are summarized in Table 1.

2.2. Nanoclay Dispersion. A good dispersion is a key challenge to achieve the best possible combination of matrix-nanoparticles. For that reason, a new dispersion technique using high shear mixer was investigated in this work and compared to sonication and manual mixing. The choice of the 3% wt Cloisite 30B was notably based on previous researches [31, 32] showing that these type of nanoparticles are easier to disperse due to their chemistry providing general improvement on the matrix properties. Furthermore this type of nanoclay was investigated in an earlier study focusing on composites manufacturing [31], and the mass fraction of 3% wt was considered adequate for RTM processing.

The sonication and manual preparations are detailed in a previous study [31]. In these techniques, the nanoclays were incorporated and predispersed in styrene first. The resin was then added to the mix and the styrene in excess was evaporated using high-speed mechanical stirring. The amount of each component was weight controlled. Since the HSM is performed in a sealed chamber, the nanoclays were directly incorporated into the resin. The gap in the geometry interaction chamber has been fixed between 50 and 100 microns, and the resin was circulated inside the chamber at high pressure. The size of particles agglomerates was significantly reduced with this mixing technique resulting in a well-dispersed and homogeneous blend. Figure 2 summarizes the blend preparation procedures with the different approaches. The letter A refers to sonication technique whereas the letters B to high shear mixing, Pure and nanocharged resins were mixed by three methods in order to take into account the possible effect of the mixing technique on the resin itself. The temperature and pressure were maintained at 23°C and 1 atm for (A0) and (A1) blends to limit the possible formation of microgels or styrene evaporation.

2.3. Rheology and Electron Microscopy. Rheology experiments were conducted using a combined motor and transducer (CMT) rheometer MCR501 from Anton-Paar. Both steady shear and small amplitude oscillatory shear (SAOS) measurements were carried out using two types of geometries: parallel plates of 25 cm diameter with a constant gap of 1 mm for high-viscosity blends and concentric cylinders for low-viscosity blends. All experiments were conducted at 23°C and before measurement the sample was left in the geometry for stabilization for 20 minutes. Plastic paraffin films (parafilm) were placed above the mixtures to limit styrene evaporation during the stabilisation stage. This film limits the air contact of the liquid resin and styrene evaporation can be neglected. The steady shear measurements were performed using a shear rate varying from 0.1 to 1000 s⁻¹.

For steady shear viscosity, the non-Newtonian viscosity η is defined as follows [19, 33]:

\[ \eta(\dot{\gamma}) = \frac{-\tau_{21}}{\dot{\gamma}}, \]

where \( \tau_{21} \) is the shear stress, and \( \dot{\gamma} \) the shear rate applied. The SAOS experiments were carried out under strain amplitudes of 1%, inside the linear viscoelasticity (LVE) domain. Frequency sweep test measurements were performed at frequencies varying from 0.1 to 100 Hz. A strain wave was imposed to the sample and the SAOS shear stress response \( \tau_{21}(t) \) defined as follows [19, 33]:

\[ -\tau_{21}(t) = (\tau_0 \cos(\delta)) \sin(\omega t) + (\tau_0 \sin(\delta)) \cos(\omega t), \]

where \( \tau_0 \) is the stress amplitude, \( \omega \) the angular frequency, and \( \delta \) the phase difference between the strain wave and the stress response. For elastic solids, the shear stress is directly proportional to the strain imposed as defined by Hooke’s law:

\[ \tau_{21} = -G \cdot \gamma_21 = -G \cdot \gamma_0, \]

where \( \gamma_0 = \dot{\gamma}_0/\omega \) is the strain amplitude and \( G \) the elastic modulus. Equations (2) and (3) can be combined to obtain the SAOS material function:

\[ -\frac{\tau_{21}}{\gamma_0} = G' \sin(\omega \cdot t) + G'' \cos(\omega \cdot t), \]

\[ \tan(\delta) = \frac{G''}{G'}, \]

where \( G' \) is the storage modulus and \( G'' \) the loss modulus, related, respectively, to the energy stored (elastic contribution) and dissipated (viscous contribution). The ratio between modulus is defined as the damping factor \( \tan \delta \). The complex viscosity \( \eta(\omega) \) is finally defined as

\[ \eta(\omega) = \frac{-\tau_{21}(t)}{\dot{\gamma}_{21}(t)}, \]

where \( \dot{\gamma}_{21}(t) \) is the SAOS strain rate.
In order to quality the size of the agglomerates and thus, evaluate the quality of the nanoreinforced blends, scanning electron microscopy (SEM) technique is used. A field emission gun scanning electron microscope (FEG-SEM-4700) from Hitachi is used at an operation voltage of 2 kV. FEG-SEM was used to investigate the distribution of clay in the Epoxy matrix. The smooth and flat surface of specimens were prepared using Ultracut FC microtome (Leica) with a diamond knife and then coated with platinum. This device cuts thin slices of nanocomposite to the desired thickness and the observed area is around 100 μm.

2.4. Calorimetry. The addition of nanoclay particles in the UP resin can cause premature cross-linking of the matrix which could be the source of composite part failure and restrict the manufacturing capabilities. Consequently, it is important to understand the effects of nanoclays on the resin cure kinetic. In order to see the possible catalytic effect of nanoparticles, calorimetry experiments were performed on blends A and B and on the resin alone. First, the samples were isothermally maintained at 70°C for different periods of time up to 24 hours in order to study the effect of the storage conditions on the nanocharged mixtures. Afterwards, the MEKP catalyst was added to the blend and samples were cured on the DSC at a dynamic ramp of 10°C/min from room temperature to 250°C. The instrument used on this study is a modulated differential scanning calorimeter (M-DSC) Q1000 from TA Instrument. This instrument has the great advantage of being able to separate the heat flow related to the cure kinetic from the changes in thermodynamic heat capacity. The instantaneous heat generated during the polymerization reaction can be expressed as follows [26, 34]:

\[
\frac{dH}{dt} = \Delta H_R^\alpha \frac{d\alpha}{dt},
\]

(6)

where \(\Delta H_R^\alpha\) is the total heat of reaction measured by M-DSC and \(d\alpha/dt\) the reaction rate. If the diffusion of chemical species is neglected, the reaction rate is assumed to be a unique function of the degree of conversion \(\alpha\) and temperature \(T\) and this expression takes the form of

\[
\frac{d\alpha}{dt} = f(T, \alpha), \quad \alpha = \int_0^t \frac{d\alpha}{dt} \cdot dt
\]

(7)

and the total heat of reaction corresponds to the area under the nonreversing heat flux curve measured by M-DSC [26, 34].

3. Results and Analyses

3.1. Rheology Analyses. In this work, the quality of the dispersion of nanoclays in the liquid resin was evaluated from rheology tests knowing that a high initial viscosity and strong shear thinning behavior are often associated to a high level of exfoliation of the nanoclay platelets [1, 6]. This rheological behavior illustrates the ability of the nanoclays to “interact” with the polymer matrix and is intimately related to the clay type, its surface treatment, and affinity with the polymeric matrix [6, 32].

Figure 3 illustrates the viscosity change with shear rate for eight samples dispersed by three different mixing techniques. As shown, the resin viscosity increases after the addition of the nanoparticles with the three mixing techniques studied in this work. The initial viscosity can rise over 3 decades. The mixing techniques does not influence the pure resin viscosity as illustrated by the A0 and B0 curves showing the same Newtonian behavior with a viscosity around 0.20 Pa.s, close to the manufacturer data of 0.1804 Pa.s. For the manually mixed blend, the size of particles is probably still quite large due to high probability of aggregates. The sonication mix (A1) results in a slight non-Newtonian shear thinning behavior with an increase of the initial viscosity to 0.80 Pa.s. This can be attributed to a better dispersion and possible intercalation of the nanoclay platelets. On the other hand, the high shear mixing (B1) blends show a strong shear thinning behavior with an initial viscosity varying from 70 to 250 Pa.s. The initial viscosity increases with the number of passes in the high shear chamber. However, at higher shear rates (20 s⁻¹ and over), the viscosity of the nanoreinforced resin is below 10 Pa.s. This value has been previously estimated as the processability limit to manufacture composite parts by RTM [31]. In the case of the high shear mixing blends, the number of passes will have an important influence on the initial viscosity of the nanocharged resin. After two passes, the nanoclays platelets seem to reach their maximum possible exfoliated state, the interactions between the platelets are

![Diagram](image_url)
to be noted that no accurate values of $G'$ were detected for frequencies below 0.1 rad/s due to the precision of the rheometer. In Figure 4(b), small variations of tan $\delta$ for the HSM mixtures at low frequency are observed, which confirms the gel-like structure of these blends. However, these are considered to be weak gels, because the ratio between the dissipative ($G''$) and elastic energy ($G'$) is low, between 0.1 and 0.7. So the viscous nature still dominates the elastic behaviour in opposite to a strong gel (tan $\delta$ below 10$^{-2}$) [35]. At almost $\omega = 10$ rad/s, the tan $\delta$ value reaches a maximum for the (A1) and manual blends (peak of the bell shape). This marks a transition to a more dissipative behaviour, meaning a transition from a gel-like state to a liquid-like state. This behaviour can be related to the alignment of the platelets in the flow direction. For the (B1) blends, a slight increase of tan $\delta$ and $G'$ at 10 rad/s can be observed, indicating the beginning of a morphological change. At higher frequencies ($\omega > 62.8$ rad/s), $G'$ seems to tend to the same value for all systems. It was finally found that at high frequencies, the $G'$ response is dominated by the matrix since the values of all mixtures trend to 1000 Pa at 628 rad/s [3].

To validate the rheology results showing the differences in dispersion of the nano-platelets, visual observation was performed using SEM and micrographs of the blends as illustrated in Figure 5. Note that the unidirectional etchings on the micrographs are due to the blade of the diamond knife during sample preparation. The manual mixing (Figure 5(a)) shows important agglomerations in the order of 10 microns whereas they are diminishing around 3 microns for sonication (Figure 5(b)). Finally, the high shear mixing seems to be a relevant technique for nano-dispersion showing agglomerates smaller than 1 micron (Figure 5(c)). Figure 6 presents the initial storage modulus as function of the size of the agglomerates obtained after the dispersion by manual mixing, (A1), and (B1) techniques. The first method shows a high level of aggregation compared to the (B1) 5 passes blend which probably has a fully exfoliated morphology. As illustrated by the $G'$ data, there is an increase in elasticity of more than two decades between (A1) and (B1) blends. This is attributed to an important change in the internal structure of the nanoclay platelets and also confirming the gel-like 3D network structure.

3.2. RTM Shear Simulation and Process Limitations. In order to simulate an RTM injection and relate it to the reorganization of the nanoclays with the resin flow, a rheology cycle was imposed to the most viscous (B1) blend that underwent 10 passes through the HSM system. The choice of this particular blend was based on the fact that it is the one showing the most important morphological changes. The imposed shear cycle combines shear and rest periods as illustrated in Figure 7. This simulation is carried out by varying the shear rate three times. At the initial state (quasistatic), the shear rate is kept constant at 0.1 (1/s). The initial shear rate was chosen in order to reproduce the initial static state of the resin before injection. This step is followed by a sudden increase of the shear rate to 1000 s$^{-1}$ representing the resin injection into the mold cavity taking less than 25 seconds.
After this period, the resin is maintained on a static state (i.e., 0.1 (1/s)) for 25 minutes. During this quasistate period at constant shear rate, the resin viscosity increase from 1.8 Pa.s to 200 Pa.s. This increment is assumed to be linked to the nanoclays reorganization, since the shear rate is constant and no chemical reaction takes place (no hardener was added to the resin).

At the very beginning of the cycle, the nanocharged resin is at rest and the initial exfoliated structure of the platelets randomly disorganized. This leads to a very high viscosity of 250 Pa.s on the first two minutes. When the resin is injected at a shear rate of 1000 s\(^{-1}\), the viscosity decreases abruptly to 1.8 Pa.s and the original nano-structure is destroyed. The particles at this point are oriented in the fluid flow direction. After the filling is completed, at around 3 minutes, the shear is released and the viscosity gradually increases until it reaches its initial value after 10 minutes. There is a restructuration of the nanoparticles in the blend which disorganized themselves to their original exfoliated structure after this period. This rheology simulation gives precious...
In determining the gel time at the maximum of the tan δ study was successfully used for epoxy prepregs, consisting of the system being studied as well as the stoichiometry of the monomer necessitates the property of the final part. The gel point is another important feature of the polymer. The calculation of the gel time with this criterion results in a viscosity increase between 2 to 5 times from the initial value. At this point the resin will not flow and the resin viscosity curve has a slope of 10% \[26\]. This criterion corresponds to a high resin viscosity. In LCM process analysis, the processing gel time may be defined as the point at which the resin viscosity curve has a slope of 10% \[26\]. This criterion results in a viscosity increase between 2 to 5 times from the initial value. At this point the resin will not flow under normal injection pressures \[26\]. Figure 9(b) illustrates the evolution of the complex viscosity with time during cure. This processing gel time calculated from complex viscosity is in good agreement with the data obtained from actual manufacturing process. To obtain consistent results, the frequency of all experiments was set to 1 Hz in the linear viscoelastic region. Experiments were then conducted in time sweep mode as shown in Figures 8 and 9.

Figure 8 illustrates the evolution of $G'$, $G''$, and tan δ during the cure of the B1-HSM 5 passes blend at 23°C. It can be seen that $G'$ is always higher than $G''$ because of the gel-like structure of the blend. As a consequence, no crossover is observed between $G'$ and $G''$. At around 9 minutes from the beginning of cure, there is a sudden increase of the storage modulus to infinity which indicates an important change in the polymer structure. At this point, the tan δ curve reaches a maximum close to 1. This is an indication of the gel formation which denotes the beginning of the cross-linking of the polymer. The calculation of the gel time with this method was carried out on the resin/nanoparticles blends mixed with different techniques as illustrated in Figure 9(a) at the same temperature and frequency. As shown, the blends mixed with a high shear mixer result in the lowest gel times, between 3 and 15 minutes. Moreover, it is observed that the gel time decreases proportionally with the number of passes in the mixer for the HSM technique. As of the hand mixed and sonicated samples, their gel point is slightly inferior to the pure resin due to the supposed catalytic effect of the nanoclay. Since the nanoclay are not well dispersed with this techniques compared to HSM mixtures, the catalytic effect is limited.

The gelification point is often defined as the time at which the $G'$ exceeds $G''$ (i.e., tan δ = 1), but this point corresponds to a high resin viscosity. In LCM process analysis, the processing gel time may be defined as the point at which the resin viscosity curve has a slope of 10% \[26\]. This criterion results in a viscosity increase between 2 to 5 times from the initial value. At this point the resin will not flow under normal injection pressures \[26\]. Figure 9(b) illustrates the evolution of the complex viscosity with time during cure. This processing gel time calculated from complex viscosity is in good agreement with the data obtained from the nanocharged mixtures and it is defined as the required time passes).
3.3. Catalytic Effect of the Nanoclays. As illustrated by the gel time results, the addition of nanoclays has a nonnegligible impact on the polymerization process. In order to better understand this catalytic effect on the cross-linking reaction of the UP resin, samples of the different blends were isothermally maintained at 70°C for various dwell times. The catalyst was then added to the blend and the heat of reaction was analyzed with an M-DSC using a constant heating rate process. The residual heat of reaction was compared to the total heat of reaction generated during the cure of the pure resin. In addition, the samples exposed at 70°C were compared to the samples stored at −18°C, so-called control samples. Neat resin samples were exposed to the same isothermal condition to take into account the styrene evaporation which has an influence on the released heat of reaction. For that reason, a correction of the total heat is made for all stored samples. The nonreleased heat due the evaporation of styrene for the pure resin sample, \( \Delta H_{\text{pure resin}} \), during storage at 70°C was calculated as follows:

\[
\Delta H_{\text{pure resin}} = H_{\text{pure resin at } -18^\circ C} - H_{\text{pure resin at } 70^\circ C}. 
\]

The evaporation of the cross-linking agent (styrene) during storage lowers the total heat or reaction of the resin a quantity the maximum of \( \tan \delta \). The catalytic effect of the nanoclay tends to decrease the gel time due to the increased interactions between nanoclay platelets and the polymer resin. Table 2 resumes the gelification time for each system according to the mixing method. Figure 9(c) summarizes the gelification times calculated from the maximum of \( \tan \delta \) according to the mixing method. The gelification time has been reduced in 17 minutes between (A1) and (B1) blends. This is mainly due to an enhanced dispersion state and probable exfoliation, which results in a greater surface for interaction between the polymer matrix and the clays. This interfacial region acts as a catalyst and accelerates the curing reaction.

**Figure 9**: Gel time at 23°C UP resin with 3% wt of C30B mixed with various dispersion techniques: (a) method of maximum \( \tan \delta \), (b) 10% slope of \( \eta^* \), and (c) gel point tendency obtained with the maximum \( \tan \delta \).
Table 2: Gel time results using two different calculation methods.

<table>
<thead>
<tr>
<th></th>
<th>Maximum tan δ (min)</th>
<th>10% slope (min)</th>
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<tbody>
<tr>
<td></td>
<td>Figure 9(a)</td>
<td>Figure 9(b)</td>
</tr>
<tr>
<td>Pure resin</td>
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<td>42</td>
</tr>
<tr>
<td>Manual mixing</td>
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<tr>
<td>3% cloisite 30B</td>
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</tr>
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<td>1 pass</td>
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</tr>
<tr>
<td>5 passes</td>
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</tr>
<tr>
<td>10 passes</td>
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Table 3: Storage temperature influence on nanocharged resin using HSM and sonication dispersion techniques.

<table>
<thead>
<tr>
<th>Isothermal temperature (°C)</th>
<th>Dwell time (h)</th>
<th>Pure resin</th>
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<tr>
<td></td>
<td></td>
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<td>$ΔH_{res}$ (J/g)</td>
<td>$H_{res}$ (J/g)</td>
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<td>−18</td>
<td>24</td>
<td>279.8</td>
<td>—</td>
<td>259.7</td>
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<td>260.1</td>
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<td>12</td>
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<td>24</td>
<td>239.3</td>
<td>40.5</td>
<td>197.1</td>
</tr>
</tbody>
</table>

1 Calculation based on $H_{res}$ of the corresponding pure resin storage.

equals to $ΔH_{pure\ resin}$. As shown in Table 3, exposing the pure resin at 70°C, for long periods, results in a decrement of the heat of reaction of 9.5% and 14.5% after 6 and 24 hours, respectively. Therefore, for the nanocomposite samples maintained at 70°C, the total heat of reaction has to be compared to the one of the pure resin under similar thermal treatment ($H_{pure\ resin}$ at 70°C). Thus, the degree of cure associated to the catalytic effect of the nanoclays $α_{nano}$ was calculated based on the total heat of reaction generated during cure of the nanocomposites and the one of the pure resin:

$$α_{nano} = 1 - \left( \frac{H_{blend}}{H_{pure\ resin}} \right),$$

where, $H_{blend}$ is the heat of reaction of the nanocharged blend and $H_{pure\ resin}$ the heat of reaction for the pure resin, both at 70°C and at the same dwell time. Figure 10 shows the results of two MDSC tests of the sonicated blend (A1) maintained at −18°C and 70°C during 24 hours. It can be seen that the residual heat of reaction after thermal exposure has decreased by 10% at −18°C and by 17.6% at 70°C. This clearly indicates the influence of the nanoclays on the cure reaction of the unsaturated polyester resin. Table 3 presents the resulting degree of cure $α_{nano}$ after exposing the nanocomposites at these temperatures.

As for the HSM blend (B1), this catalytic effect of nanoclays is very important. The degree of cure $α_{nano}$ reaches 18% after 1 h of exposition and 38.5% after 2 h at 70°C. After this point, the resin gels and it is no longer possible to be add the catalyst. The differences in cure reaction between the (A1) and (B1) blends can be attributed to an increased dispersion and possible exfoliation of the HSM blend. This catalytic effect of nanoclay platelets is probably...
due to the existence of attractive forces between the clay and the polymer matrix. The cation-exchange capacity of the nanoclay influences the cure reaction. This acceleration is induced by the alkylammonium ions contained in the surface treatment of the nanocharges [20].

Figure 11 shows the results of the measured degree of cure $\alpha_{nano}$ for (A1) and (B1) blends as a function of exposition time at 70°C. A phenomenological autocatalytic model was used to model the cure reaction of both blends. The degree of cure was evaluated by an nth-order model as follows [26, 34]:

$$\frac{d\alpha}{dt} = K_A \cdot (1 - \alpha)^n,$$

(10)

where $\alpha$ is the degree of polymerization, $n$ the reaction order, and $K_A$ the rate constant given by an Arrhenius temperature dependence defined as

$$K_A = K_0 \exp\left(\frac{-E_A}{R \cdot T}\right),$$

(11)

where $K_0$ is the Arrhenius constant, $E_A$ is the activation energy, and $R$ is the ideal gas constant. Equations (10) and (11) are widely used to model the cure of many polymer systems such as polyester and epoxies. This model is used on the assumption that only one kind of reaction may represent the whole cure process [34]. The parameters of the proposed cure kinetics model were obtained by fitting the experimental data extracted from M-DSC tests and presented in Table 3. The coefficients of the two models are shown in Table 4. As illustrated in Figure 11, there is a good agreement between the M-DSC experiments and the proposed models. This indicates that an autocatalytic reaction of the polymer can be started due to the presence of nanoclay particles. Moreover, these results demonstrate that the degree of exfoliation accelerates the autocatalytic cure reaction. It is then concluded that the exfoliation of MMT changes the cure rate of the UP resin tested on this work. Similar results were reported in the past for epoxy resins [37]. This could also indicate that the interfacial region between the clay and the polymer may dominate the properties of the blend.

### 4. Conclusion

The scope of this work was to study the nanoclays dispersion in an UP resin and by means of three different mixing techniques. Rheology tests based on shear and SAOS experiments revealed the differences between the manual mixing, sonication, and high shear mixing techniques. Notably, the shear thinning behavior is a result of the morphological change of the blend due to the rearrangement of the silicate nanoclays. Furthermore, the intercalated and exfoliated structure has shown a gel-like structure of high shear mixed blends. This is, however, a weak gel since $\tan \delta$ is in the order of $10^{-1}$. This gel-like structure is a direct consequence of the interaction between the polymer and the nano-platelets leading to the creation of a 3D network in the liquid.

The high shear mixing has the highest elastic modulus (100 Pa) and initial viscosity (up to 250 Pa.s compared to other mixing techniques). However, the viscosity decreases at high shear rate due to the reorientation of the nanoclays. The viscosity reaches 2 Pa.s at high shear which makes it possible to process the blend by RTM. The dispersion was also verified by SEM microscopy showing agglomerates of 10 microns for the manual mixing and less than 1 micron for the high shear technique. The nanoclays seem also to have a catalyst effect on the resin system, showing a reduction of the gel time due to their exfoliation state. The neat resin showed a gel time of 45 minutes whereas the 10 passes HSM blend jellified at around 3 minutes. This phenomenon has to be taken into account for proper composites molding. The DSC study confirms the catalytic effect of the nanoclays showing a degree of cure of 38.5% for the high shear mix after 2 hours of exposition at a 70°C.

This study illustrates how well-dispersed nanoclays in UP resin bring many interesting structural changes in the polymer matrix. This work also shows that processing may be achievable since the time to orient the nanoparticles in the direction of fluid flow is within seconds and the restructuration to their initial random disorganized structure takes few minutes. Future work will focus on the use of HSM technique for manufacturing bio-composite parts improved by nanoclay reinforcing of the biorein.
Abbreviations

FEG: Field emission gun (scanning electron microscope)
G': Elastic modulus
G'': Loss modulus
HSM: High shear mixing
LCM: Liquid composite molding
LVE: Linear viscoelastic
M-DSC: Modulated differential scanning calorimetry
MMT: Montmorillonite
phr: Parts per hundred
RTM: Resin transfer molding
SCRIMP: Seeman composite resin infusion molding process
SEM: Scanning electron microscopy
TEM: Transmission electronic microscopy
UP: Unsaturated polyester resin
VARI: Vacuum assisted resin infusion
% wt: Weight percentage
XRD: X-Ray diffraction
ω: Frequency (rad/s)
tan δ: Damping factor.

Acknowledgments

The authors acknowledge the support provided by the Chair of High Performance Composite (CCHIP) and the Center for Applied Research on Polymers and Composites (CREPEC). The financial contribution of the Natural Sciences and Engineering Research Council of Canada (NSERC) is also greatly appreciated.

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