Effective surface functionalization of fumed silica with NBR telechelic oligomer for high-voltage polyolefin-based dielectric composites

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Abstract

In this study an effective surface functionalization of fumed nano-silica for dielectric insulation composites is developed. A three step modification procedure was applied: 1. Silanization of the silica surface with (3-glycidoxypropyl)trimethoxysilane (GOPTMS); 2. Grafting of an acrylonitrile-butadiene telechelic oligomer terminated with two carboxylic acid groups (o-NBR); 3. Hydrogenation of the unsaturated polybutadiene segments for better compatibilization with the saturated polyolefin matrices and for higher dielectric performance.

The deposition of the coupling system was monitored by thermogravimetric analysis (TGA) revealing 6.4 wt. % of GOPTMS and 13.6 wt. % of o-NBR bond to the silica, resulting in an exceptionally high total deposition (20.0 wt. %). The o-NBR grafting was analyzed qualitatively by Fourier transform infrared spectroscopy (FTIR) showing successful deposition of o-NBR and effective hydrogenation.

The silica modified according to this procedure is a promising functional nano-filler for high-voltage insulation composites allowing improved dispersion and distribution in polyolefine matrices and therefore enhancing their electrical properties.

1. Introduction

Recent progress in diversification of electric energy generation sources and development of smart grids has triggered the need of designing innovative materials for high-voltage current transmission applications.

Thermoplastic polyolefins, such as polyethylene (PE), polypropylene (PP) or poly(ethylene-co-octene) (POE), exhibit good dielectric and insulating properties, which make them ideal candidates for high voltage applications. Moreover, they show very good processability, mechanical properties and aging resistance. However, these polymers are not entirely free from drawbacks: for example, the melting temperature of PE is relatively low, which necessitates cross-linking (XLPE) for high-voltage applications. Under service conditions, a local temperature rise could be dangerously close to the melting temperature of non-cross-linked PE. This results in inability of effective material recycling of the XLPE insulator after its use [1]. PP exhibits a significantly higher melting temperature than PE (~160 °C versus ~110 °C, respectively), however its elasticity is limited necessitating its copolymerization or blending with more flexible polyolefins (like POE) in order to achieve a satisfactory level of elasticity.

Incorporation of nano-particle fillers into polymer matrix in order to improve their electrical properties is reported in literature [2-4]. R. C. Smith et al, showed that incorporation of nano-fillers consisting of particles of a size lower than 100 nm results in a significant improvement of the dielectric strength and voltage endurance of the polymer composites [5]. This, results from increasing of the interfacial area formed between the polymer matrix and the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6], but also from altering the interphase properties by incorporating specific chemical groups attached to the nano-filler surface area [6].

In order to improve the dispersion and distribution of nano-fillers in the polyolefin matrix, a surface treatment of the filler is applied that reduces the physicochemical incompatibility between the polar nano-filler and the non-polar matrix. The nano-filler/polymer compatibilization effectiveness is very important, as poor dispersion and distribution of the filler can lead to counter-productive results like an early electrical breakdown of the dielectric composite [7]. Therefore, new approaches leading to effective nano-filler surface modifications are widely needed and developed.

In this study we present a novel approach for an effective surface functionalization of fumed silica for polyolefin-based nano-dielectric composite preparation. Such a surface functionalization of fumed silica is a challenging task, since its surface contains a significantly lower amount of silanol groups. To overcome this disadvantage, a three step procedure was applied: 1. Formation of oligomerized silane layer on the silica surface – this is the crucial step allowing high yield grafting as it introduces a high amount of ethoxy groups reactive towards the carboxy-terminated oligomer; 2. grafting of telechelic oligomer macromolecules onto the silane layer; 3 hydrogenation of the oligomer for better compatibilization with the saturated polyolefine matrix.

2. Materials & Methods

Fumed silica Aerosil 200 was obtained from Evonik Industries, Germany. (3-Glycidoxypropyl)trimethoxysilane (GOPTMS; 98 % purity), tripropylamine (98 % purity) and p-toluene sulfonyl hydrazide (98 % purity) were purchased from AWC BCR GmbH, Germany. Dicarboxy terminated acrylonitrile-butadiene oligomer (o-NBR; 8-12 wt. % of bonded acrylonitrile, 26th Nordic Insulation Symposium on Materials, Components and Diagnostics (NORD-IS 19), Tampere, Finland, 2019
Mₘ ~3800 g/mol) was originated from Sigma-Aldrich Merck, Germany. Technical grade toluene was obtained from BOOM B. V., The Netherlands.

In the first step, the silica was silanized with (3-glycidoxypropyl)trimethoxysilane (GOPTMS). 10 g of the silica with 1 ml of GOPTMS and 100 μl of an acidic catalyst were placed in three-neck round bottom glass flask equipped with thermometer and water cooler. At last, 300 ml of the toluene was added to make a stable suspension. The flask was immersed in a silicone oil filled heating bath, which was placed on a magnetic heater/stirrer. A teflon-covered magnetic stirring bar was placed inside the flask to stir the suspension vigorously (500 rpm). The ingredients were heated up to 110 °C in an air atmosphere and stirred for 14 hours. Afterwards, the silica was removed from the flask and filtered with the help of a low-pressure pump and rinsed two times with fresh toluene.

In the second step, the silica silanized in the first step was once again placed in a similarly equipped round bottom flask with 3 g of o-NBR. The mixture was treated under similar conditions as step 1 for 14 hours. Afterwards the modified silica was removed from the flask, filtered, rinsed with toluene and placed into a laboratory oven heated up to 120 °C for 6 hours. Then the modified silica was purified from ungrafted o-NBR by a Soxhlet extraction with boiling toluene for 14 hours.

In the third, final step, 1.5 g of the o-NBR grafted silica was hydrogenated with a mixture of 1.35 g of p-toluene sulfonyl hydrazide and 0.85 g of tripropylamine. All the ingredients were placed in the previously used round bottom flask equipped with thermometer, cooler and this time also a nitrogen inlet unit to implement an inert atmosphere for the reaction. The flask was filled with toluene and stirred with a magnetic bar for 8 hours in inert nitrogen atmosphere at 105 °C. Afterwards the modified silica was purified by Soxhlet extraction with toluene for 14 hours.

The amount of the deposited silane and grafted o-NBR were measured by means of a TA Instruments 550 thermogravimeter (TGA). The samples were heated from room temperature to 650 °C under synthetic air atmosphere with a heating rate of 20 °C/min. From this test the samples mass loss in function of temperature (TGA) and their derivatives (DTGA) were obtained. The o-NBR grafting results and the hydrogenation effect were qualitatively analyzed by Fourier transform infrared spectrometry (FTIR) using a Perkin Elmer Spectrum 100 machine equipped with an attenuated total reflectance (ATR) unit. The scanned wavelength spectrum was within 650 – 4000 cm⁻¹. The resulting spectrum obtained was an average of 12 measurements.

3. Results & Discussion

The approach of silica surface modification to improve the compatibility with polyolefine matrices bases on the methodology proposed by Z. Huang et al. [8]. This method comprises the following reactions: 1. Silanization of a surface with (3-glycidoxypropyl)trimethoxysilane (Figure 1); 2. Addition reaction between the epoxy ring from the silane and carboxylic acid groups positioned at both ends of a telechelic oligomer molecule (Figure 2). In order to improve the compatibility between the grafted oligomer and a polyolefin matrix and to improve the dielectric properties of the composites, hydrogenation of the oligomer backbone was performed (Figure 3).

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**Figure 1.** Scheme of silica surface silanization with GOPTMS.

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**Figure 2.** Scheme of o-NBR grafting onto the silica surface.
The first two steps of the modification were monitored by TGA showing the amount of the coupling agent chemically anchored to the silica surface (Figure 4). The way that fumed silica is produced results in a very low moisture content, which makes it promising for application in dielectric nanocomposites as a functional filler. The mass loss at 650 °C measured for the reference Aerosil 200 sample was only 0.6 %, whereas the modified silica samples exhibit 6.4 % and 20.0 % of mass loss after silanization and o-NBR grafting respectively. This shows a very high effectiveness of the modification, not only in comparison to commercially available modified fumed silicas (up to 4.5 % of deposited coupling agent), but even to more sophisticated approaches, for example living polymerization of polyethylene on a silica surface [9].

The silica grafted with o-NBR exhibits two step kinetics of thermal decomposition of the deposited organic layer. The high value of the temperature of the second highest decomposition rate (T2 = 557.4 °C) observed for the o-NBR-grafted silane suggests formation of a carbonaceous residue from o-NBR during the thermal decomposition. The acrylonitrile mers present in the o-NBR macromolecular backbone are considered
to be able to trigger carbonization of NBR-based composites [11, 12]. Under synthetic air atmosphere the carbonaceous char burns out completely leaving only silica as non-flammable residue (Figure 4).

In order to confirm the o-NBR presence on the silica surface, FTIR spectra were taken (Figure 6). The reference silica does not show any specific bands of organic compounds. But after grafting of the o-NBR via GOPTMS, four characteristic bands appear: bands at 2857 cm\(^{-1}\) and 2929 cm\(^{-1}\) originate from >CH\(_2\) symmetric and antisymmetric stretching respectively; the band at 911 cm\(^{-1}\) from =CH vinyl groups and the band at 969 cm\(^{-1}\) originates from =C-H bending from butadiene mers. The presence of the former bands at 2857 cm\(^{-1}\) and 2929 cm\(^{-1}\) are associated with saturated hydrocarbon chains, whereas the latter (911 cm\(^{-1}\), 969 cm\(^{-1}\)) come from unsaturated chain fragments.

### Table 1. Parameters of thermal stability of the surface modified silica samples: T\(_{\text{on}}\) – onset temperature of thermal decomposition (temperature of 1 % mass loss); T\(_1\) – temperature of the first highest decomposition rate; T\(_2\) – temperature of the second highest decomposition rate.

| Parameter      | Sample description | Silica | Silica + silane | Silica + silane + o-NBR |
|----------------|--------------------|--------|----------------|-------------------------|
| T\(_{\text{on}}\) (°C) | -                  | 243.7  | 274.7          |                         |
| T\(_1\) (°C)    | -                  | 254.4  | 349.4          |                         |
| T\(_2\) (°C)    | -                  | -      | 557.4          |                         |
| Residue at 650 °C (%) | 99.4              | 93.8   | 80.0           |                         |

Simultaneously, incorporation of nitrogen-containing polar acrylonitrile mers is expected to alter the charge trapping properties of the insulation composites by improving both, the dielectric breakdown strength and the space charge distribution as is suggested in literature [2].

### 4. Conclusions

A novel approach for an effective surface modification of fumed silica was presented. A straightforward three step modification procedure resulted in deposition of a significant amount (20.0 %) of the organic coupling system. The silica with the grafted and hydrogenated o-NBR oligomer exhibits two main functionalities: 1. High compatibility with polyolefine matrices like XLPE, PP and POE (88-92 wt.% of hydrogenated polybutadiene segments); 2. Altered electrical properties of the nano-dielectric composites by introduction of nitrogen-containing polar groups into the polymer/nano-filler interface (8-12 wt.% of acrylonitrile segments present in o-NBR).

Application of this nano-filler for polyolefine-based insulating composites is expected to result in a significant improvement of their dielectric performance.

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### References

[1] Y. Sekiguchi, N. Ohkawa, H. Nojo and S. Hashimoto, “Development of recycling technology of XLPE”, JICABLE’97, 2007.
[2] D. Ma, T. A. Hugener, R. W. Siegel, A. Christerson, E. Martensson, C. Ormneby and L. S. Schadler, “Influence of nanoparticle surface modification on the electrical behaviour of polyethylene nanocomposites”, IOP Nanotechnology, vol. 16, no. 6, 2005, pp. 724-731.
[3] X. Huang, P. Jiang and Y. Yin, “Nanoparticle surface modification induced space charge suppression in low density polyethylene”, Applied Physics Letters, vol. 95, no. 24, 2009, p. 242905.
[4] X. Huang, Y. Zheng, P. Jiang, and Y. Yin, “Influence of nanoparticle surface treatment on the electrical properties of cycloaliphatic epoxy nanocomposites”, IEEE Transactions on Dielectrics and Electrical Insulation, vol. 17, no. 2, 2010, pp. 635-643.
[5] R. C. Smith, C. Liang, M. Landry, J. K. Nelson, and L. S. Schadler, "The mechanisms leading to the useful electrical properties of polymer nanodielectrics", IEEE Transactions on Dielectrics and Electrical Insulation, vol. 15, no. 1, 2008, pp. 187-196.
[6] T. J. Lewis, "Nanometric dielectrics", IEEE Transactions on Dielectrics and Electrical Insulation, vol. 1, no. 5, 1994, pp. 812-825.
[7] T. P. Schuman, S. Siddabattuni, O. Cox, and F. Dogan, "Improved dielectric breakdown strength of covalently-bonded interface polymer–particle nanocomposites", Composite Interfaces, vol. 17, no. 8, 2010, pp. 719-731.
[8] Z. Huang, H. Ji, J.W. Mays and M.D. Dadmun, “Understanding the grafting of telechelic polymers on a solid substrate to form loops”, Macromolecules, vol. 43, no. 3, 2008, pp. 1099-1101.
[9] P. Xiang, K. Petrie, M. Kontopoulou, Z. Ye and R. Subramanian, “Tuning structural parameters of polyethylene brushes on silica nanoparticles in surface-initiated ethylene “living” polymerization and effects on silica dispersion in a polyolefin matrix”, Polymer Chemistry, vol. 4, no. 5., 2013, pp. 1381-1395.
[10] A. M. Tiefenthaler and M. W. Urban, “Thermal stability of silane coupling agents on Nextel fibres”, Composites, vol. 20, no. 2, 1989, pp. 145-150.

[11] G. Gao, Z. Zhang, Y. Zheng and Z. Jin, “Effect of fiber orientation angle on thermal degradation and ablative properties of short-fiber reinforced EPDM/NBR rubber composites”, Polymer Composites, vol. 31, no. 7, 2010, pp. 1223-1231.

[12] D. I. Chukov, A. A. Stepashkin, A. I. Salimon, S. D. Kaloshkin and I. S. Pyatov, “Mechanical properties and chemical resistance of new composites for oil pump impellers”, Applied Sciences, vol. 8, no. 5, 2018, pp. 750.