Studies on Thermal and Morphological Behavior of 3-
Aminopropyltriethoxysilane Grafted Epoxidized Ethylene-
Propylene-Diene Terpolymer

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Abstract: A novel and new copolymer of 3-aminopropyltriethoxysilane (APTES) on epoxidized ethylene-
propylene-diene terpolymer (eEPDM) via Sol-Gel Process. EPDM was epoxidized with an in situ formed per
formic acid to prepare epoxidized EPDM (eEPDM). The effect of eEPDM concentration, APTES concentration,
reaction time and reaction temperature on the graft copolymerization was studied. The grafting efficiency of
APTES on eEPDM, eEPDM was confirmed by Fourier Transform infrared (FT-IR) spectroscopy. The
Thermal behaviors of EPDM, eEPDM, g-EPDM were studied by Thermo gravimetric analysis and DSC. The
Morphological properties of Epoxidation of EPDM and eEPDM-g-APTES are compared.

Keywords: eEPDM, eEPDM-g-APTES, Grafting, process, sol-gel

I. INTRODUCTION

It has been reported that ethylene-propylene-diene terpolymer (EPDM) has good resistances to heat,
light, oxygen and ozone, because it has low contents of the nonconjugated diene component (1-4). Epoxidation
is the simple and efficient method for introducing a new reactive group into polyolefins, leading to new and
useful properties and wide use in a variety of applications (5). Epoxidation of olefinic compounds using organic
peracids has been extensively studied since oxiranes were prepared by reacting ethylenic compounds with
perbenzoic acid. Oxiranes are generally prepared using perbenzoic acid (9), chloroperbenzoic acid (10) and
monoperphthalic acid (11). Performic acid is an extremely active organic peracid (12) used for the
epoxidation, but, owing to its instability (17), it has to be prepared in situ.

The development of grafted hybrid polymeric materials yields new materials with the balance of
properties for high performance applications (18-22). The grafting of silane onto ethylene-propylenediene
terpolymer (EPDM) is a novel technique used for the commercial production of cross linked EPDM for cable
insulation (23). Tanida and Sato prepared heat shrinkable tubes by grafting VTMO with EPDM (24). Kawada et
al. synthesized silica filled silane grafted EPDM and EPDM composites (25) and found that the heat resistance
of silane grafted EPDM was considerably higher than that of ungrafted EPDM. Umada et al. investigated the
thermal stability of silane grafted EPDM and EPDM (26) and they concluded that the heat grafted EPDM exhibits
better thermal stability than those of ungrafted EPDM.

In the present study, a novel and new graft copolymer of 3-aminopropyltriethoxysilane onto
epoxidized ethylene-propylene-diene terpolymer (eEPDM-g-APTES) has been synthesized in toluene. The
effect of eEPDM concentration, APTES concentration, reaction time and reaction temperature on the graft
copolymerization was studied. The grafting mechanism of APTES onto eEPDM was confirmed by FT-IR
spectroscopy. The mechanical, morphological and thermal properties of eEPDM and eEPDM-g-APTES are
compared.

II. EXPERIMENTAL

2.1 Materials

The EPDM (ENB) elastomer used in this study was a commercial grade Nordel IP 4770P
(ethylene/propylene/ 5-ethylidene-2-norbornene = 70/25/5 by wt. %, Mooney viscosity, ML (1+4) at 125°C is 70
and density of 0.87 g/cm³) of DuPont Dow elastomers, USA. 3-aminopropyltriethoxysilane (APTES), (Mn =
221.3, boiling point = 217°C, density = 0.946 g/cm³,) was procured from Aldrich Chemicals, USA.
Dibutyltindilaurate (DBTDL) (density = 1.066 g/cm³, freezing point >110 °C, Mw = 631.56) was purchased
from Merck, Germany. Analytical grade formic acid (88%), hydrogen peroxide (30%), toluene, n-hexane were
used as received.

2.2. In situ epoxidation of ethylene-propylene-diene terpolymer

The EPDM was first dissolved in toluene in a three necked flask equipped with a mechanical stirrer and
thermometer, and maintained at 50°C in a water bath. Under continuous stirring, the EPDM solution was

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Studies on Thermal and Morphological Behavior of 3-aminopropyltriethoxysilane grafted epoxidized acidified stepwise with 88% formic acid to pH 2–3. The epoxidation was performed by dropping the required amount of H₂O₂ (30%) for 30 min. A rapid introduction of this reagent is not recommended, because it causes excessive development of oxygen due to the decomposition of hydrogen peroxide at high temperature. The reaction was continued for 7 h at 50°C. After epoxidation, the rubber was coagulated in acetone, thoroughly washed with distilled water, soaked in 1% w/v Na₂CO₃ solution for 24 h, and finally rinsed with distilled water. The rubber prepared was dried in a vacuum oven at 40°C to a constant weight.

2.3. Preparation of eEPDM-g-APTES

The grafting reactions were carried out in 500 ml three necked round bottom flask equipped with a reflux condenser, a Teflon-coated magnetic stirring bar and a nitrogen inlet. A given amount of eEPDM was dissolved in 200 ml toluene in the flask. APTES at a given mole ratio was dissolved in 50 ml of toluene and then added with dibutyltindilaurate (1%). The reaction was carried out under various experimental conditions (Table 1). After the completion of reaction, the products were precipitated with methanol, filtered and dried in vacuum.

2.4. Isolation of graft terpolymer

In the synthesis, the products obtained consists of ungrafted eEPDM, poly(aminopropyltriethoxysilane) and eEPDM-g-APTES. The ungrafted eEPDM was separated from rest of the products by extraction using n-hexane. The 3-aminopropyltriethoxysilane grafted eEPDM (eEPDM-g-APTES) was separated from poly(aminopropyltriethoxysilane) using DMF where the former is soluble and latter is insoluble.

The total conversion (%) was calculated from the ratio of the total weight of products formed to the weight of eEPDM charged. The grafting ratio and grafting efficiency were determined on the basis of changes in polymer weight during the reaction process and the total amount of eEPDM-g-APTES formed respectively. The grafting ratio and grafting efficiency were calculated from the following equations:

\[
\text{Total conversion} \% = \frac{\text{Total weight of polymers formed}}{\text{Weight of eEPDM charged}} \times 100
\]

\[
\text{Grafting ratio} \% = \frac{\text{Weight of eEPDM-g-APTES}}{\text{Weight of eEPDM charged}} \times 100
\]

\[
\text{Grafting efficiency} \% = \frac{\text{Weight of eEPDM-g-APTES}}{\text{Total weight of polymers formed}} \times 100
\]

2.5. Measurements

EPDM, eEPDM and eEPDM-g-APTES were characterized with the help of IR spectra obtained from Shimadzu-1800S, using solvent casted thin films. The tensile tests were carried out using a universal testing machine (ZWICK-1484) at 27 °C with a cross head speed was 500 mm/min, using dumbbell shaped tensile specimens according to ASTM D 412-87. The hardness of the samples was measured according to ASTM D 2240-86. The DSC was performed on a NETZSCH DSC 200 thermal analyzer under nitrogen flow in the temperature range -100 to 400 °C, with a heating rate of 10 °C/min. Thermo gravimetric analysis (TGA) were carried using a NETZSCA TG 209 instrument under nitrogen atmosphere at a heating rate of 10 °C/min. A Quanta-200, FEG, scanning electron microscope was employed to examine the morphology of the gold coated surfaces of cryogenically fractured samples.

III. RESULTS AND DISCUSSION

The tentative grafting reaction mechanism of APTES onto eEPDM is presented in Scheme 1. The grafting mechanism proceeds via aminolysis reaction where the eEPDM (ENB) polymeric chain covalently bonds with the silane after sol-gel processing.
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3.1 Fourier-transform infrared spectroscopy

Fig. 1 shows the FTIR spectra of EPDM, eEPDM and eEPDM-g-APTES. The IR spectra of EPDM (Fig. 1(a)) shows the C–H stretching vibration (aliphatic) at 2911 cm⁻¹, –CH₂ rocking vibration at 1451 cm⁻¹, CH₃ symmetric bending vibration at 1367 cm⁻¹ due to the presence of propylene group, –(CH₂)ₙ– wagging vibration at 721 cm⁻¹ due to the presence of polyethylene chain, C–C stretching vibration at 2851 cm⁻¹, and the unsaturation band (>C=CH–) at 811 cm⁻¹ due to the presence of ENB content. The FTIR spectra of eEPDM (Fig. 1(b)) was characterized by the presence of a epoxide band at 870 cm⁻¹ due to asymmetric epoxide ring stretching. Furthermore, the intensity of the >C=CH– band at 811 cm⁻¹ decreases because of the epoxidation of EPDM, which demonstrates that the C=C double bond in EPDM was converted to the epoxy functional group in eEPDM. The conversion of double bonds to epoxide was obtained as 50% (ca.2.4 mol %). To take advantage of relative change of absorbance at 811 and 870 cm⁻¹, a quantitative analysis was performed by area measurement of methyl deformation band at 1369 cm⁻¹ as internal standard (22). Fig. 1(c) illustrates the IR spectra of eEPDM-g-APTES which reveals: O-H stretching vibration at 3250 cm⁻¹, C-H stretching vibration at 2900 cm⁻¹ and C-H deformation at 1443 cm⁻¹. The disappearance of the absorption band at 870 cm⁻¹ clearly indicates that the reaction at the epoxide groups has proceeded. A new band has been formed at 1080 cm⁻¹ that is characteristic of Si-O bonds derived from APTES.

3.2 Effect of reaction conditions on graft copolymerization

3.2.1 Effect of eEPDM content

The effect of eEPDM content on the graft copolymerization is presented in Fig. 2. The grafting reaction was carried out at 80°C for 4 h with 2.5 wt.% of APTES. The grafting efficiency increase linearly with increase in concentration of eEPDM, due to the availability of epoxy ring present in eEPDM. The grafting efficiency for 10 wt.% eEPDM is 20% and that of 30 wt.% is 30%. With higher concentrations of eEPDM, more active centers are generated in the system and thus the grafting efficiency increases. On the other hand, at lower concentrations of eEPDM, the fewer the active centers, and thus higher degree of homopolymerization of APTES.
3.2.2 Effect of APTES concentration

Fig. 3 shows the effect of APTES concentration on the grafting efficiency. The grafting reactions were carried out at 80°C for 4 h with 1, 1.5, 2, 2.5 and 3 wt.% of APTES onto eEPDM. The grafting efficiency increases with increase in APTES concentration and attains the maximum and then declines down. The grafting efficiency for 2 wt.% of APTES is 23.83 whereas the highest grafting efficiency of 25.01 is observed for 2.5 wt% of APTES. But, when the grafting was carried out using 3 wt% APTES, the grafting efficiency decreases to 20%. The dramatic decline of grafting efficiency at higher concentration of APTES is due to the formation of homopolymer. These homopolymer successfully hinder the rate of penetration of APTES molecules to eEPDM macro radicals, resulting in decrease in grafting efficiency.

3.2.3 Effect of reaction time

The efficiency of grafting of APTES onto eEPDM with respect to reaction time, is given in Fig. 4. The reactions were carried out at 80°C with 0.1 wt.% of DBTL for 2, 4, 6, 8 and 10 h. It is observed that the grafting efficiency increases with increase in reaction time upto 4 h, and beyond 4 h there was no appreciable effect observed on the grafting efficiency. However, the total conversion increases with increase in reaction time due to the formation of the homopolymer.

3.2.4 Effect of reaction temperature

The influence of temperature on the grafting is presented in Fig. 5. The reaction was carried out with 2.5 wt.% of APTES and 0.1 wt.% of DBTL at different temperatures, namely, 40, 50, 60, 70 and 80°C. The grafting efficiency increases with increasing reaction temperature from 40 to 80°C due to increase in the rate of decomposition of epoxidation group. However beyond 80°C there was no appreciable change in the grafting efficiency observed, which indicates the completion of the reaction.
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![Graph of grafting efficiency against reaction temperature](image)

Fig. 5. Plot of grafting efficiency against reaction temperature (APTES, 2.5 wt%; reaction time, 4 h).

<table>
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<th>Condition</th>
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**Table 1** Conditions of graft copolymerization

3.3 Thermal properties

The effect of APTES grafting on thermal properties was determined by thermo gravimetric analyzer. Thermograms of EPDM, eEPDM and eEPDM-g-APTES are given in Fig. 6. The inception decomposition temperature of EPDM, eEPDM and eEPDM-g-APTES are 440, 444 and 449 °C respectively. Similarly the final decomposition temperatures of EPDM, eEPDM and eEPDM-g-APTES are 482, 486 and 490 °C respectively. The higher thermal stability of eEPDM compared to EPDM can be explained by the substitution of the double bond on the EPDM side chain with the epoxy rings (25). The grafting of APTES onto eEPDM also increases both inception and final decomposition temperatures because of the inherent high heat resistances of EPDM and APTES (26).

The glass transition behaviour of the hybrid material is associated with cooperative motion of large chain segment. The DSC results of EPDM, eEPDM and eEPDM-g-APTES are shown in Fig. 7. The midpoints of the transition temperature curve in the DSC curve are recorded as Tg values. The Tg values EPDM, eEPDM and eEPDM-g-APTES are -43, -40 and -36 °C. The glass transition temperatures Tg, increased due epoxidation. The increase in Tg as a result of epoxidation may be caused by the presence of the polar epoxide group, which gives more rigid backbone than the unsaturated group (27, 28). The results also indicate that the Tg value for eEPDM-g-APTES is higher than eEPDM and EPDM, due to the silane content. Similar behaviour was obtained for PMMA-MSMA hybrid system (29).

![TGA curves](image)

Fig. 6. TGA curves of (a) EPDM (b) eEPDM (c) eEPDM-g-APTES
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3.4 SEM Analysis

The morphological characterization of the films was carried out through SEM analysis. Fig. 8 and 9 shows the SEM photographs of EPDM and eEPDM which indicates a plane and smooth surface with some bigger particles. However, the surface of eEPDM-g-APTES is smoother with some orientations as seen in Fig. 10. The orientation appears on the surface of eEPDM-g-APTES can be probably due to chemical bonding between APTES and eEPDM chains.

IV. CONCLUSION

EEPDM-g-APTES graft copolymer was synthesized in toluene. The following conclusions are made based on the data resulted from different experimental studies. The grafting of APTES onto eEPDM proceeds via aminolysis reaction where the eEPDM polymeric chain covalently bonds with the silane which has been confirmed from FTIR spectra. It was ascertained that the threshold grafting efficiency was maximum at 2.5 wt. % of APTES. The grafting efficiency increases with increase of temperature and eEPDM content. The epoxidation of EPDM as well as grafting of APTES onto eEPDM improve thermal characteristics.
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REFERENCES