Nanostructured Epoxy/POSS composites: Enhanced Materials for High Voltage Insulation Applications

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ABSTRACT

In this study, the dielectric and thermal properties of nanostructured epoxy/POSS (Polyhedral Oligomeric Silsesquioxanes) composites were investigated, using a reactive Triglycidylisobutyl-POSS (TGIB-POSS) additive from 1 up to 10 wt%. POSS has been successfully dispersed at a molecular level for low content composites, which show a remarkably improved resistance to corona discharges, with up to 60% less eroded sample volume, along with significantly increased dielectric breakdown strengths and thermal conductivities. Epoxy/POSS composites containing 5 wt% and more of the POSS additive exhibit agglomerations, which have been observed by SEM. Furthermore, dielectric spectroscopy revealed additional interfacial loss peaks for such composites containing 5 wt% POSS and more, in addition to the α- and β-peaks known for epoxy.

Index Terms — AC breakdown strength, dielectric spectroscopy, epoxy, electrical discharge, interface, polymer composites, POSS, surface erosion, thermal conductivity.

1 INTRODUCTION

EPOXY resins are an important dielectric material in the high voltage insulation industry, and they are widely used in insulation systems for bushings, rotating machines and cable terminations, to name a few. However, as an organic material, it is prone to degradation due to electrical discharges and usually features rather low thermal conductivities, which can constitute a limiting factor for power ratings. These deficient properties have been counteracted in the past by the addition of micrometric inorganic filler materials, such as silica, which can render the resulting composite more resilient to electro-thermal stresses and improve parameters such as the thermal conductivity.

In the last decade though, much has been reported about a novel class of dielectric materials, namely polymer nanocomposites or nanodielectrics, which feature nanometric filler particles [1, 2]. However, such nanodielectrics can only unfold their full potential, when good dispersion and distribution of those filler particles within the matrix are achieved. Albeit, due to the incompatibility of inorganic particles with the organic polymer, dispersion and distribution are regularly compromised, resulting in agglomerations of nano particles, often leading to agglomerates with submicrometric or micrometric dimensions. In order to enhance the interaction between inorganic filler particles and the organic matrix, and hence, improve the dispersion of such particles in polymers, their functionalization has become rather common. Surface treatments with coupling agents, such as silane [3-5], or more complex processes, including grafting organic brushes on particle surfaces [6], have been explored with varying degrees of success.

As an alternative to nanometric silica, Polyhedral Oligomeric Silsesquioxanes (POSS) could be of interest for electrical insulation applications, as it is a hybrid material, consisting of an inorganic silica-like core surrounded by organic side-groups. These organic groups can be either reactive or non-reactive and therefore offer compatibility with a multitude of polymers, depending on the specific organic side-group configuration. POSS, being a hybrid inorganic-organic material, further combines the beneficial material properties of ceramics, such as inertness and electrical

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insulating properties, with the processability of polymers. It was previously denoted as a “Nano-building-block”, because of its crystalline structure [7, 8], enabling POSS to organize a polymeric matrix through its reactive groups on a nanometric scale. POSS molecules, commonly with sizes below 2 nm, are assumed to be able to dissolve within a polymer without forming agglomerations known from conventional nanoparticles [9].

The successful incorporation of POSS into epoxy has been shown to significantly improve the composite’s dielectric performance, in terms of corona resistance [10-12] and dielectric breakdown strengths [13-15]. Furthermore, higher thermal stabilities of POSS composites have been reported in [8, 16], as well as improved thermo-mechanical properties of such nanostructured materials [7].

The aim of this study was to investigate the dielectric and thermal properties of epoxy/POSS composites, by incorporating a reactive Triglycidylisobutyl-POSS, which we intended to disperse at a molecular level within the matrix.

2 MATERIALS AND SAMPLE PREPARATION

DER332 epoxy resin (DOW Chemicals) and Jeffamine D230 curing agent (Huntsman Corp.) were used as the base epoxy system in a ratio of 1000:344 parts (epoxy to curing agent). Liquid Triglycidylisobutyl-POSS (Hybrid Plastics) was used as an additive to fabricate composites. Triglycidylisobutyl-POSS (TGIB-POSS) has three mono-directionally oriented epoxy groups which are attached to the cage-like silica core (see Figure 1). These epoxy groups are compatible with those of the epoxy system used and can therefore form covalent bonds with the matrix, as schematically shown in Figure 1. Due to the fact that each TGIB-POSS molecule features three additional epoxy groups that will consume curing agent during the reticulation process, a distinct stoichiometric ratio for each epoxy/POSS composite was established, in order to match the amount of reactive epoxy groups with the amine groups of the curing agent within the composite.

During the compounding process, the liquid TGIB-POSS, whose viscosity is similar to that of the epoxy system, was dispersed in the epoxy resin using a magnetic stir on a hot plate at 40 °C. Subsequently the curing agent was added to the mixture. The complete mixing procedures were conducted under vacuum, in order to ensure proper degassing of the liquid composites before they were cast in stainless steel molds. All test specimens were cured for six hours at 100 °C under vacuum, followed by a gradient cool down to ambient temperature. A release agent (QZ13) was applied on the mold before the epoxy was cast to facilitate the removal of the samples after the curing process. Subsequently, the samples were post-cured for 12 hours at 130 °C, also under vacuum. More details about the fabrication process and the stoichiometry can be found in [17].

Along with neat epoxy resin specimens, epoxy/POSS composites with TGIB-POSS contents of 1, 2.5, 5 and 10 wt% were fabricated. The POSS composites are further referred to as e.g. ePOSS 1wt%, for the epoxy/POSS composite containing 1 wt% of TGIB-POSS, and so on.

3 EXPERIMENTAL METHODS

3.1 MICROSTRUCTURE ANALYSIS

The test specimens were mounted in epoxy resin and subsequently polished with grain sizes down to 50 nm to assure a smooth sample surface for Scanning Electron Microscope (SEM) observations. A TM3030 tabletop SEM (Hitachi High Technologies America) was used for low magnification pictures (x5k), whereas a SU-70 high resolution SEM (Hitachi High Technologies America) was used for micrographs with magnifications of x20k and x50k, to further investigate some features found on the composites with higher POSS contents.

3.2 DIFFERENTIAL SCANNING CALORIMETRY

Glass transition transition temperatures were measured by Differential Scanning Calorimetry (DSC) with a heat flow calorimeter (Q20, TA Instruments) on samples which had not previously been exposed to the thermal post-treatment. Each sample was subjected to two thermal cycles with a heating rate of 10 °C/min from 20 °C to 250 °C, followed by a gradual cool down to 20 °C at 10 °C/min. The glass transition temperatures Tg were obtained from the second heating cycle to ensure no thermal history of the samples.

3.3 AC BREAKDOWN STRENGTH

The dielectric breakdown strength was measured based on the ASTM D 149 standard [18]. Samples were immersed in transformer oil (Luminol TR-i) and placed between two symmetrical ball tip electrodes with diameters of 4 mm. The short-term test was used, where a 60 Hz voltage with a ramp rate of 2 kV/s was applied between the electrodes until breakdown occurred. On each sample type, 15 breakdowns
were performed, after which the ball tips were exchanged and replaced with new ones to avoid pitting. The breakdown test specimens had an average thickness of 130 μm ± 5%.

3.4 RESISTANCE TO CORONA DISCHARGES

To evaluate the resistance to corona discharges, each sample was subjected to partial discharges (PD) using a point-to-plane geometry operated in open air. A sinusoidal voltage of 4 kVrms at a frequency of 300 Hz was applied between the high voltage rod electrode, bearing a 4 mm diameter tungsten carbide ball tip, and the ground electrode. The samples were placed centrally on the ground electrode framed by an acrylic base plate, with the tip of the rod electrode facing the sample surface. The high voltage electrode rod and the sample surface were separated by a 200 μm air gap (see Figure 2). All test specimens were exposed to electrical discharges for a duration of 30 hours, which is equal to exposures for 150 hours at 60 Hz and 180 hours at 50 Hz equivalent time, respectively.

The resistance to erosion due to corona discharges was subsequently evaluated by the use of a mechanical profilometer, based on the eroded volume calculated from the obtained data. The values of the eroded sample volumes can thus be seen as inverse reference values of the resistance to partial discharge. Before the samples were analyzed with the profilometer, they were carefully cleaned in an ultrasonic bath in order to remove the debris caused by the PD.

3.5 THERMAL CONDUCTIVITY

The thermal conductivity was obtained with a DTC-25 guarded heat flow meter (TA Instruments) in accordance with the ASTM E 1530 standard [19]. Test specimens with 50.8 mm (2 inches) diameter were subsequently placed between the upper heating plate and the lower cooling plate of the device. A pressure of 20 psi applied on the upper plate ensured intimate contact between the sample, the heating and the cooling plates. A heat flux transducer in the lower plate measured the heat flow Q through the sample. The thermal conductivity λ is defined by (2), with the sample surface A, its thickness t and the temperature drop over the sample volume ΔT. The measurements were conducted for a medium sample temperature of 25 °C. In order to assure the thermal stability of all test specimens during the measurements, each sample was left inside the test chamber for two hours before the thermal conductivity values were obtained.

\[ \lambda = \frac{Q/A}{\Delta T/t} \]  \hspace{1cm} (1)

3.6 DIELECTRIC SPECTROSCOPY

The frequency- and temperature-dependent dielectric responses of the samples were obtained by Broadband Dielectric Spectroscopy (BDS, Novocontrol) for a temperature range from 20 °C to 100 °C (± 0.1 °C), in several distinct isothermal steps. The test specimens were subjected to an excitation voltage of 1 V with a frequency ranging from 0.1 Hz to 1 MHz. The BDS measurements were performed on 40 mm diameter samples on a parallel plate electrode setup. The complex permittivity ε* as given in (1) was retrieved, with the real permittivity ε' representing the storage part, and ε'' the imaginary or dielectric loss part.

\[ \varepsilon^*(f, T) = \varepsilon'(f, T) - j\varepsilon''(f, T) \]  \hspace{1cm} (2)

4 EXPERIMENTAL RESULTS AND DISCUSSION

4.1 MICROSTRUCTURE ANALYSIS

In Figure 3, the SEM micrographs of neat epoxy and the POSS-composites are presented. The left column shows the microstructure obtained with low magnification (x5k) on the TM3030 SEM. The next two columns represent SEM micrographs obtained with the high resolution Hitachi SU-70 microscope using x20k and x50k magnification, respectively.

For the ePOSS 1wt% and ePOSS 2.5wt% composites (Figure 3b and Figure 3c), the micrographs obtained up to a magnification of x50k did not show any particular trace of the presence of POSS additive, and hence, the revealed microstructures resemble those obtained for the neat epoxy (Figure 3a). The POSS additive thus appears to be dispersed into the epoxy matrix at a molecular level or to very small agglomerates of less than 10 nm if any, which allows us to consider these specific materials as actual nanocomposites.

However, for the composites with higher filler contents, viz. ePOSS 5wt% (Figure 3d) and ePOSS 10wt% (Figure 3e), agglomerations of submicrometric dimensions (< 1 μm) were found. Further, both the size and the occurrence of these agglomerated structures increased with the POSS content.

4.2 DIFFERENTIAL SCANNING CALORIMETRY

The resulting graphs obtained from the DSC experiments are presented in Figure 4. The derived glass transition temperatures Tg are summarized in Table 1. It was found that incorporating POSS into the epoxy matrix led to a slight diminishment of Tg for all composites, compared to the neat polymer. The highest difference was found for the ePOSS 10wt% composite, which exhibited a glass transition temperature of almost 6 °C lower than that of neat epoxy.
Figure 3. SEM micrographs with increasing magnification from left to right, for a) neat epoxy, b) ePOSS 1wt%, c) ePOSS 2.5wt%, d) ePOSS 5wt% and e) ePOSS 10wt%.
It is common understanding that $T_g$ is closely related to the mobility of the molecular chains of the epoxy network. The mobility of the chains will further depend on factors such as the degree of cross-linking, chain lengths, flexibility of side-groups and chains, as well as free volume, as has been elaborated in detail in this extensive review [20]. Considering the slight decrease in $T_g$ for the POSS composites compared to the neat polymer, it is assumed that the cross-linking of the epoxy matrix is hindered in the presence of Triglycidylisobutyl-POSS. Additionally, the mono-directional alignment of epoxy groups on these particular POSS molecules might lead to shorter polymer chains, given that the chain “growth” during the cross-linking process is limited to one side of the TGIB-POSS molecule. Also, auto cross-linking between TGIB-POSS molecules themselves could further contribute to the reduction of chain lengths and cross-linking degree of the resulting epoxy/POSS network. Lastly, it should be mentioned that for the composites containing 5wt% and 10wt%, it is possible that the involved POSS molecules in the agglomerates shown by the SEM micrographs (see Figure 3d and 3e), will not, or not completely react with hardener molecules. This would lead to free volume due to excess hardener in those composites with high POSS content, which would support the lowest values of $T_g$ for the ePOSS 10wt% composite.

### Table 1. Glass transition temperatures of neat epoxy and TGIB-POSS composites.

<table>
<thead>
<tr>
<th>Sample type</th>
<th>$T_g$ in °C</th>
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<tbody>
<tr>
<td>Neat epoxy</td>
<td>92.0</td>
</tr>
<tr>
<td>ePOSS 1wt%</td>
<td>88.6</td>
</tr>
<tr>
<td>ePOSS 2.5wt%</td>
<td>89.6</td>
</tr>
<tr>
<td>ePOSS 5wt%</td>
<td>88.6</td>
</tr>
<tr>
<td>ePOSS 10wt%</td>
<td>86.5</td>
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</table>

### 4.3 AC BREAKDOWN STRENGTH

The obtained AC breakdown (BD) data were treated by the two-parameter Weibull distribution in accordance with the IEEE 930 standard [21] in order to evaluate the breakdown strengths of the test specimens. In Figure 5, the Weibull plots are depicted, and the corresponding Weibull parameters $\alpha$ and $\beta$ are summarized in Table 2. The scale parameter $\alpha$ represents the breakdown strengths in kV/mm at which 62.3% of the samples experienced electric BD, while the shape parameter $\beta$ is an inverse measure for the scatter of the BD data, meaning the higher it is, the lower the variation of the data to the Weibull distribution.

Compared to neat epoxy, which had a BD strength of 214.9 kV/mm, all POSS composites, with the exception of ePOSS 10wt%, exhibited significant increase in BD strength. The highest BD value was found for ePOSS 2.5wt% (241.7 kV/mm), followed by ePOSS 5wt% (234.6 kV/mm) and ePOSS 1wt% (225.6 kV/mm), whereas the 10wt% composite (214.7 kV/mm) would perform just as neat epoxy. Thus, it seems there is an optimal TGIB-POSS content around 2.5 wt%, where this POSS additive is still homogeneously dispersed within the epoxy at a molecular level, without showing agglomerations. With a further increase of filler content, the presence of TGIB-POSS agglomerations, as observed by SEM (see Figure 3), will consecutively counteract the beneficial effect of POSS. Those agglomerations will cause local electric field enhancements, which subsequently lead to higher electric stress in the vicinity of agglomerates, and further results in a decrease of the BD strength for the composites, until the amount and size of agglomerations have completely nullified the POSS contribution as in the case of the 10 wt% composite.

Previous studies have shown increased BD strengths for epoxy composites with up to 2.5 wt% nano-silica, while incorporation of larger contents resulted in decreased BD values in comparison with the neat epoxy [22, 23]. This could most likely be attributed to conductive interfacial layers forming around silica nano-particles due to the hydrophilic nature of silica, creating conductive paths [24], and thus, corrupting the BD strength of such composites.

### Table 2. Weibull parameters as obtained from Figure 5.

<table>
<thead>
<tr>
<th>Sample type</th>
<th>$\alpha$ in kV/mm</th>
<th>$\beta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neat epoxy</td>
<td>214.9</td>
<td>14.02</td>
</tr>
<tr>
<td>ePOSS 1wt%</td>
<td>225.6</td>
<td>19.64</td>
</tr>
<tr>
<td>ePOSS 2.5wt%</td>
<td>241.7</td>
<td>19.55</td>
</tr>
<tr>
<td>ePOSS 5wt%</td>
<td>234.6</td>
<td>26.23</td>
</tr>
<tr>
<td>ePOSS 10wt%</td>
<td>214.7</td>
<td>11.72</td>
</tr>
</tbody>
</table>
The hydrophilicity of silica results from the formation of silanol groups at the silica particle surface [25]. By using reactive TGIB-POSS molecules in our study, where organic groups surround the silica core, the resulting composites are not subject to the phenomenon of conductive paths around the filler. Thus, significant improvements of the dielectric strength of the resulting TGIB-POSS composites, as compared to neat epoxy, could be also achieved for comparably higher filler contents. Even more so, when agglomerations of POSS molecules can be avoided at higher filler loadings, as presented in one of our recent studies [14].

4.4 RESISTANCE TO CORONA DISCHARGE

The results of the surface erosion experiment based on the respective eroded sample volumes after exposure to PD for 30 hours are presented in Figure 6. A decrease of eroded sample volume can be associated with an increase in resistance to electrical discharges. The experiment showed remarkable improvements in resistance to electrical discharge of all POSS composites, as compared to the neat polymer. With a 60% and 59% reduction in ablated material, for ePOSS 1wt% and ePOSS 2.5wt% respectively, the low content POSS composites revealed the best resistance to electrical discharges among all test specimens. With further increases of the POSS content, a continuous diminishment of resistance to PD was however noted for the resulting POSS composites with 5 and 10 wt% additive. This worsening of performance should be attributed to the occurrence of cumulative agglomerations with increasing POSS contents, as observed by SEM for ePOSS 5wt% and ePOSS 10wt% composites, and shown in Figure 3d and Figure 3e. Such agglomerations cause local electric field enhancements, which subsequently lead to locally increased electric and thermal stresses of the composite, resulting in intensified erosion of the surrounding organic matrix.

Similar results for improved corona resistance of epoxy composites featuring inorganic nanofillers such as silica have been reported in the past [26-28]. However, in some cases, very high nano-silica contents of 26 wt% and above [27] were needed to attain improvements comparable to those observed in this study, by incorporating 1 wt% of TGIB-POSS in epoxy. Thus, it is assumed that by dispersing POSS at a molecular level in epoxy, and because of the formation of covalent bonds between the POSS molecules with their inorganic cores and the epoxy matrix, the resulting change of morphology, and hence, the nanostructuration of the respective composite, gives rise to the significant resistance to corona discharges for the ePOSS 1wt% and ePOSS 2.5wt%. Coherent results with a different reactive POSS additive have been recently published in [12].

4.5 THERMAL CONDUCTIVITY

The thermal conductivities were measured twice on each sample type, while considering that the reproducibility of the used method used may vary by approximately 5%. Further, the respective average values of thermal conductivities from those two measurement series were taken for the following discussion. The experimental results are displayed in Figure 7.

It was found, that addition of TGIB-POSS in the epoxy matrix resulted in an increase in thermal conductivities of approximately 10% for the respective composites with up to 5 wt% POSS content. The highest thermal conductivities were exhibited by both ePOSS 1wt% (0.159 W/m-K) and ePOSS 2.5wt% (0.159 W/m-K), followed by ePOSS 5wt% (0.158 W/m-K). However, ePOSS 10wt% (0.149 W/m-K) did not reveal a significant change in thermal conductivity, as compared to the neat polymer (0.145 W/m-K).

It is well established that thermal conduction in solids is based on phonons (lattice vibrations) [29, 30]. Phonon scattering, which appears at either particle-particle or particle-matrix interfaces, or at the interfaces of the amorphous structure in epoxy resins itself [29], lead to attenuation of the thermal conductivity within a material. As a consequence, the addition of low contents (i.e. 1 wt%) of high thermally conductive filler was deemed insufficient for significant enhancements of the thermal conductivities of the resulting composites, as indicated in several studies [31-33].

However, it was established in [29] that creating ordered, crystalline-like structures on a nanometric level within epoxy resins, which are characterized by their amorphous structure, would cause a diminishment of phonon scattering, and hence, significantly improve the thermal conductivity of the polymer.
Accordingly, by adding POSS in the epoxy resin, which forms covalent bonds with the epoxy matrix, we succeeded in nanostructuring the resulting composites, which further resulted in reduced phonon scattering within the composite. In conclusion, the thermal conductivity of the POSS composites could be improved as compared to neat epoxy.

Nevertheless, there seems to be a threshold limit in the case of TGIB-POSS, due to agglomerations seen for the 5 wt% and especially the 10 wt% POSS composites. These agglomerations cause an augmentation of phonon scattering at matrix-particle interfaces, and hence, consecutively nullify the effect of nanostructuring. Consequently, this results in more or less equivalent thermal conductivities of ePOSS 1wt%, ePOSS 2.5wt% and ePOSS 5wt%. Moreover, in the case of ePOSS 10wt%, the numerous agglomerations might cause pronounced phonon scattering, which suppresses the beneficial effect seen for the composites with lower TGIB-POSS contents.

4.6 DIELECTRIC SPECTROSCOPY

The dielectric responses obtained at 20 °C for all the investigated materials are presented in Figure 8. Both the real part of the complex permittivity (Figure 8a) and its imaginary component (Figure 8b) show that all composites exhibit dielectric responses similar to that of neat epoxy for power frequencies and above. For lower frequencies though, additional interfacial loss peaks were found for ePOSS 5wt% and ePOSS 10wt% (Figure 8b), being more pronounced for the 10 wt% composite. The corresponding steps in real permittivity could also be seen for lower frequencies in Figure 8a. However, ePOSS 1wt% and ePOSS 2.5wt% did not reveal such interfacial loss peaks, and hence, their dielectric responses are similar to that of neat epoxy.

The appearing interfacial loss peaks found for the composites containing 5 or 10 wt% POSS can be linked to the aforementioned agglomerations observed using SEM microscopy, as presented in Figure 3d and Figure 3e. These agglomerations give rise to interfacial polarization for the composites with high filler content. This is further supported by the lack of an additional loss peak for the 1 and 2.5 wt% composites, as well as by the microstructure analysis where no agglomerations were found for these two composites (Figures 3b and 3c). It therefore seems analogous to the results already presented in the previous subsections, that for low TGIB-POSS contents a good dissolution in epoxy is possible, allowing a dispersion of POSS at a molecular level. Consequently, no additional interfacial polarization occurs [34].

Figures 9a to 9e show the temperature- and frequency-dependent dielectric response of all test specimens. The α- and β-relaxation peaks could be located for neat epoxy and the epoxy/POSS composites, for both the real and imaginary parts of the complex permittivity. In amorphous polymers, the α-relaxation is correlated to relaxation processes of the main polymer backbone due to chain movement as a result of the glass transition phenomenon. The β-peak however, is associated with secondary relaxation processes due to local crankshaft movements of hydroxyl ether groups in the case of epoxy resin [35].

For all investigated materials, the β-peak was located for temperatures below 80 °C, in the vicinity of 100 kHz. The α-peak was found at 100 °C, with a loss maximum around 100 Hz for neat epoxy, ePOSS 1wt%, ePOSS 2.5 wt% and ePOSS 5 wt% (see Figures 9a’ to 9d’). With the incorporation of 10 wt% POSS, the α-peak was shifted slightly towards higher frequencies though, as shown in Figure 9e’. This result is in good correlation with the glass transition values presented in Section 4.1. Given that the 10 wt% composite has the lowest Tg, the α-relaxation process is initiated at lower temperatures, as compared to neat epoxy, and hence, the peak is shifted to slightly higher frequencies at 100 °C.

Analogous to the observations made at 20 °C (see Figure 8), at temperatures of 80 °C and below, the BDS analysis revealed similar ε’ for neat epoxy, ePOSS 1wt% and ePOSS 2.5wt%, while ePOSS 5wt% and ePOSS 10wt% exhibited steps in ε’ in the low frequency domain (see Figures 9a to 9e). Those steps were much more distinct for the 10 wt% composite. According to these steps in ε’, additional interfacial loss peaks in ε” could be seen for ePOSS 5wt% and ePOSS 10wt%, with higher magnitudes for the 10 wt% composite (Figure 9d’ and Figure 9e’). This interfacial loss peak was more difficult to detect at a temperature of 80 °C in the case of ePOSS 5wt% and it was
Figure 9. Temperature-dependent real part (left column) and imaginary part (right column) of complex permittivity for a) neat epoxy, b) ePOSS 1wt%, c) ePOSS 2.5wt%, d) ePOSS 5wt% and e) ePOSS 10wt%
completely overshadowed by low-frequency dispersion due to charge fluctuations for a temperature of 100 °C for both, ePOSS 5wt% and ePOSS 10wt%. The increasing dielectric losses at 100 °C for lower frequencies with a slope ≠ -1 on the log-log scale, accompanied with the increasing real part of the permittivity for frequencies below 1 Hz, translates a more complex behavior than simple DC conductivity of being responsible for the increasing dielectric losses [23]. The low frequency dispersion at 100 °C can be explained by interfacial polarization on a macroscopic level at the electrode/sample interface, the so called electrode polarization [36], possibly superimposed by electronic conductivity. This is a rather common behavior for solid insulating materials at elevated temperatures [5].

4.7 RELAXATION BEHAVIOR

For further understanding of the relaxation behavior of polymer composites, their dielectric responses are frequently fitted by an empirical equation (3) proposed by Havriliak and Negami [36], as it is known to be valid for polymers as well as inorganic materials:

\[ \varepsilon^*(\omega) = \varepsilon_\infty + \frac{\Delta\varepsilon}{1 + (i\omega\tau)^{\alpha\text{HN}}} \beta_{\text{HN}} \]  

The frequency-dependent complex permittivity \( \varepsilon^*(\omega) \) can be approximated by (3), where \( \varepsilon_\infty \) is the high frequency permittivity, \( \Delta\varepsilon \) the dielectric relaxation strength, \( \omega \) the angular frequency (\( \omega = 2\pi f \)), \( \tau \) the relaxation time, \( \alpha_{\text{HN}} \) and \( \beta_{\text{HN}} \) (with \( 0 < \alpha_{\text{HN}} < 1 \) and \( \beta_{\text{HN}} > 0 \)) the Havriliak-Negami parameters which designate, respectively, the symmetric and asymmetric broadening of the relaxation time distribution function. When the shape parameters \( (\alpha_{\text{HN}} \text{ and } \beta_{\text{HN}}) \) are equal to 1, equation 3 is simplified, and the relaxation law proposed by Debye [37] is obtained.

In this study, the relaxation behavior of our test specimens was further analyzed by fitting the \( \beta \)-relaxation peaks of the imaginary part of the complex permittivity in order to obtain the corresponding relaxation times. After the corresponding relaxation times were obtained for all the investigated temperatures, the molecular mobility of polymer chains participating in the \( \beta \)-relaxation process was investigated, as described by the Arrhenius law:

\[ \tau = \tau_0\beta \exp \left( \frac{E_a}{k_B T} \right) \]  

According to the Arrhenius law, the relaxation time \( \tau_0\beta \), which is inversely proportional to the molecular mobility, can be described with the pre-exponential factor \( \tau_0\beta \), the activation energy \( E_a \) of the \( \beta \)-relaxation process and the Boltzmann’s constant \( k_B \). The global error, obtained after both the fitting of the \( \beta \)-relaxation and the derivation of the parameters of the Arrhenius law, was below 10%.

The relaxation plots and activation energies for the samples are depicted in Figure 10 and Table 3, respectively. It can be seen that neat epoxy and all POSS composites, with the exception of the 2.5 wt% composite, show very similar \( \beta \)-relaxation behaviors and activation energies. However, ePOSS 2.5wt% differs significantly, resulting in over 65% increased activation energy. This means, that by addition of 2.5 wt% POSS to the epoxy matrix, the molecular movements during the \( \beta \)-relaxation process are notably hindered. It appears that due to the good dispersion of POSS, along with the formation of covalent bonds between the filler and the epoxy, a more rigid epoxy/POSS network is obtained. Such an increase in the rigidity of the epoxy/POSS network is the result of incorporating the rigid silica-like cage structure of POSS into the matrix. However, if only 1 wt% POSS is added to the epoxy, this effect might not be that significant, and similarly, for the higher filler content composites with 5 wt% and 10 wt% POSS, where due to the formation of agglomerations not enough POSS could be integrated into the epoxy network, and hence, no distinct change in the rigidity of the molecular web resulted. These findings are supported by the aforementioned improvements in BD strength, erosion resistance and thermal conductivity, where the ePOSS 2.5wt% composite out-performed neat epoxy and the other POSS composites, significantly in some cases.

Table 3. Activation energies obtained by fitting the \( \beta \)-relaxation process.

<table>
<thead>
<tr>
<th>Sample type</th>
<th>( E_a ) in eV</th>
</tr>
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<tbody>
<tr>
<td>Neat epoxy</td>
<td>0.526</td>
</tr>
<tr>
<td>ePOSS 1wt%</td>
<td>0.526</td>
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<tr>
<td>ePOSS 2.5wt%</td>
<td>0.875</td>
</tr>
<tr>
<td>ePOSS 5wt%</td>
<td>0.523</td>
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<td>ePOSS 10wt%</td>
<td>0.504</td>
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</table>

5 CONCLUDING REMARKS

The aim of this study was to investigate the dielectric and thermal properties of epoxy/POSS composites, along with the effect of nanostructurization of composites, by incorporating a reactive TGIB-POSS and dispersing it at a molecular level. It has been shown that the addition of low amounts of TGIB-POSS (≤5 wt%) leads to significantly improved dielectric breakdown strengths, corona resistances, as well as thermal conductivities of the resulting epoxy/POSS composites. It is assumed that the formation of covalent bonds
between the filler and the epoxy matrix significantly improves the filler/matrix interaction, leading to intimate interfaces instead of undefined interphases, and hence, contributes to a distinct improvement in performance for the resulting composites. This is supported by the superior performances of the lower content composites ePOSS 1wt% and ePOSS 2.5wt%, where no agglomerations were found, and hence, the dispersion of POSS at a molecular level in the epoxy has been successfully achieved.

In conclusion, although we used a reactive additive, agglomerations could not be avoided for higher filler contents. This prohibited the incorporation of a larger POSS content in the epoxy network, which should be beneficial for further improvements in dielectric performance, such as the resistance to corona discharges. The limited dissolubility of the used TGB-POSS in epoxy can be attributed to its low degree of reactivity (only 3 epoxy side groups around the silica-like core). In fact, when a different POSS additive with a higher degree of reactivity, and hence, a higher amount of epoxy side-groups, was used to fabricate epoxy composites, we successfully incorporated even larger contents of POSS into the epoxy matrix, without finding agglomerations. Consequently, the corona resistance of these composites could be enhanced even more significantly [14].

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REFERENCES


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