



Impact of Alumina Trihydrate and Silica on Mechanical, Thermal and Electrical Properties of Silicone Rubber Composites for High Voltage Insulations

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Submission: 📅 November 07, 2017; Published: 📅 December 01, 2017

Abstract

In recent years, silicone rubber (SiR) based composites have been widely investigated for outdoor applications due to its promising insulating properties. However, mechanical, thermal and tracking properties of pure silicone rubber are very low, which restrains its application for long-term performance. In this research work, the influence of micro-sized alumina-tri-hydrate (ATH) and micro/nano-sized silica (SiO₂) fillers on mechanical, thermal and electrical properties of room temperature vulcanized (RTV) SiR has been studied. SiR-blends with varying amounts of ATH and SiO₂ particles were prepared, by following blending in a two roll mixing mill, compression moulding and post curing processes sequentially. In order to evaluate relative tracking and erosion resistance of SiR-blends, inclined plane test (IPT) was conducted in accordance with ASTM-D-2303 standard procedure. Surface temperature distribution was recorded using Fluke-Ti25 infrared (IR) camera during IPT experiments. Thermogravimetric analysis (TGA) was carried out to analyze thermal stability of ATH/SiO₂ filled SiR Composites. Surface morphology, tensile strength, percent elongation at break (%EAB), hardness, erosion, tracking resistance and thermal properties were also investigated and discussed. Results showed that the mechanical, thermal and tracking/erosion performance of SiR-blends improves by the incorporation of ATH/silica particles, which is governed by filler type, size and wt% in polymer matrix.

Keywords: ATH; IPT; Characterization; SiR; SiO₂

Introduction

Since 1970s, silicone rubber has been used for outdoor insulation [1]. Due to light weight, unique hydrophobicity, better pollution/contamination performance, easy installation and excellent insulation properties (resistivity > 10¹² Ohms), SiR composite insulators are broadly employed in High Voltage (HV) transmission and distribution [2]. However, these polymeric materials are affected by various types of environmental and electrical stresses which limit its use as HV insulants [3,4]. Heavy rain, ultraviolet radiation (UV), pollution, dry band arcing, corona discharge, temperature, chemicals, cyclic loads and the moisture penetration plays a significant role in SiR degradation [5]. In addition, the tracking and erosion of polymer insulants due to the dry band arcing is the other main problem which is not yet fully addressed [6]. During service, if the localized surface temperature due to dry-band arcing becomes greater than the insulants safe limit, chemical reaction may takes place, which can lead to material tracking and erosion [7].

In order to enhance mechanical, thermal and tracking properties of pure SiR insulants, making these promising candidates for HV outdoor applications, inorganic particles incorporation in the base material attained focused attention of the researchers. Improvement in the physical and electrical properties, due to incorporation of micro and nano sized particles motivated researchers to try different types as well as sizes of fillers, so as to optimize its performance as insulants [8-11]. Furthermore, inorganic particles build the barrier to avoid surface tracking and hinder the insulator surface erosion as a result of heat produced during dry-band arcing, thus preventing the material degradation [12]. Previously, Fe₂O₃ [13], Al₂O₃ [14], carbon black, titanium dioxide, magnesium oxide, calcium carbonate and ZnO etc. were dispersed in SiR matrix and investigated [15-17]. However, these fillers are costly and susceptible to acid attack. ATH and natural silica have also been frequently incorporated into SiR matrix to tweak mechanical, thermal, tracking and erosion resistance

[7,18,19] This reinforcement is the function of filler size, shape and dispersion as well as molecular interactions [20].

Recently, synthesis, design, thermal degradation and erosion resistance of SiR composites was reported in number of studies [11,21-29]. Du et al. [21] reported that material degradation depends on leakage current magnitude and arc mobility. Likewise, Ansoorge et al. [22] found that at high temperature (350 °C for pure SiR), thermal degradation of SiR composites is due to depolymerization and formation of low molecular weight (LMW) components, which varies with pollution density and polymer functional groups. According to Meyer et al. [23] the Particle size, shape and particle-polymer bond potential are the utmost significant features of inorganic fillers to improve the polymer characteristics (electrical, mechanical and thermal). Furthermore, the authors of [24], investigated the impact of particle size, its surface modification and tracking resistances. In other work Du et al. [25] studied the influence of thermal conductivity on dc resistance to erosion of SiR/BN nano-composites. They inferred that with the increase in particle concentration from 0 to 7wt% thermal conductivity enhances, thereby reducing the weight loss and erosion. Likewise, the authors of [26], examined the effect of barium titanate (BaTiO₃) on surface degradation and tracking resistance of room temperature vulcanized (RTV)-SiR composites. Isaias et al. [11], investigated the inclined plane tests on ATH and silica filled composites with SiR and EPDM to evaluate the effect of inorganic particles on erosion/tracking resistance of SiR and EPDM composites. According to Refat et al. [27], the dry-band arcing and erosion of SiR can be suppressed by incorporating ATH and silica particles in the DC-IPT using wavelet based multi resolution analysis of leakage current. Likewise, it is studied that particles networked dispersion in polymer matrix can improve the composites properties [28].

This work is the continuation of our previous work in which we investigated the impact of silica particles on mechanical, thermal and electrical properties of EPDM, Epoxy and SiR composites [29]. However, AC-IPT based analysis of ATH and silica particles filled SiR Composites has not been thoroughly investigated before to the best of our knowledge. Investigated SiR composite insulants showed high tracking and erosion resistance as well as enhanced mechanical and thermal properties, indicating great potential for electronic and electrical engineering applications, such as high voltage transmission and substation insulation. Samples of

SiR mixed with micro-sized ATH, micro-sized SiO₂, micro-sized (ATH+SiO₂) and (micro-sized ATH+nano-sized SiO₂) particles with different wt% ATH/SiO₂ were prepared and investigated using IPT analysis in this study. The performance metrics studied includes: tensile strength, %elongation at break, hardness, weight loss with temperature and erosion/ tracking resistance properties of SiR novel blends. The synergetic impact of micro-ATH and micro/nano-sized-SiO₂ fillers and their wt% on the overall performance of SiR blends are discussed.

Experimental

Materials

In this work the base material, RTV-SiR (70% Poly-di-methylsiloxane(PDMS and 30% vinyl) with density 1.15g.cm⁻³ was supplied by Wacker Chemie Germany. And the micro-ATH, micro-SiO₂ and nano-SiO₂ fillers were produced by Huber Engineering materials, US-silica and Sigma-Aldrich, respectively. In addition, different physical properties of ATH and Silica particles are tabulated in Table 1. Moreover, various auxiliary chemicals like sulphur, ZnO, stearic-acid etc, were commercially available and purchased from local market. Furthermore, the chemical structures of ATH, SiO₂ and SiR are shown in Figure 1 respectively.

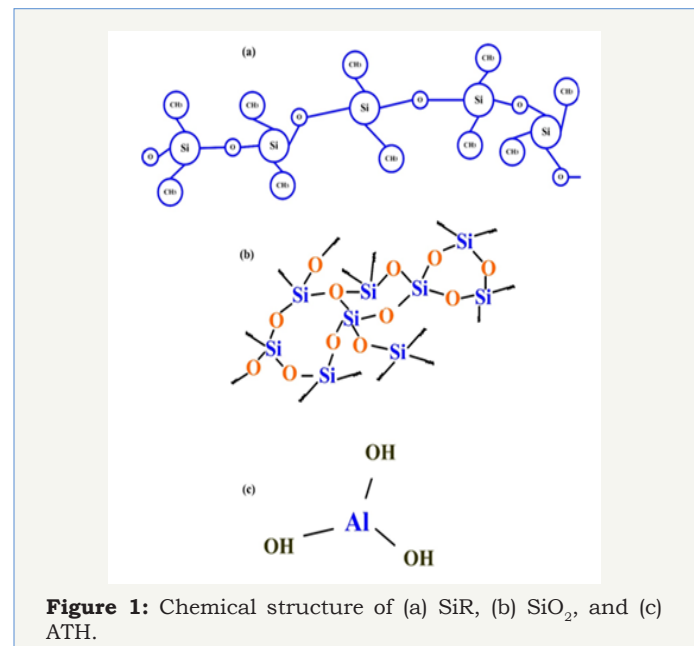


Table 1: Particles specifications.

Particle	Average Particle Size	Surface area/gram (m ² /g, BET)	Density (g/cu.cm)	Ph	Supplier
Micro-ATH	5μm	3.5	2.42	9	Huber Engineering Materials
Micro-SiO ₂	3μm	5	0.58	6.2	US Silica
Nano-SiO ₂	10nm	390	2.2	6.5	Sigma-Aldrich

Blends preparation

Preparation of RTV-SiR composites were performed in a high shear blender (HSM-100LSK) via simple blending followed by

hot-compression moulding and post curing technique. Blending of SiR and inorganic-particles (ATH/SiO₂) with known quantity of activator and accelerators was carried out in a two roll mixing

mill using ASTM-D 1418-10a standard procedure [30]. During mixing the mixer speed, mixing duration and temperature were kept as 50rpm, 20 minutes and 150 °C, respectively. Thereafter, the amalgamated mixture was poured into pre-heated steel mold (circular shape, 3.5mm thick and 85mm dia), following by compression molding in hydraulic press at 20MPa for 30 minutes.

Table 2: Prepared samples chemical composition.

Blend Code	Description	Base Material	Inorganic Particles		
			Type	size	Concentration[wt%]
A	Neat SiR	RTV-SiR	-	-	-
B	15% μ ATH+SiR	RTV-SiR	ATH	5 μ m	15
C	20% μ ATH+SiR	RTV-SiR	ATH	5 μ m	20
D	15% μ ATH+5% μ SiO ₂ +SiR	RTV-SiR	ATH	5 μ m	15
			SiO ₂	3 μ m	5
E	10% μ ATH+2% nSiO ₂ +SiR	RTV-SiR	ATH	5 μ m	10
			SiO ₂	10nm	2
F	15% μ SiO ₂ +SiR	RTV-SiR	SiO ₂	3 μ m	15

Blends characterization

Microstructural characterization: To develop polymer composites with optimized properties, the uniform dispersion of filler particles in polymer matrix is a critical dilemma. In this work a Hitachi-S4700 (USA), scanning electron microscope (SEM) was used to image the blends cross-sectional morphology. Moreover, before microstructural studies all blends were coated with gold to control the charging effects. All micrographs of square-shape samples (size $\sim 10 \times 10 \text{mm}^2$) were recorded at a magnification of $\sim 1000\times$.

Mechanical characterization: Mechanical properties such as Tensile Strength (TS) and %-Elongation at break (%EAB) were measured using universal testing machine (Instron-4465) with a 25mm/min cross head speed at room temperature according to ASTM D-412 [33]. Samples used were dumped shaped as per standard requirements. The Tensile strength and elongation at break can be calculated using the following equations [34].

$$TS \text{ (MPa)} = P / A \text{ (1)}$$

$$EAB \text{ (% strain)} = E / L \times 100 \text{ (2)}$$

Where, P, A, E and L represents the applied load, cross sectional area, strain at failure and gauge length, respectively.

Shore A Duo (Durotech) equipment was used to measure hardness as per ASTM-D2240 standard [35]. All hardness tests were performed on square shaped samples (50 \times 50 \times 3.5mm) and average of three blends of same composition is reported.

Thermal characterization: Percentage weight loss and thermal stability of prepared blends was performed using a thermal analyzer (Shimadzu-TGA-50H Japan) at 20 °C/min heating rate in temperature range 0 °C-850 °C. All tests were conducted

ZnO, Tetra-Methyle-Thiuram-Disulphide (TMDTD), stearic-acid and Mercapto-Benzo-Thiazole (MBT) were used as auxiliary additives. Moreover, further details about blends preparation can be seen from our previous studies [29,31,32]. Chemical composition of doped and un-doped SiR samples are summarized in Table 2.

on 20mg samples in air ambience by fixing the sample between two temperature controllable flat plates to provide uniform heat flow. The 10%, 50% and final residual weight temperature were measured under steady state condition for each sample.

Electrical characterization:

a. Inclined plane test (IPT) setup for tracking/erosion analysis

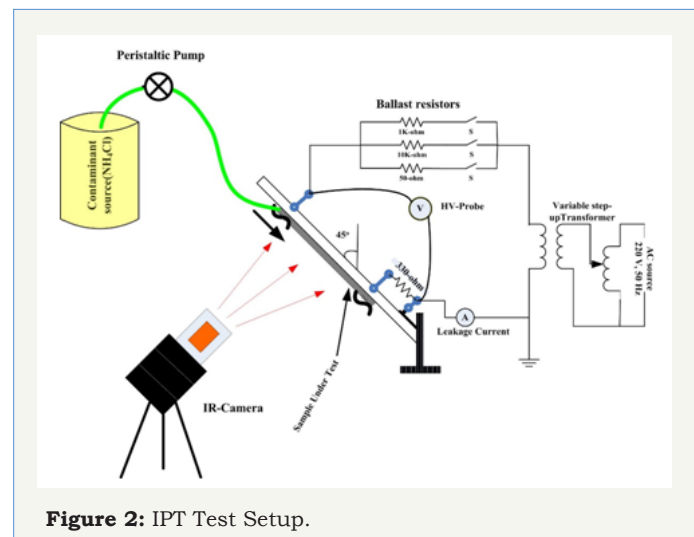


Figure 2: IPT Test Setup.

Various accelerated tests were employed to predict the performance and life of SiR-blends housing on HV transmission and distribution insulators. To evaluate the capacity of SiR composites to resist surface tracking and erosion, ASTM-D 2303 Inclined Plane Tracking (IPT) and erosion test is employed [36]. Figure 2, shows the test setup of IPT test with variable AC voltage source as per standard requirements. Each sample of size 65 \times 25 \times 3.5mm was energized from 220V, 50Hz AC voltage source with 70kVA/10kV variable transformer. In this test-setup, the composite samples

were mounted at an angle of 45 °C with respect to horizontal line along with contamination (NH₄CL with concentration of 1g/dm³ of distilled water) flow rate of 0.20ml/minute. The sample under test was observed after each passing hour, if the sample under test did not track, then the applied voltage was increased by 250V. In subsequent stages, the same procedure is followed until a continuous tracking of 25mm deep on insulator surface or leakage current of more than 60mA for 2s is observed [37]. Further, if the sample did not track in 6 hours, the test is stopped as required by the ASTM D2303 standard. In addition, measurement of the surface temperature distribution was recorded by Fluke-Ti25 infrared (IR) camera at equal interval of time, whereas leakage current (LC) was measured using Fluke-digital multi-meter between sample and ground terminal. As proposed by standard, before and after testing, all the specimens were cleaned with distilled water and alcohol (isopropyl) and weighted. The percent eroded mass was calculated using following equation [38].

$$\% \text{ eroded mass} = \left[\frac{\text{final mass} - \text{initial mass}}{\text{initial mass}} \right] \times 100 \quad (3)$$

Results and Discussion

Morphology results

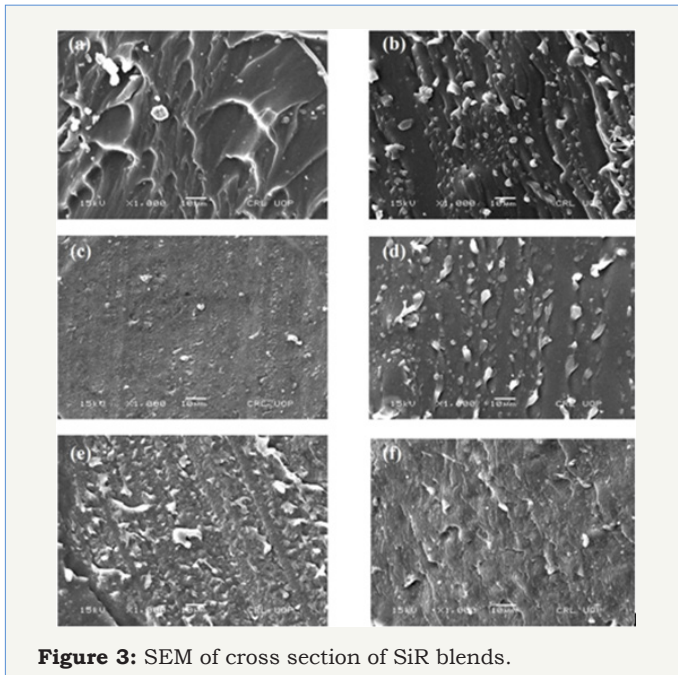


Figure 3: SEM of cross section of SiR blends.

The filler particles distribution, geometry, size and concentration in the polymer matrices have greatly influenced the mechanical, thermal and electrical properties of resultant composites [3,17,39,40]. Figure 3(e-f) presents, the SEM micrographs (on cross-section surface) of ATH/SiO₂ filled silicone rubber blends. From all micrographs no major dents and cracks were seen on surface of all blends. Moreover, among unfilled and filled SiR blends significant difference can be seen. Compared to SEM micrographs of blend-B and C, the neat SiR blend (Figure 3a) and filled SiR blends (Figure 3d-f) showed large agglomerates and in homogeneities. Furthermore, the 15% ATH blend (Figure 3b)

and 20%ATH blend (Figure 3c) with silicone rubber showed higher filler homogeneity and reduced filler agglomerations. Additionally, from SEM images it is obvious that each blend exhibited dissimilar micrographs with different phase difference among particles and base material. Consequently, each sample will exhibit different behavior in mechanical, thermal and electrical analysis. Hence, based on microstructural results, it can be inferred that the overall performance of micro-ATH filled SiR composited would be higher than that of the SiR hybrid (ATH+SiO₂ co-filled) composites.

Mechanical Results

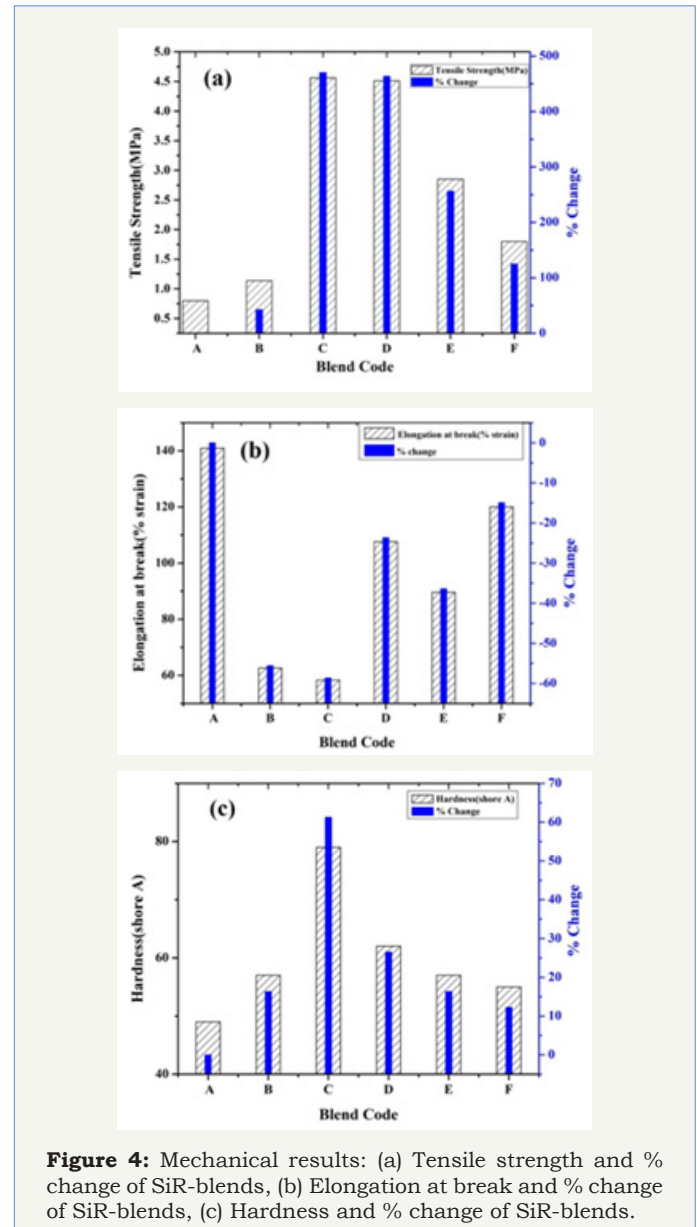


Figure 4: Mechanical results: (a) Tensile strength and % change of SiR-blends, (b) Elongation at break and % change of SiR-blends, (c) Hardness and % change of SiR-blends.

Mechanical properties like average tensile strength, % elongation at break and hardness of SiR-blends are compared in Figure 4(a-c). It can be seen that blending of ATH and SiO₂ particles with SiR can result in enhanced mechanical features. The improved tensile strength of ATH/SiO₂ and ATH-SiO₂-cofilled SiR-blends are presented in Figure 4a. This improvement is 42.5% for blend-B,

470% for blend-C, 463% for blend-D, 256.25% for blend-E and 125% for blend-F. Among all blends, only blend-C showed higher tensile strength of ~4.56 MPa as depicted in Figure 4a, which may be due to higher cross-linking (intense Hydrogen-bond between -OH groups of base rubber and filler surface groups) among SiR matrix and ATH particles. Another reason is may be the smaller free volume as well as particle uniform dispersion in polymer matrix which restricts chain mobility and consequently improves tensile property.

Likewise elongation at break (which act as insulants degradation indicator) of all prepared blends was decreased with the addition of both ATH and silica particles in different wt% as summarized and compared in Figure 4b. This decrease in EAB is 55.60% for blend-B, 58.65% for blend-C, 23.68% for blend-D, 36.4% for blend-E and 14.9% for blend-F. Maximum EAB was exhibited by blend-C as shown in Figure 4b. This decreasing trend in EAB of SiR blends is attributed to smaller inter-particle distances, strong bond energy and enhanced cross-linking, which restrain chain scission and molecular instabilities.

Similarly all SiR-blends showed improvement in hardness with the addition of ATH/SiO₂ particles in different wt%. This hardness improvement is 16.32% for blend-B, 61.22% for blend-C, 26.5% for blend-D, 16.32% for blend-E and 12.24 % for Blend-F as shown in Figure 4c. From Figure 4c, it is obvious that higher hardness was exhibited by blend-C (~79, shore A) compared to neat and all other SiR samples. This enhancement is attributed to particles small size, uniform dispersion, higher molecular interactions and improved cross-linking among particles and base matrix. Hence generally all SiR-blends showed improved mechanical properties, but blend-C (20%ATH) exhibited excellent mechanical properties.

TGA Studies

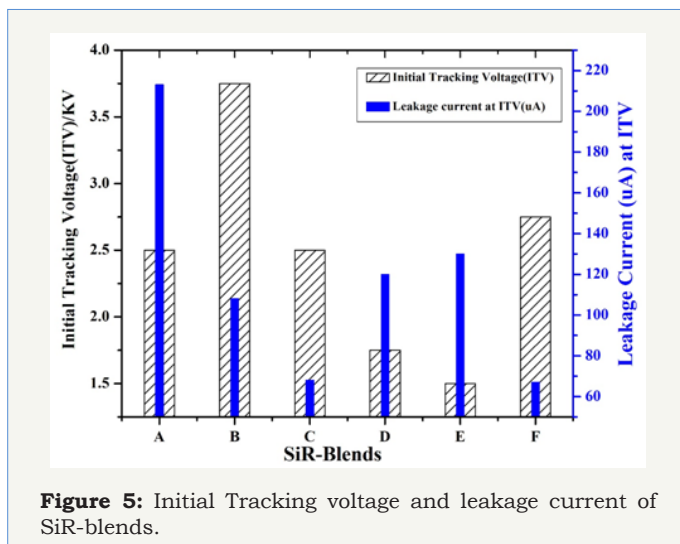


Figure 5: Initial Tracking voltage and leakage current of SiR-blends.

To understand the impact of ATH and SiO₂ particles on thermal performance of SiR-blends, TGA was conducted as shown in Figure 5. From TGA curves (Figure 6), it is obvious that some samples exhibit superior performance and some have degraded behavior than neat SiR for given temperature range (from 0 °C to 850 °C).

Thermal stability of SiR is increased for blend-B and blend-D, whereas blend E, C and F showed degraded performance than neat SiR. For instance, at 850 °C, the final residual wt left for blend A, B, C, D, E and F are 69.24%, 73.92%, 62.19%, 74.31%, 68.81% and 59.98%, respectively (Figure 6). Among all composites, blend-D exhibited highest residuary mass (74.31%), whereas lowest mass was found for blend-F (59.98%). This variation in mass loss and final residuary is due to the material decomposition, chain scission, evaporation of volatile species, moisture and gas particles. Blends thermal degradation may be due to CH₃-groups oxidation, de-polymerization, increased molecular interactions between SiR matrix and ATH/SiO₂ particles at elevated temperature and covalent bonds amalgamation (H-bond). Consequently, reduction in cross-linking as well as Vander wall forces can be exhibited by SiR blends at higher temperatures. Hence from TGA analysis, we infer that hybrid-particles (ATH+SiO₂) in certain proportion have significant impact on SiR thermal characteristics as validated by blend-D composition [39,40].

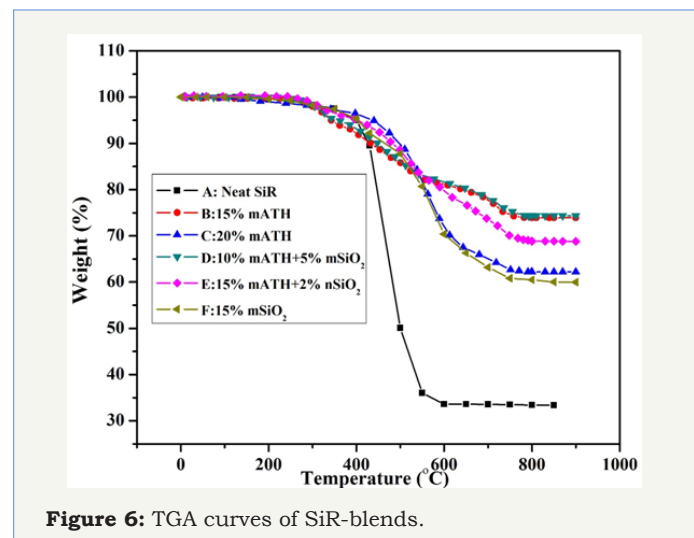


Figure 6: TGA curves of SiR-blends.

Electrical properties

Tracking/Erosion analysis results: The initial tracking voltage (ITV) and corresponding leakage current (LC) obtained for polymer samples are figured as function of SiR-blends as shown in Figure 5. It can be noted that ITV is independent of particles concentration (wt. %) and depends on energy dissipation in SiR-blends [41]. From Figure 6, it can be seen that ITV and LC varies in discrete proportion to ATH/SiO₂ filled SiR-blends. Its measured values are 2.5kV, 3.75kV, 2.56kV, 1.75kV, 1.5kV and 2.75kV for blends A, B, C, D, E and F, respectively. Among all the investigated samples maximum ITV is exhibited by blend-B compared to other SiR-blends. During test, execution chalking and hydrophobicity variation was also noticed in direct proportion with the applied voltage and test execution time on the top surface of SiR-blends. In addition, the Amonium-chloride (NH₄CL) contaminant during IPT test, bridges the top and bottom electrodes, which causes surface conductivity and joule-heating effect. As a consequence, the dry-band-arcing (DBA) is formed as soon as the input voltage exceeded than the sample breakdown voltage, as validated by

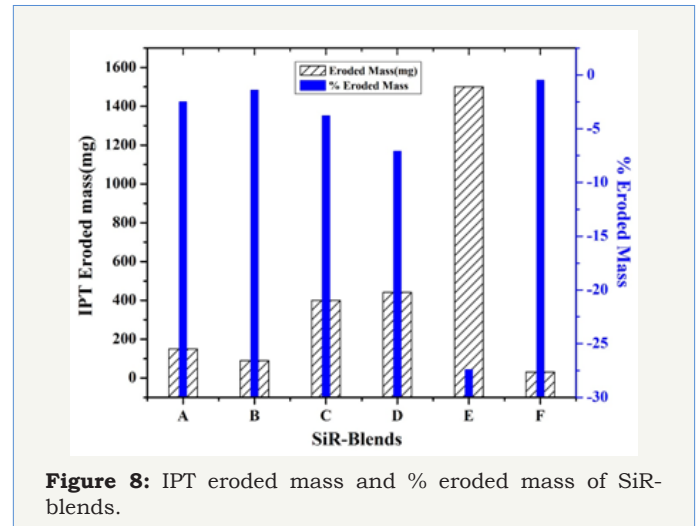
the blends-thermographs (Figure 7). During arcing period the generated heat is conducted from arc to specimen surface, which yields caloric/thermal deterioration and transforms samples nature from hydrophobic to hydrophilic as demonstrated in Figure 7. These thermographs of SiR-blends were recorded during IPT to investigate surface tracking, thermal profile and hydrophobicity variation with time. From IR images, it is obvious that DBA varies along sample surface with time, which eventually increases surface as well as surroundings temperature through diffusion of heat. Amongst all IR images of fabricated blends, it is inferred that upon gradual increase in voltage with time, minimum time-to-track was exhibited by blend-E (in 60 minutes), whereas maximum time-to-track was noticed for blend-B (in 300 minutes). From IR images, the red and bright spots denote higher temperature whereas, blueish spot indicates lower temperature.

Blend type	IR-image	Tracking Time(min)	Tracking Temperature
A		180	132 degree-F
B		300	103 degree-F
C		240	110 degree-F
D		120	162 degree-F
E		60	173 degree-F
F		240	121 degree-F

Figure 7: TGA curves of SiR-blends.

Moreover, in consequence of continuous arcing, enough energy can be transferred to the sample surface, which accelerates material decomposition as well as erosion as depicted in Figure 8. Herein, it can be seen that SiR-blends mass loss varies with filler type and concentration, which is 2.5% for blend A, 1.43% for blend B, 3.81% for blend C, 7.1% for blend D, 27.4% for blend E, and 0.5% for blend F. Among all investigated samples, maximum erosion takes place in blend E (~1500mg), whereas minimum mass loss was found in blend-F (~31mg) as shown in Figure 8. The SiR-blends with ATH (B, C, D, E) showed higher mass loss compared to SiR-SiO₂ blends. It is attributed to variation in bond energies among ATH/SiO₂ and SiR matrix. In addition, the heat generated during arcing and its conduction from tracked region to surrounding environment also influences the erosion of the polymeric blends, which varies for silica as well as ATH samples as validated by infrared (IR) thermographs. Furthermore, while performing erosion/tracking

test, gradual decrease in sample hydrophobicity with time was noticed for all blends, but rapid hydrophobicity loss was found for blend-E as indicated by IR images. This hydrophobicity loss may be due to evaporation and diffusion of low molecular weight (LMW) particles from bulk of material to outer surface.



Correlation between electrical and mechanical properties: Hydrophobicity, hardness, tensile strength, dielectric strength, erosion and tracking resistance are the material physical properties. These properties vary even among identical polymer samples due to internal defects, filler type/concentrations and molecular conformations [28]. In literature there is no direct mathematical relationship between mechanical and electrical properties. However, on the basis of literature it can be correlated as follows.

The dielectric breakdown (i.e. the maximum voltage that an insulation material can sustain) property of any insulating material depends upon the intensity of applied electric field (voltage), which can be calculated using the following formula [29,30].

$$\text{Dielectric strength (kV/mm)} = \frac{\text{Breakdown voltage (kV)}}{\text{Sample thickness (mm)}} \quad (4)$$

Likewise Swanson et al. [31] correlated the dielectric strength with volume/surface resistivity, relative permittivity and loss tangent according to the following formulas.

$$\text{Dielectric strength (kV/mm)} = A + [B * \text{Log} (\rho v / (\zeta r * \tan \phi))] \quad (5)$$

Where ‘ ρv ’ is the volume resistivity, ‘ ζr ’ is the relative permittivity and ‘ $\tan \phi$ ’ is the loss tangent.

$$\text{Volume resistivity } (\rho v) = [1 / (n * \mu * e)] \quad (6)$$

Where, ‘ n ’ is the number of charge carriers, ‘ μ ’ is the charge mobility and ‘ e ’ is the particle charge

It can be seen from Eq. (6), that there is an inverse relationship between volume resistivity and charge mobility. Now as soon as the applied electric field (applied voltage) across the polymer samples increases, the corona discharge generates ionic particles and energetic electrons, ultraviolet, ozone etc. Consequently these particles will cause the degradation of macromolecular

chains into low weight molecules (LWM) due to chemical bonds breaking [32,33]. Hence, with the increase in surface discharges due to applied voltage, the increased crosslinking process prevents the low weight molecules to transfer from bulk of the material to surface, which led to a reduction in hydrophobicity and mechanical stability [34]. This reduction in surface hydrophobicity due to electrical and non-electrical stresses could speed up the surface deterioration and physical defects (pores and flaws), thereby reducing surface/volume resistivities. As a consequent, it induces leakage current (LC) and dry band arc discharges on SiR Specimens [35]. This LC gives us good indications about mechanical and surface degradation process, because there is a direct relationship between LC and surface hydrophobicity (resistivity) as was observed during tracking/erosion investigations (section 3.3.1) of SiR blends.

In addition, the electrical (i.e. arcing, corona discharge, leakage current) and environmental stresses (i.e. Heat, UV-radiation, wind, biological degradation, snow, acid rain) have significant effect on insulation mechanical, thermal and electrical properties [36,37]. As the electrical stress occurs on insulator surface, it leads to the degradation of electrical characteristics, which is due to the loss of low molecular weight components and volatile particles. Likewise, the environmental stresses such as UV, Heat, snow, acid rain and biological factors de-polymerizes polymer insulants and scission of Si-O-Si chains. Consequently, loss of elasticity, erosion and surface roughness takes place which leads to reduction in mechanical stability. Hence it can be inferred that due to electrical and environmental stresses there is a direct relationship between electrical and mechanical properties as validated by IPT test and IR images. Because during IPT operation gradual decrease in hydrophobicity (resistance to water absorption and flow) followed by increase in LC was observed. As soon as sample lost their surface resistivity, tracking was occurred followed by erosion and weak mechanical performance [38-41].

Conclusion

In summary, the RTV-SiR composites filled with micro-ATH, micro- and nano-sized silica particles were prepared and investigated for surface morphology, mechanical, thermal, tracking and erosion properties. The amalgamation of micro/nano particles into SiR base matrix imparted enhanced mechanical, thermal and electrical properties. The major conclusions are summarised as follows.

a. Mechanically: Compared to all other SiR composites, the blend-C showed higher tensile strength (~4.56MPa), reduced elongation at break (~58.3% strain) and improved hardness (~79).

b. Thermally: The hybrid blend-D exhibited excellent thermal stability by leaving higher residual mass (74.3%) at 850 °C than that of other SiR blends. In addition, at 850 °C the neat SiR sample showed inferior thermal stability by leaving lower residual mass (33%).

c. Electrically: The enhancement in tracking and erosion resistance was found both for ATH and silica filled blends. However,

comparison to the ATH/SiO₂ filled SiR composites, the maximum tracking resistance was demonstrated by blend-B~3.75kV. Likewise, the maximum erosion resistance (lower eroded mass) was computed for blend-F (31mg), which is too lower than all SiR specimens.

Hence from silicone rubber novel blends, it is inferred that micro-ATH blends showed enhanced mechanical and tracking performance, while hybrid blend (D) and SiO₂-filled blend (F) exhibited higher thermal stability and erosion resistance, respectively. These characterization results will satisfy the applications of SiR insulants in the field of electronic and electrical engineering.

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