Properties of SBR/NR Blend: The Effects of Carbon Black/Silica (CB/SiO₂) Hybrid Filler and Silane Coupling Agent

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Abstract
In the current research, investigation of natural rubber (NR)/styrene butadiene rubber (SBR) blend reinforced with carbon black (CB)/silica (SiO₂) and with and without silane coupling agent (SCA) was analyzed. The total hybrid filler (CB/SiO₂) concentration in the composite was fixed at 50 phr. Cure characteristics, mechanical properties and surface morphology were examined. The results revealed that as the SiO₂ content increase, the optimum cure time, scorch time, minimum torque, and maximum torque of NR/SBR-CB/SiO₂ composites with and without SCA increased. Except for maximum torque, the optimal cure time and minimum torque of NR/SBR-CB/SiO₂ composites with SCA were lower than those without SCA. Compared with NR/SBR composites with SCA, addition of 0/50 CB/SiO₂ resulted in 53% decrease of tensile strength and 81% increase of elongation at break, superior to that of NR/SBR composites without SCA. The results revealed that as the SiO₂ content increase, the optimum cure time, scorch time, minimal torque, and maximum torque of NR/SBR-CB/SiO₂ composites with and without SCA increased. Except for maximum torque, the optimal cure time and minimum torque of NR/SBR-CB/SiO₂ composites with SCA were lower than those without SCA.

Keywords NR/SBR · Carbon black/silica · Silane coupling agent · Cure characteristics · Mechanical properties

1 Introduction
The development in industries and advancement in economic need results in never-ending demand for newer as well as low-cost materials. The polymer blending was a universal technology commonly used in the development of a newer product with superior properties. Because of its ability to adjust its shape crystallise under stretching, natural rubber (NR) has exceptional tensile properties [1, 2], as well as better stress relief, electrical insulation, high abrasion, and fatigue resistance. NR, on the other hand, was vulnerable to ozone and had a lower resistance to petroleum-based fuels and oils [3]. Because of its low cost, styrene butadiene rubber (SBR) was a commonly used rubber and was the first synthetic rubber considered for manufacturing rubber components. Despite their strong mechanical properties, SBRs have low abrasion resistance when exposed to conditions such as moisture, ozone, and sunlight. The presence of a double bond in the main polymeric chain was responsible for this [4–6]. Due to their excellent mechanical and other engineering properties, NR and SBR blends are very appealing nowadays. Natural rubber (NR) offers stronger strength, reduced heat build-up, and better low-temperature performance than styrene butadiene rubber (SBR). SBR has superior crack resistance, wet grip, and weather resistance than NR. As a result, tyre manufacturers frequently use NR/SBR mixtures. However, there are no defined principles for creating a compound for an NR/SBR blend in order to achieve a specific set of mechanical properties [7]. SBR and NR are partially miscible. In SBR–NR systems, the largest amount of interface was observed at a 50:50 blend composition [8]. Blends of natural rubber and butadiene rubber (NR/BR) or natural rubber and styrene-butadiene rubber (NR/SBR) are used in standard tyre formulations for cars and other vehicles. The blends of NR and SBR have been in use for a longer time period in many applications which also reduces the compounds cost [9, 10]. Studies have been made by many researchers of such blends and the different properties are examined in detail [11, 12].

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Despite NR and SBR are proven to have a wide range of desirable qualities, reinforcing fillers are often required to provide the desired properties for industrial applications. In the rubber industries, particle fillers are widely used for different functions, the most important of which being reinforcements, material cost reduction, and processing improvements [13]. More than 200 different types of organic and inorganic fillers have been examined and reported over the years [14, 15]. Unfortunately, not all of them are reinforcing material, but some have limited use. Different filler materials, such as carbon black (CB) and silica (SiO₂), are commonly used to enhance the mechanical properties of rubber composites [16, 17], but talc and calcium carbonates can only be used for reducing the cost of compound [18]. CB and SiO₂ have been commonly used as reinforcement materials in most engineering applications to improve the degree of rubber compounds. Tensile strength, tear strength, abrasion resistance, and modulus are all improved when CB is reinforced with either natural or synthetic rubber [19].

As compared to silica reinforcement, CB reinforced rubber composites have a higher modulus. The interaggregate interactions of CB, the nature of the polymer, hydrodynamic effects due to CB, and extra cross-linkage due to filler-rubber linkage all influence the elastic modulus of a rubber composite filled with CB. Size and surface area, surface activity and chemical properties, and form and structure are the key characteristics of carbon black filler. When it comes to interacting materials with rubber compounds, particle size and surface area are crucial. Carbon black particles range in size from 8 nm for furnace blacks to 300 nm for thermal blacks. Finer particles improve tensile strength, reinforcing, and abrasion resistance [20]. Whereas, silica offers properties such as improved tear strength, adhesion behavior, abrasion and aging resistance [21]. In comparison to CB reinforcement, silica reinforcement reduces rolling resistance, wet grip, and abrasion resistance in tyre treads [22]. In general, tyre tread compounds have a high filler percentage (above 40 phr), and silica has been included in numerous tread compounds in the previous two decades [23].

Since carbon black and silica both have their own set of benefits, using hybrid fillers (a mix of CB and silica) in rubber compounds can give you the best of both worlds. Wires and cloth coatings, tyre tread, rubber-covered belts, conveyor belts, hoses, engine mounts, cable jackets, and bumper strips can all be made with CB-reinforced rubber composites that have silica added to them [24, 25]. The reinforcing efficiency of these fillers is known to be dependent on the kind of rubber matrix, filler parameters such as size, structure, and surface functioning, and mixing conditions [26]. The surface chemistry of carbon black and silica is vastly different. On the carbon black surface, there are many functional groups such as hydroxyl, carboxyl, lactone, pyrone, ketone, quinone, and phenol, but the number is modest [27]. Silica is a filler for rubbers with particle sizes ranging from 10 to 40 nm. Chemically bound water makes up 25% of the silica, with adsorbed water accounting for the remaining 4–6%. In the polymeric industry, silica served as non-black reinforcing filler. In general, silica has a lot of polar groups on its surface, which helps the filler-filler interaction by hydrogen bonds to be stronger [28]. As compared to CB, adding silica to the rubber composite causes a slew of problems, most notably poor dispersion. Because the intermolecular hydrogen interactions between hydroxyl groups on silica’s surface are especially strong, it can aggregate tightly [29, 30]. Its property can result in a silica-filled rubber compound’s scorch time being prolonged and its delta torque being reduced due to poor dispersion of curatives by silica. The introduction of a particular coupling agent [31] will overcome the problems associated with silica incorporation. Silane coupling agents (SCAs) like bis(3-triethoxysilyl)-propyl)-tetrasulfide (TESPT) are commonly employed to increase filler dispersion and minimize curative adsorption on the silica surface [32–35]. A siloxane bond is formed when the silane coupling agent combines with silanol on the surface of silica. A crosslink between the silica and the polymer is formed when a silane molecule bound to the silica surface reacts with a rubber molecule.

CB–silica dual phase filler is a popular commercial hybrid filler (CSDPF). Cabot Corporation [36] produces it by pyrolyzing petroleum- and silicon-based feedstocks in a novel co-fuming process. CSDPF has a strong carbon domain surface activity and a high silica surface [37]. In compared to traditional CB and silica fillers, it has been demonstrated to provide improved overall mechanical qualities. A typical truck-tread compound has some features that are superior to the vulcanizate filled with CB N110, such as abrasion resistance and low rolling resistance. Because of the high polymer–filler interaction and low filler–filler interaction, this is the case. The CSDPF is reported to have strong abrasion resistance and tear strength advantages over silica [38]. Despite this, the CB and silica ratio in CSDPF has remained unknown. The types and contents of fillers have an impact on the curing characteristics of filled rubber compounds [39–42]. Silica-filled rubber compounds cure differently than carbon black-filled rubber compounds [35, 43–45]. The structure and surface chemistry of the reinforcing filler have an impact on crosslink density. A filled rubber compound cures faster than an unfilled rubber compound. Carbon black improves reversal resistance by increasing crosslink density [46].

The area between carbon black agglomerates decreases as carbon black loading increases, resulting in significant filler-filler interactions. This could be because gum vulcanizates have a lower cross-linking density than filled vulcanizates, which have a higher crosslinking density that increases as carbon black loading increases. This demonstrates carbon black’s potential to provide filled vulcanizates more rigidity, which lowers the rubber chains’ mobility [47].
On the other side, Bandyopadhyay et al. [48] found that adding silica to the rubber matrix reduces tensile characteristics due to less efficient interaction at the rubber/silica interfaces due to the aggregated silica particles’ smaller surface area. The silane coupling agent is prevented from interacting with the silica by the adsorption of acrylonitrile-butadiene rubber on the silica surface. As a result, there will be less interaction between the filler and the rubber [49].

To improve the mechanical properties of silica-filled rubber composites, silane coupling agents are used. It is required to neutralize silanol groups on silica’s surface while also increasing silica distribution in rubber blends. Many studies have been done on the impact of silane coupling agents on the mechanical and physical properties of rubber composites [50–55]. From the literature, it was clear that not much systematic research work have been reported with CB/SiO₂ hybrid filler on the properties of NR/SBR composites. Replacement of CB with SiO₂ in passenger car tire tread compounds, after the patent of Michelin in 1992 [56], offers tire compounds with lower rolling resistance and higher wet grip [57], so less fuel consumption and better driving safety. The present study employs high specific surface area and structure CB N330 in synergy with SiO₂ in NR/SBR tire tread composites. In the present work, the hybrid filler of SiO₂/CB-filled NR/SBR with various ratios is investigated keeping the total hybrid filler content constant at 50 phr. The effect of CB N330/SiO₂ on the properties of NR/SBR composites with and without SCA is investigated with respect to cure characteristics, tensile properties and morphology, all of importance for tire performance. CB/SiO₂ is not added beyond 50 phr due to the formation of cluster in rubber matrix and cost factor.

2 Experimental

2.1 Materials and Methods

The natural rubber (NR-RSS1) was purchased from Asian Rubber, Ambattur, Chennai, India and SBR(SBR-1502) was procured from Arihant Reclamation Pvt. Ltd., Delhi, India. The reinforcement (silica (SUPERSIL 140), carbon black (N330)), silane coupling agent (SCA), and curatives (zinc oxide, stearic acid, sulphur, mercaptozothiazylidisulphide (MBTS), tetramethyliuramdisulphide (TMTD) andprocessing oils) were of commercial grade and purchased from Vignesh Chemicals, Chennai, India.

The alteration of silica with TESPT was carried out in solution to see how SCA affects silica reactivity and to eliminate any confusing effects of the NR/SBR matrix. In a three-necked flask, 10 g of silica were dispersed in 300 mL of toluene, followed by 1.0 g of SCA. The silanization reaction was carried out at 80 °C for 12 h with constant stirring. After that, the result was centrifuged and washed with ethanol and toluene multiple times to remove the ungrafted TESPT before being dried at 50 °C.

The NR/SBR composite was formulated with a blend ratio of 50/50 as shown in Table 1. A two-roll mill at 50 °C was used to make the rubber compounds. NR was masticated for 4 min before being blended with SBR for another 4 min in the first stage. Before incorporating constituent or reinforcing materials, it was important to soften the compound blend. CB/SiO₂ was incorporated in the NR/SBR blend with two major portions. Half of the constituents (zinc oxide and stearic acid) will be induced together after half of the CB/SiO₂ is added. The other half of CB/SiO₂ and its constituents were treated in the

| Sl. No | Constituents (phr) | SBR | NR | CB | SiO₂ | SCA | Zinc oxide | Stearic acid | sulphur | MBTS | TMTD | Processing oil |
|-------|------------------|-----|----|----|------|-----|------------|-------------|---------|-------|------|----------------|
| 1     | Without SCA      | 50  | 50 | 50 | 0    | -   | 5          | 2           | 2       | 1     | 1    | 5              |
| 2     |                  | 50  | 50 | 40 | 10   | -   | 5          | 2           | 2       | 1     | 1    | 5              |
| 3     |                  | 50  | 50 | 30 | 20   | -   | 5          | 2           | 2       | 1     | 1    | 5              |
| 4     |                  | 50  | 50 | 20 | 30   | -   | 5          | 2           | 2       | 1     | 1    | 5              |
| 5     |                  | 50  | 50 | 10 | 40   | -   | 5          | 2           | 2       | 1     | 1    | 5              |
| 6     |                  | 50  | 50 | 0  | 50   | -   | 5          | 2           | 2       | 1     | 1    | 5              |
| 7     | With SCA         | 50  | 50 | 50 | 0    | 3   | 5          | 2           | 2       | 1     | 1    | 5              |
| 8     |                  | 50  | 50 | 40 | 10   | 3   | 5          | 2           | 2       | 1     | 1    | 5              |
| 9     |                  | 50  | 50 | 30 | 20   | 3   | 5          | 2           | 2       | 1     | 1    | 5              |
| 10    |                  | 50  | 50 | 20 | 30   | 3   | 5          | 2           | 2       | 1     | 1    | 5              |
| 11    |                  | 50  | 50 | 10 | 40   | 3   | 5          | 2           | 2       | 1     | 1    | 5              |
| 12    |                  | 50  | 50 | 0  | 50   | 3   | 5          | 2           | 2       | 1     | 1    | 5              |
same way. In the final stage, the sulphur was gradually added followed by MBTS and TMTD into the blend respectively. The total compounding time was done approximately from 25 to 35 min and compound dumping temperature were range from 50 to 80 °C. The prepared samples cure characteristics were investigated using a Monsanto Oscillating Disc Rheometer (ODR 2000) at 160 °C. The rubber compound was then compressed and moulded at 160 °C in an electrically heated hydraulic press in accordance with the cure times [58–60]. A die cutter was used to make the dumbbell and nicked tab end-samples.

2.2 Cure behaviors

The cure behaviour of the unvulcanized rubber samples was carried out as per ASTM D-2084 standard by oscillating disc rheometer at 160 °C, oscillation arc of 0.5° and 100 cycles per minute (1.66 Hz) conditions.

2.3 Mechanical properties

According to the requirements, dumbbell and nicked tab end shaped samples are cut from the moulded sheets. Tensile and tear checking was carried out in accordance with ASTM D 412 and ASTM D 624. Dak System Inc is used to perform tensile and tear tests at a crosshead speed of 500 mm/min (model: T-72102). The tensile strength, elongation at break and modulus were determined by Universal testing machine at 23 °C. The average values of six replicate measurements were used to arrive at the final outcome.

The samples hardness was determined using an ASTM D 2240 Shore-A Durometer. The hardness of composites was evaluated by the penetration of the Durometer tester into the rubber material, kept under formerly set conditions. The sheets with an effective thickness of 6 mm were used to test hardness.

The rebound resilience tests were carried out using a vertical rebound resilience tester in accordance with ASTM D 2632 guidelines. The rolling slide test was used to assess the abrasion resistance, which was determined as the relative volume loss from the wear test using a Zwick DIN Abrader (model: 6102), as per the ASTM D 5963 standard testing technique. A 0.1 kN force was given to the rolling sliding cylinder on SiC paper in this experiment. A minimum of five measurements were taken, with the average values reported.

2.4 Crosslink density

Cure test pieces of 20 X 20 X 2 mm were weighed and immersed in a glass vessel containing toluene (50 mL) at 23 °C. To avoid oxidation, the vessel was kept in a dark place. The samples were removed from the glass diffusion vessel and tissue paper was used to remove the excess toluene from the sample. To prevent toluene evaporation, the samples were stored in a closed vessel and the weights of the swollen samples were calculated. The sample was then re-immersed in toluene and the process was repeated until the sample reached a constant swollen weight. For the volume fraction of swelling rubber, the sample was de-dried in a vacuum at room temperature to a constant weight. The Flory-Rehner equation [61] was used to compute the molecular weight between two crosslinks (Mc) using the swelling data.

\[
M_c \left( \frac{g}{mol} \right) = \frac{-\rho_p V_r V_r^{1/3}}{\ln(1 - V_r) + V_r + \chi V_r^2} \quad (1)
\]

\[
V_r = \frac{1}{1 + Q_m} \quad (2)
\]

where, \( \rho \) = density of the rubbers, \( V_s \) = molar volume of the toluene, \( V_r \) = volume fraction of the swollen rubber, \( \chi \) =interaction parameter of the NR/SBR rubber, and \( Q_m \) = weight swell of the SBR/NR rubber blends in toluene.

The degree of crosslinking density (v) is given by:

\[
v (mol/cm^3) = \frac{1}{2M_c} \quad (3)
\]

The degree of crosslinking density of NR/SBR composites was determined using the following constant values.

\[
\rho \text{ (SBR)} = 0.921 \text{ g/cm}^3, \\
\rho \text{ (NR)} = 0.912 \text{ g/cm}^3, \\
\chi \text{ (NR/SBR)} = 0.3557 \quad [62], \text{ and} \\
V_s = 106.29 \text{ cm}^3/mol.
\]

2.5 Scanning electron microscopy (SEM)

With an acceleration voltage of 3 kV and a film of gold over it, a scanning electron microscope (Hitachi S-2500, Japan) was used to research the surface morphology of the tensile fractured surface.

3 Results and discussion

3.1 Cure behavior

Figures 1, 2, 3 and 4 depicts the cure behaviours (optimum cure time, scorch time, minimum torque, and maximum torque) of CB/SiO2 filler filled NR/SBR composites with and without silane coupling agent. The graph for the cure time is shown in Fig. 1. The optimal cure time for both composites increases as the silica content increases. This was due to the presence of the silanol group in it [63]. Silica’s surfaces are normally hydrophilic and polar. It also
has a high tendency for absorbing moisture, which has a negative impact on the rubber composites’ cure properties [64]. Figure 1 shows that when a silane coupling agent was added to an NR/SBR composite with a similar composite ratio, the optimum cure time \( t_{90} \) was shorter than when the silane coupling agent was not added. The composites with the presence of silanol groups which results in causing less hydrophilic characteristic, thus ensures a faster optimum cure time [65]. The silanol groups in sulfur-cured rubbers can cause accelerator adsorption, resulting in a delay in the vulcanization reaction and, as a result, a loss in crosslinked density [66].

The graph for scorch time was shown in Fig. 2. The scorch time of hybrid filler-filled NR/SBR composites with and without SCA and without silane coupling agent increases as the concentration of silica in the composite increases. When comparing composites with and without SCA, it was discovered that composites with Si69 have a longer scorch time \( t_2 \), indicating that there is good reaction between the SCA-silica-NR/SBR compounds in the composite. When the silica content is high, the amount of SCA utilised in this study is thought to be sufficient (as the SBR, NR, and SCA are constant except for silica content). As a result, the reaction between silica-SCA-NR/SBR was slightly increased. The NR/SBR tends to adsorbed on the silica surface in mixes without SCA, preventing the silica from adsorbing the accelerator and thereby reducing scorch time [67]. When silica concentration is high, especially greater than 30 phr, the scorch time...
increases steadily. The steady increase in scorch time when silica concentration is larger than 30 phr could be due to the fact that these compounds have had reduced heat history during mixing.

The incorporation of high specific surface area and high structure CB N330 that has better compatibility with NR/SBR compared to polar silica, causes an increased interfacial contact and consequently higher viscosity of the composites. The graph for the minimum torque was shown in Fig. 3. With an increase in silica concentration, the minimum torque of NR/SBR composites filled with hybrid filler with and without silane coupling agent increases. However, when comparing the composites with and without silane coupling agent, the composites with silane coupling agent exhibit a lower torque at all ratios, indicating that they are more processable [68–70].

The graph for the maximum torque was clearly shown in Fig. 4. From the figure it was clear that the maximum torque for NR/SBR hybrid composites with and without SCA was enhanced with an increase in silica concentration. Incorporation of hybrid fillers reduces the movement of the macromolecular chains in the composites, henceforth increases the maximum torque. However, when compared to NR/SBR composites without silane coupling agent, hybrid filler reinforced NR/SBR composites with silane coupling agent have a higher maximum torque value. The increased interaction between the hybrid filler and the NR/SBR matrix, especially in the presence of a silane coupling agent, may be responsible for the increased maximum torque [71–73].

### 3.2 Mechanical properties

From Fig. 5, 6, 7, 8, 9 and 10, the mechanical properties of CB/SiO₂ hybrid fillers reinforced NR/SBR composites with and without silane coupling agent are shown. Figure 5 shows the tensile strength of NR/SBR composites filled with hybrid fillers with and without a silane coupling agent. The figure clearly shows that both composites follow the same pattern, with the tensile strength of the composite decreasing as the silica content rises. The poor interaction of hybrid filler with NR/SBR rubber, as well as the rise in silica-silica interaction, resulted in a decrease in tensile strength at higher silica concentrations. Furthermore, as compared to the dispersion of carbon black, the dispersion of silica in rubber blends was very poor. This was due to the fact that silica forms a tight hydrogen bond, causing reinforcing material to clump together. The tensile strength of the hybrid composites was improved to a greater extent with the addition of a silane coupling agent. It was also apparent from the results that the composite with silane coupling agent had higher tensile strength than the composite without silane coupling agent at a comparable ratio of hybrid fillers (especially when the concentration of silica was more). The reinforcing material and the silane coupling agent formed a stable bond (Si–O–Si) when the silane coupling agent was incorporated into the composite [74]. The presence of a silane binding agent improved the interaction of hybrid fillers in the NR/SBR matrix, resulting in better dispersion. By lowering the interfacial energy between the stages, the hybrid filler’s interfacial adhesion to the NR/SBR matrix improves. The tensile strength of carbon black and NR/SBR matrix will be higher than silica due to good interaction and dispersion. In comparison to silica filler, carbon black filler has a stronger reinforcing effect [75].

The graphs for 100 percent modulus and elongation at break of hybrid composites with and without silane coupling agent are shown in Figs. 6 and 7. With increasing silica
concentration, the 100 percent modulus tends to decrease, while the elongation at break tends to increase. The matrix-filler interaction decreases as the concentration of silica increases. In contrast, the modulus of hybrid composites with silane coupling agent is higher than that of composites without silane coupling agent at equivalent composite ratios. This was due to the presence of a silane binding agent, which increased cross-link density and improved filler dispersion. The elongation at break values increase with increasing silica concentrations when a silane coupling agent is added. This is because the silane coupling agent aided chemical bonding between the hybrid filler and the NR/SBR matrix, increasing elongation at break values [76]. Elongation at break increment due to the interfacial adhesion force and movement of rubber molecular chains. Because of the poor dispersion of filler in NR/SBR composites filled with CB/silica, the interfacial adhesion force between rubber and particles is weakened, then lots of rubber molecular chains can move easily and get sufficient extension when composite is stretched, and that induces a high elongation at break.

Figures 8 and 9, clearly indicates the hardness and rebound resilience of hybrid fillers reinforced NR/SBR composites with and without silane coupling agent. The hardness of the composite tends to decreases whereas the rebound resilience increases with the increase in the concentration of silica in it. Lower crosslink density (as illustrated later in Fig. 11) of the composite was responsible for the minimum hardness. This was because, as the crosslink density of the
composite decreases, the harder matrix turns into a softer one \cite{77,78}. The increase in the concentration of silica leads in decrease in the rubber-filler interaction. Nevertheless, at a similar composite ratio, the hardness and rebound resilience of hybrid composite with silane coupling agent shows a greater value as compared to the composite without silane coupling agent \cite{79}.

The Fig. 10, show the graph for the abrasion loss of hybrid NR/SBR composites with and without SCA. The graph shows that as the concentration of silica increases, the abrasion resistance decreases. The abrasion resistance of the composites was measured by the friction coefficient and by the modulus of the composite \cite{80}. The lower frictional coefficient and higher modulus indicates that the composite have a better abrasion resistance. The differences among the composites that are filled with CB/SiO$_2$ and CB/silica/SCA could match the theory of modulus and dispersion of filler well according to what we have discussed in Fig. 5. As compared to composites with mediocre filler dispersion, the composites with excellent filler dispersion were said to have improved abrasion properties \cite{81–84}. As a result, the histrionic reduction in abrasion resistance at high silica content was expected to be due to weak silica dispersion or a heavy filler–filler relationship \cite{85}. As to the CB/SiO$_2$ composites with this experimental conditions, the friction coefficient might have a great influence on the wear resistance of composites \cite{86}.

The Fig. 11, show the graph for the crosslinking density of hybrid NR/SBR composites with and without SCA. Increases in silica concentration result in a decrease in filler-rubber interaction (increases in silica-silica interaction), affecting the crosslinked density value \cite{87–90}. The tensile strength, tensile modulus, elongation at break, and hardness of hybrid fillers filled NR/SBR blends with SCA are higher than those of NR/SBR blends without SCA at similar blend ratios. In the presence of SCA, this is simply owing to enhanced filler dispersion and greater crosslinked density. The SCA significantly reduced filler-filler interaction, resulting in enhanced rubber-filler interaction and improved filler dispersion.

### 3.3 Scanning electron microscopy

The Fig. 12a-b shows the SEM micrographs of tensile fractured specimen reinforced with hybrid fillers with and without silane coupling agent at 20/30 composite ratio. Figure 12a shows a rougher surface with numerous tear lines, indicating that more energy was needed to break the composite than Fig. 12b, which was nearly smooth. This is because the presence of a silane coupling agent improves the interaction of the CB and silica with the NR and SBR rubber matrix. In SEM micrographs of SBR/NR composites, the black region represents the rubber matrix and CB/SiO$_2$ hybrid filler, while the white line represents the silane coupling agent. The cross-section of silane coupling agent is represented by the white lines or areas. The uniformly distributed hybrid fillers in NR/SBR matrix have an ability to resist the crack initiated and propagates which results in better tensile property of the composite \cite{91}. The addition of the silane binding agent to the composite improves filler dispersion and decreases the silica-silica interface, resulting...

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**Fig. 11** The crosslinking density of CB/SiO$_2$ filler filled NR/SBR composites with and without SCA

**Fig. 12** The tensile fracture surfaces of hybrid fillers filled NR/SBR composites with ×300 magnification, (a) with silane coupling agent at 20/30 composite ratio, (b) without silane coupling agent at 20/30 blend ratio
in improved matrix-reinforcing material interaction. Further, due to the decrease of hydrogen bonds between the silanol groups on the silica surface, the composites without SCA are non-uniformly dispersed in SBR/NR matrix with agglomerates [92]. In contrast, in composites containing SCA, practically all of the silanol groups on the surface of the silica particles were reacted, causing the silica filler to be compactable with the SBR/NR rubber matrix and well-dispersed in SBR/NR composites, as illustrated in Fig. 12a.

4 Conclusions

The optimum cure time of hybrid fillers reinforced NR/SBR composites with silane coupling agent was less than the composite without silane coupling agent, according to the report. This was attributed to the decrease in curative adsorption phenomena observed on the silica surface. The scorch time of the hybrid filler reinforced NR/SBR composites with silane coupling agent is higher. With increasing silica concentration, the minimum and maximum torque for the composite with and without silane coupling agent increases. Hybrid filler reinforced NR/SBR composites with silane coupling agent had higher tensile strength, modulus, elongation at break, hardness, rebound durability, and abrasion resistance than composites without silane coupling agent. This was due to the composite’s improved matrix-filler interaction due to the presence of the silane coupling agent. The presence of silane coupling agent in hybrid fillers reinforced NR/SBR composites is revealed by SEM micrographs, which reveal better dispersion of the hybrid fillers in the NR/SBR matrix.

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Author contributions 1. G. Anand – Corresponding Author – First Author – Fabricated the composite samples and conducted the testing as per the ASTM standard and have written the manuscript, both authors read and approved the manuscript.
2. S. Vishvanathperumal – Second Author – Helped in Fabrication the samples and conducted the testing as per the ASTM standard and helped in writing the manuscript, both authors read and approved the manuscript.

Data Availability All data generated or analyzed during this investigation are included in this published article.

Competing Interests None of the authors have any relevant financial or non-financial competing interests and any conflict of interest to publish the article.

Conflict of Interest The authors declare no conflict of interest.

Compliance with ethical standards No, all testing are conducted in this study as per ASTM standards.

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