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A step towards a green and sustainable method to understand the effect of glucose on a silica filled natural rubber composite

In the present scenario, the world is concerned about producing more environmentally friendly and sustainable products by reducing the carbon footprint, especially in elastomer products. Therefore, the incorporation of maximum naturally produced white fillers like silica into a naturally originated elastomer, *i.e.*, natural rubber (NR), is the ultimate aim of most of the rubber industries. However, adding a bio-based carbohydrate (glucose) into a silica-filled NR matrix facilitates the dispersion of the silica filler into NR, making the product greener and more sustainable. The interaction between the functional groups (amine) of NR and carbonyl groups of glucose is known as the Maillard reaction, which interestingly exhibited a significant improvement in both mechanical and dynamic properties of the NR vulcanizates due to the superior dispersion of silica in the NR matrix, as confirmed by detailed FIB-SEM and AFM analyses. The contribution of the Maillard reaction to crosslinking and network formation between the polymer–polymer and polymer–filler is validated by an in-depth rubber state ¹³C NMR study.

1. Introduction

Natural rubber (NR) is purely a sustainable material since it is a green elastomer derived from mother nature (rubber tree: Hevea brasiliensis). This naturally occurring elastomer has superior dynamic and mechanical properties compared to synthetic elastomers, including green strength, tear strength, tensile strength, excellent growth resistance, and minimal heat buildup.^{1,2} The main factor contributing to the superior mechanical capabilities of NR is the development of strain-induced crystallization in the elastomer during the tensile test.3,4 According to the detailed analysis of the NR elastomer, the structure of rubber hydrocarbon starts with the ω -terminal coupled to two units of trans 1,4isoprene, followed by a lengthy series pattern of cis 1,4-isoprene units (>1000 units), and finally the α -terminal. The long chain of fatty acid ester groups forms a chemical link between the phospholipid and α -terminal. On the other hand, the physical links make a bridge to join the protein and ω-terminal.⁵⁻⁸ In the case of the NR matrix, integration and dispersion of polar fillers like silica are exceedingly difficult due to the presence of the functional

Many researchers have proposed various ideas and technologies over the past few decades to solve this critical problem and produce green tyres. One method involves forming a chemical attachment between silica and NR using alkoxyl, poly-sulphide, or alkoxyl-poly-sulphide bonds. NX using alkoxyl, poly-sulphide, or alkoxyl-poly-sulphide bonds. NX using alkoxyl, poly-sulphide, are TESPT bis(triethoxysilylpropyl) tetrasulphide, APTES (3-amino-propyltriethoxysilane), NXT (3-octanoylthio-1-propyltriethoxysilane), 3-mercaptopropyl-ethyoxyl-di(tridecylpentamethoxy) (Si747), superlink bis(tridecyl-pentaethoxy-siloxane), OCTEO (triethoxyoctyl-silane) and TESPT in combination, etc.], particularly silane coupling agents, are used for this purpose. 15-21 However, the cost of silane coupling agents is a concern for the rubber industry, and they often cause pre-scorching of the rubber compound due to sulfur release at processing temperatures. It is reported that

groups of proteins and phospholipids. The surface of amorphous silica particles is covered with silanol groups and siloxane bridges, causing them to aggregate through hydrogen bonding. Due to poor compatibility, silica does not mix with NR as easily as carbon black. To reduce flocculation and improve dispersion and filler-polymer interaction, coupling agents are used. These agents create bonds between silica and rubber, acting as interconnecting bridges. P11 A more environmentally friendly and long-lasting rubber product is one wherein the silica filler has been used to replace as much carbon black (CB) as possible. Manufacturing green tyres, in particular, benefits greatly from the combination of improved dispersion and greater silica loading in terms of essential rubber product properties.

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HTNR (hydroxyl-terminated natural rubber), ENR (epoxidized NR), guayule rubber, and deproteinized NR show better silica dispersion, but they cannot replace pure natural rubber compounds when considering key physicomechanical properties. 9,22-27 However, during the silanization process with silica particles, the functional groups of non-rubber elements compete with the coupling agent, compromising the silica filler's homogeneous dispersion in the NR matrix. 9,28 Instead of getting rid of the proteins and phospholipids, the notion of introducing glucose as a blocking agent at the beginning of mixing was presented. Given that one of the internal sugars in tapped natural rubber latex is glucose, this monosaccharide sugar, the most common carbohydrate, plays a part in the biosynthesis of natural rubber (NR) latex from rubber trees and other NR latex-producing plants. ^{29,30} In the past, Nimpaiboon et al. investigated how adding different sugars to fresh natural rubber, deproteinized natural rubber, and synthetic isoprene rubber affected their viscosity.² The incorporation of sorbitol and sorbic acid was examined to facilitate silica dispersion in silica-filled natural rubber composites.^{7,13} However, due to the presence of an aldehyde group in glucose, it was not suitable for use in natural rubber. The aldehyde group of glucose can interact with the amine group present in natural rubber, causing the Maillard reaction. This reaction may result in the hardening of the rubber compound and deterioration of the properties of the final product. Nonetheless, a different hypothesis was proposed to further explore this assumption: using glucose as a blocking agent of the functional groups of natural rubber. This approach could also contribute to the uniform dispersion of silica. The Maillard reaction might also improve the composite's strength and modulus in a homogeneously silica-dispersed NR system. It is possible that the functional groups of glucose will partially block the functional groups of NR in the NR molecules at an early mixing stage. The functional groups of glucose will then act as dispersing and plasticizing agents during the inclusion of silica to create a silica NR composite.

In the current work, we will compare the mechanical and dynamic characteristics of silica-filled NR composites with a detailed morphological investigation of the elastomer vulcanizates to examine the impact of glucose on those materials. To understand the variations in crosslinking junctions at different loadings of glucose and the silica filler in the NR elastomer compound, a comprehensive rubber state ¹³C NMR investigation was carried out. Moreover, a detailed morphological analysis was executed to perceive the dispersion of silica in the NR matrix.

2. Materials and methods

2.1. Materials

The used compounding materials are as follows: natural rubber (NR) - Technically Specified Rubber (TSR-10) purchased from EQ Rubber. Highly dispersible silica (HD silica) [grade-160007/N₂SA-182.54 (m² g⁻¹)] was supplied by Madhu Silica. Stearic acid, zinc oxide (ZnO), the curative - soluble sulfur, and the curing accelerator - N-tertiarybutyl-2-benzothiazole sulfenamide (TBBS) were purchased from local suppliers (analytical grade). Glucose, a 6carbon carbohydrate used as the modifying or blocking agent, was procured from Sutaria Chemicals, Mumbai.

2.2. Compound preparation

2.2.1. Rubber compounding formulation

2.2.2. Mixing conditions. The fill factor of the internal mixer with a tangential rotor type was around 0.65. After the third mixing stage in a Banbury mixer, the formulated (Table 1) compound was rolled through a two-roll mill to produce an 8 mm thick sheet, which was then let to rest at room temperature for a full day. The composites' optimum cure time was estimated by using the Moving Die Rheometer (MDR 2000 Alpha Technologies, USA) at 160 °C for 30 minutes. In compliance with the optimum cure time (T_{c90}) , the molding was run at 160 °C using a compression moulding machine (Moore Press, United Kingdom) and cooled to room temperature.

2.3. Testing and characterization of samples

The NR composite's rheological properties were tested at 160 °C for 30 minutes using the MDR (Alpha Technologies) to understand the cure characteristics. The hardness of all rubber vulcanizates was measured using a durometer A and an IRHD combined model hardness tester (Gibitre Instruments, Italy) according to the ASTM D 2240 test method. The results were reported as an average of four observations. Tear (angular) and tensile (dumbbell-shaped) specimens were punched out from the molded sheets using a hollow die punch (CEAST, Italy). The tests were performed in a universal testing machine (Zwick Roell Z010, Germany) at room temperature (25 \pm 2 $^{\circ}$ C), at 500 mm min⁻¹ crosshead speed. Tear and tensile tests were carried out in line with the ASTM D 624-99 and ASTM D 412-98 methods, respectively. The crosslink density of rubber vulcanizates was measured using the Flory-Rehner method using eqn (1) and (2). The samples were cut into round shapes (12.5 mm diameter discs) and weighed before and after being soaked in toluene for 72 hours, which ensured a swelling equilibrium. The crosslink density was calculated by using the Flory-Rehner equation (eqn (1)), given as:

$$v_{\rm e} = \frac{\rho_{\rm d}}{M_{\rm c}} = -\frac{\ln(1 - v_{\rm r}) + v_{\rm r} + \chi_1 v_{\rm r}^2}{M_{\rm c} \left(v_{\rm r}^{\frac{1}{3}} - \frac{v_{\rm r}}{2}\right)} \tag{1}$$

where χ_1 is the Flory-Huggins interaction parameter between rubber and toluene (0.391), M_c is the molecular weight of

Table 1 Formulation of natural rubber compounds

Ingredients	phr^a
NR	100
Silica	10/30/50
Glucose	10% of silica filler
ZnO	5
Stearic acid	3
Soluble sulphur	1.5
TBBS	1.5

^a Parts per hundred grams of rubber.

polymer between crosslinks, ν_r is the molar volume of toluene (106.3 cm³ mol⁻¹), and ν_e is the crosslink density (mol cm⁻³). The below-mentioned equation (eqn (2)) was used to determine the volume percentage of rubber (ν_r) in the swollen network.

$$v_{\rm r} = \frac{\frac{w_1}{\rho_{\rm d}}}{\frac{w_1}{\rho_{\rm d}} + \frac{(w_2 - w_1)}{\rho_{\rm c}}} \tag{2}$$

where w_1 and w_2 represent the rubber's mass before and after swelling, respectively. Toluene's density is expressed as $\rho_{\rm s}$ (0.8669 g mL $^{-1}$), and the rubber's pre-swelling density is expressed as $\rho_{\rm d}$. Using the rectangle-shaped samples measuring 25 mm \times 10 mm \times 2 mm, a dynamic mechanical analyzer (DMA) (Plus1000, Metravib, France) was used to perform the dynamic mechanical thermal analysis in the tension mode. To realize the dynamic mechanical properties, the full temperature sweep was performed for the NR vulcanizates.

Using rubber-state NMR (nuclear magnetic resonance) spectroscopy, the crosslinking junction structure of the natural rubber vulcanizates was determined. Prior to NMR tests, the vulcanizates of natural rubber were extracted using acetone for 48 hours using a Soxhlet system, and then they were dried for 72 hours in a vacuum oven. A JEOL ECA400 FT-NMR running at 99.55 MHz for $^{13}\mathrm{C}$ was used to conduct the rubber-state NMR experiments. The $^{13}\mathrm{C}$ NMR had a spinning rate of 18 \pm 0.02 kHz, an accumulation of 40 000 scans, and a pulse repetition duration of 5 s.

A FIB-SEM (SII SMI-3050SE) was used to observe the morphology of the silica-filled natural rubber vulcanizate at a 3 kV accelerating voltage. An extremely thin segment of the natural rubber vulcanizates was also formed using FIB-SEM (Focused Ion Beam-Scanning Electron Microscopy). A Sorvall Instruments MT6000 ultramicrotome operating at $-80~^{\circ}\text{C}$ was also utilized to prepare the ultra-thin portion of the polymer composite. After annealing the thin slices for 30 minutes at 353 K, they were dyed with either ruthenium tetroxide (RuO₄) or osmium tetraoxide (OsO₄).

Atomic force microscopy (AFM) analysis was performed using an Agilent 5500 (USA) instrument. The properly cleaned molded NR sheet samples were investigated to study the surface roughness of the vulcanizates and delineated as a phase morphology. The scanning of a 5 \times 5 μ m area was carried out using a SiN₄ (silicon nitride) tip in tapping mode at a resonance frequency of 150 kHz and 42 N m⁻¹ force constant, respectively.

Results and discussion

3.1. Cure characteristics

The cure curves and data of the natural rubber compounds are depicted in Fig. 1 and Table 2, respectively. It is well-known and proven that the presence of silica in rubber vulcanizates exhibits a retarding effect during vulcanization, which is attributed to the adsorption of curing accelerators and activators on the acidic surface of silica particles. 31,32

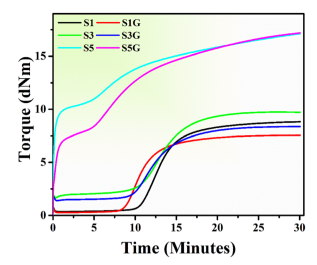


Fig. 1 Torque vs. time plot.

Therefore, the silica-incorporated standard compounds (*i.e.*, S1, S3, and S5) displayed a comparatively higher optimum cure time ($T_{\rm c90}$ – optimum cure time) compared to the glucose-treated compounds. The incorporation of glucose in silica-filled NR compounds interestingly resulted in a reduction in the $T_{\rm c90}$ value, which may be due to the interaction between the amine groups of proteins present in NR and aldehyde groups of glucose, known as the Maillard reaction. Moreover, due to the maximum silica loading in samples S5 and S5G, the obtained initial torque was so high. Because of this, an initial hump can be observed in the cure curve of these 50 phr silica-loaded samples. As glucose has a plasticization effect, the initial torque of S5G was comparatively less than that of the S5 compound.

3.2. Mechanical and dynamic properties

The hardness and modulus of the vulcanizates are depicted in Fig. 4a and b. A gradual increment in both hardness and modulus was observed with the loading of silica, except for a sudden drop in the hardness value of the glucose-treated 50 phr silica loaded sample. This might be because of the plasticization effect of glucose at higher silica loading, as the phr of the embedded glucose was enhanced with the increment of silica loading. Concurrently, the modulus of the same vulcanizate (S5G) interestingly followed its own path towards a higher value.

This may be owing to the Maillard reaction (Fig. 2), which resulted in well-networked NR vulcanizates in comparison with

Table 2 Cure characteristics of the rubber compounds

Sampl	es T_{c90} (min	$(n)^a T_{s2} $ (min	$M_{\rm H} (dNr)$	m) ^c M_L (dN:	$(m)^d M_H - M_L$	(dNm)
S1	17.82	11.64	8.83	0.34	8.49	
S1G	15.43	9.71	7.56	0.27	7.29	
S3	17.9	11.54	9.74	1.67	8.07	
S3G	17.85	11.07	8.37	1.39	6.98	
S5	20.94	0.24	17.11	5.86	11.25	
S5G	20	0.39	17.19	3.14	14.05	

 a $T_{\rm c90}$ – optimum cure time. b $T_{\rm s2}$ – scorch safety. c $M_{\rm H}$ – maximum torque. d $M_{\rm L}$ – minimum torque.

Fig. 2 Maillard reaction.

the untreated standard samples (the mechanism of a wellnetworked natural rubber in the presence of glucose as well as the silica filler is revealed in Fig. 3). Also, the formation of a Schiff base complex due to interaction between MBT (2-mercaptobenzothiazole) originating from TBBS during vulcanization and the carbonyl group of glucose might be another reason.²

The stress vs. strain plot is shown in Fig. 5, where the glucose-treated compounds' higher strength was evident. The tensile strength of the untreated vulcanizates was found to be maximum for the 10 phr silica-loaded compound. Thereafter, deterioration in the tensile strength was observed upon further addition of the silica filler due to the self-agglomeration of the silica particles in the NR matrix (Fig. 6a). 20,33 However, on the other hand, the incorporation of a few phr of glucose exhibited a remarkable improvement in tensile strength compared to the untreated standard compounds, without much difference in elongation at break. This is attributed to a better polymer-filler interaction and silica dispersion in the presence of glucose. 34-36

A gradual rise in tear strength can be observed (Fig. 6c) in the case of glucose-incorporated NR vulcanizates. This improvement in tear strength is indeed another proof of facilitation of silica dispersion in the presence of glucose compared to the untreated

standard compounds. Moreover, a well filler dispersed rubber composite system presents a higher crosslink density value. This is attributed to a superior polymer-filler interaction in silicafilled NR vulcanizates. Therefore, a similar kind of trend in crosslink density can be observed in Fig. 6d. 28,37,38

Essential parameters like polymer-filler and filler-filler interactions affect the reinforcement in the rubber matrix. The filler-filler interactions can be determined from the variation of storage modulus with strain, which is generally measured as the Payne effect, and the same denotes the difference between storage modulus at minimum and maximum strains. A higher difference indicates a greater Payne effect value, which means a higher filler-filler interaction.

Fig. 7a, b and c, d present the strain sweep and $\Delta E'$ (Payne effect) plots of the NR vulcanizates at room temperature and at 70 °C, respectively. As per the previous assumption from mechanical properties, it is now quite clear that the presence of glucose actually helped in dispersing the silica filler in the NR matrix. Therefore, a lower $\Delta E'$ value can be noticed for the glucose-treated vulcanizates compared to the untreated standard compounds. This is ascribed to a superior polymer-filler interaction and comparatively uniform dispersion of the silica filler in the NR matrix compared to the untreated samples. 20,39,40 The trend of the Payne effect at both room temperature and high temperature seems to be similar, which confirms the same effectiveness of glucose in the NR composite at high temperature as well.

Dynamic mechanical analysis and temperature sweep plots are depicted in Fig. 8. In the case of tyre application to predict

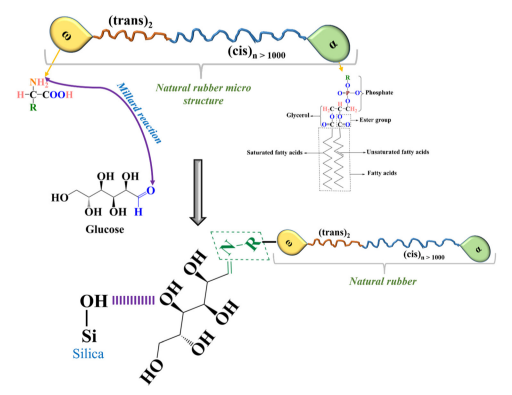
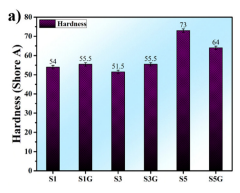
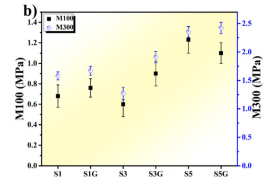
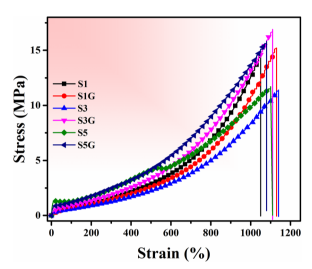


Fig. 3 Interactions between natural rubber, glucose, and the silica filler in the system.





(a) Hardness and (b) modulus plots of the NR vulcanizates.



Stress vs. strain plot of the NR vulcanizates

the rolling resistance and wet traction of the tyre compound on the lab scale, the $\tan \delta$ vs. temperature plot generated from DMA temperature sweep analysis is a general initial trend. The $\tan \delta$ at 60 to 70 °C indicates the rolling resistance of a tyre, and $\tan \delta$ at 0 °C is used to assume the wet grip of the tyre. Thus, in accordance with the tan δ vs. temperature plot (Fig. 8a) it was clearly noticed that the $\tan\delta$ at 70 $^{\circ}\mathrm{C}$ is actually reduced in the presence of glucose rather than the untreated standard sample. This signified a significant reduction in rolling resistance of the tyre tread compound. This is attributed to a better polymerfiller interaction, as confirmed by the strain sweep study of the NR vulcanizates. On the contrary, there is not much difference in tan δ at 0 °C between the treated and untreated NR vulcanizates. However, the wet grip in the case of heavy vehicle tyres is a secondary concern, unlike light weight vehicle tyres like two wheeler or passenger car tyres. On the other hand, the storage modulus (E') vs. temperature plot is shown in Fig. 8b. The stability of a heavy vehicle tyre is a very important factor, especially during the running condition of the vehicle. The storage modulus (E') at high temperature (60 to 70 °C) signifies the same thing; a high value of E' at high temperature indicates a superior stability of the tyre tread compound. However, in this

study, a gradual rise of the E' value was noticed with the enhancement of silica loading, which is owing to a higher restricted motion of the elastomer molecules in conjunction with the increment of silica loading. Moreover, the value of E'at 70 °C for 10 phr and 50 phr silica-loaded compounds did not display any difference among the treated and untreated samples. 33,39,41

Nonetheless, surprisingly the glucose-treated 30 phr silica loaded NR vulcanizate exhibited a slight upliftment of the E' value at 70 °C compared to the standard compound. The better polymer-filler interaction can build a highly networked structure into the system, signifying a higher crosslink density value, which could be the major reason behind the mentioned incident.

Elastomer molecule confinement effects in composite systems can control composites' characteristics and consequent uses. One of the main elements influencing mechanical characteristics in the presence of fillers is the orientation of elastomer chains. The confinement effect is directly related to the polymer's segmental and chain mobilities, which can be clearly understood from the temperature sweep plot (Fig. 8a) of DMA analysis. It is observed that the addition of silica filler resulted in the decrease in the magnitude of the $\tan \delta$ peak, indicating the restricted mobility of polymer chains. The NR chains' confinement by silica fillers is responsible for the decrease of the $\tan \delta$ peak values in the NR composites. The confined fraction of NR chains varies depending on the weight percentage of the silica filler. A decrease in the values of the $\tan \delta$ peak suggests that the NR-silica composites' interfacial reinforcing has improved. However, the further reduction of the $\tan \delta$ peak in the presence of glucose signifies the significant improvement in the interfacial reinforcement in the NRsilica composites, i.e., a better polymer-filler interaction compared to the composites without glucose. 42-44

Compared to the low silica-filled NR composite, the $\log E' \nu s$. temperature plot (Fig. 8b) of high silica-loaded NR composites provides valuable information concerning the rise in the storage modulus at the high-temperature area. The values of E' climb when the amount of silica filler is increased, and the addition of glucose causes the storage modulus to rise more with each silica loading. This demonstrates unequivocally that the silica filler is evenly distributed throughout the NR matrix.

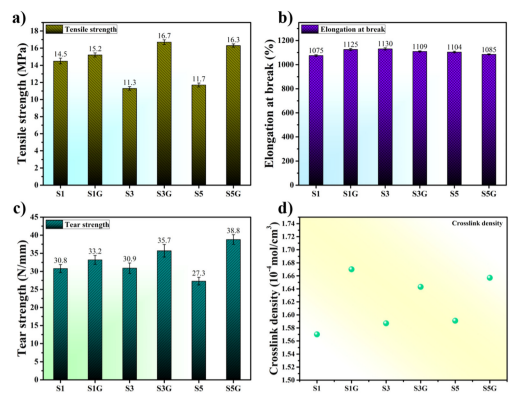


Fig. 6 (a) Tensile, (b) elongation, (c) tear strength, and (d) crosslink density plots of the NR vulcanizates

This highlights that there is a good rubber-silica interaction and that the reinforcement is reasonably strong when silica and glucose are present, which limits the mobility of the chain segments. 43-45

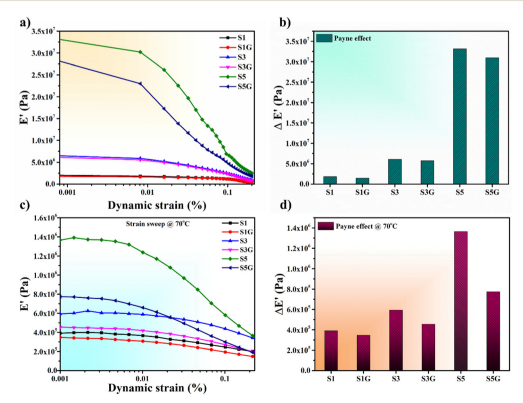


Fig. 7 (a) Strain sweep, (b) Payne effect plot at room temperature and (c) strain sweep, (d) Payne effect plot at 70 °C of the NR vulcanizates.

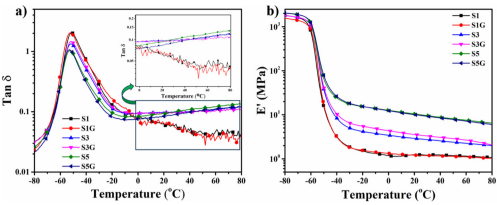


Fig. 8 DMA temperature sweep plot – (a) $\tan \delta vs$. temperature plot and (b) storage modulus vs. temperature plot of the NR vulcanizates.

3.3. Rubber state NMR

The Fig. 9 shows rubber state ¹³C-NMR spectra of silica-filled NR vulcanizates and silica-glucose filled NR vulcanizates together with the solution state ¹³C-NMR spectrum of unvulcanized NR. Large signals appeared at 23.4, 26.0, 32.0, 125.0, and 135.0 ppm (parts per million) in all spectra. These signals were assigned to C5, C4, C1, C3, and C2 of the *cis*-1,4-isoprene unit, according to previous reports. In addition, small signals appeared between 35 and 60 ppm for the NR vulcanizates, even though they did not appear for the unvulcanized NR.

Fig. 10 shows the enlarged rubber state ¹³C-NMR spectra between 35 and 60 ppm. The small signals appeared at 37.0, 40.0, 44.5, 50.0, 50.5, 57.3, 58.0, and 58.5 ppm in the rubber state ¹³C-NMR spectra of the silica-filled NR vulcanizates and silica-glucose filled NR vulcanizates. These signals were assigned to carbons linked to sulfur as crosslinking junctions and their adjacent carbons, as shown in Fig. 11.

The signals at 50.0 and 50.5 ppm were assigned to quaternary carbon linked to sulfur (quaternary C–S) and tertiary carbon

C2 C3 C4 C5

(g) 50 phr silica + 5 phr G

(h) 50 phr silica

(e) 30 phr silica + 3 phr G

(d) 30 phr silica

(e) 10 phr silica + 1 phr G

(b) 10 phr silica

(a) NR

Fig. 9 (a) The solution-state 13 C NMR spectrum of unvulcanized NR and the rubber-state 13 C NMR spectra of the vulcanized natural rubbers prepared with silica and with silica and glucose. (b) 10 phr silica, (c) 10 phr silica with 1 phr glucose, (d) 30 phr silica, (e) 30 phr silica with 3 phr glucose, (f) 50 phr silica, and (g) 50 phr silica with 5 phr glucose, together with the solution-state 13 C NMR spectrum of unvulcanized natural rubber.

linked to sulfur (tertiary C-S), respectively, which were formed by direct addition of sulfur to the carbon-carbon double bond of the cis-1,4-isoprene unit. In contrast, the signals at 37.0 ppm were assigned to methylene groups adjacent to the quaternary C-S and tertiary C-S. In contrast, the signal at 57.3 ppm was assigned to quaternary allylic carbon linked to sulfur (quaternary allylic C-S), which was formed by recombination of the sulfur radical with the tertiary allylic carbon radical generated by abstracting hydrogen from allylic carbon at the 4th position (C4) of the cis-1,4-isoprene unit followed by conjugation, whereas the signals at 58.0 and 58.5 ppm were assigned to tertiary allylic carbon linked to polysulfide (tertiary allylic C-Sx) and tertiary allylic carbon linked to monosulfide (tertiary allylic C-Sm), respectively, which were formed by recombination of sulfur radicals with the secondary allylic carbon radicals generated by abstracting hydrogen from allylic carbon at the 1st position (C1) and 4th position (C4) of the cis-1,4-isoprene unit.46-48 The signals at 44.5 ppm were assigned to methylene

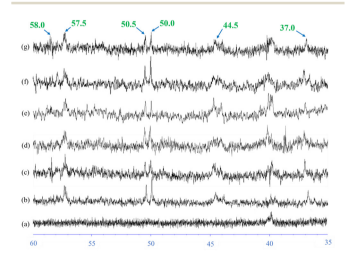


Fig. 10 Enlarged rubber-state ¹³C NMR spectrum: (a) the solution-state ¹³C NMR spectrum of unvulcanized NR and the rubber-state ¹³C NMR spectra of the vulcanized natural rubbers prepared with silica and with silica and glucose, (b) 10 phr silica, (c) 10 phr silica with 1 phr glucose, (d) 30 phr silica, (e) 30 phr silica with 3 phr glucose, (f) 50 phr silica and (g) 50 phr silica with 5 phr glucose, together with the solution-state ¹³C NMR spectrum of unvulcanized natural rubber.

Paper

Fig. 11 Assignment of signals in the rubber-state ¹³C NMR spectra of silica filled NR vulcanizates and silica-glucose filled NR vulcanizates.

groups adjacent to the quaternary allylic C-S, tertiary allylic C-Sx and tertiary allylic C-Sm, whereas the signal at 40 ppm was assigned to the methylene group (C1) of the trans-1,4-isoprene unit generated by cis-trans isomerization of the cis-1,4-isoprene unit as a side reaction. In the present study, we found that the signals at 37.0, 40.0, 44.5, 50.0, 50.5, 57.3, 58.0, and 58.5 ppm similarly appeared in the rubber state ¹³C-NMR spectra of silicafilled NR vulcanizates and silica-glucose filled NR vulcanizates. To be more exact, the signal at 58.0 ppm was more prominent for the silica-filled NR vulcanizates containing a larger amount of the tertiary allylic C-Sx, whereas the signal at 58.5 ppm was more prominent for the silica-glucose filled NR vulcanizates containing a larger amount of the tertiary allylic C-S. 46,47

Rubber state NMR spectroscopy applied in the present study makes it possible to detect the gap of the energy levels split due

to the Zeeman effect as a chemical shift value with respect to carbons and hydrogens in different chemical environments since a residual dipole-dipole interaction of the NR vulcanizates is eliminated by fast magic angle spinning of about 20 kHz. This implies that the carbons and hydrogens in the different chemical environments are quantitatively analyzed with intensities of signals appearing in the rubber state NMR spectra. 49 Therefore, we estimated the intensity ratios of the signals at 50.0, 50.5, 57.3, 58.0 and 58.5 to the signal at 23.4 ppm (C5 of the cis-1,4-isoprene unit). Fig. 12 shows the estimated values of the intensity ratios of the signals at 50.0, 50.5, 57.3, 58.0, and 58.5 ppm for the silica-filled NR vulcanizates and silica-glucose filled NR vulcanizates. First, the intensity ratio value of the signal at 57.3 ppm, which was the largest among the intensity ratio values, was independent of amounts of silica and silica-glucose; that is, the intensity ratio value was almost the same for the silica-filled NR vulcanizates and silicaglucose filled NR vulcanizates. Second, the intensity ratio value of the signal at 58.0 ppm was the highest for the 50 phr silicafilled NR vulcanizate, whereas that of the signal at 58.5 ppm was high for the 30 phr silica-3 phr glucose-filled NR vulcanizate and the 50 phr silica-5 phr glucose filled NR vulcanizate. These correspond to the values of crosslink density. $^{50-52}$

Third, the intensity ratio values of the signals at 50 and 50.5 ppm were larger for silica 30 phr and silica 50 phr filled NR vulcanizates. In the previous study, Kashihara and collaborators reported that the signals at 50 and 50.5 ppm were partly assigned to episulfides or epipolysulfides formed by the direct addition of sulfur to the carbon-carbon double bond of the cis-1,4-isoprene unit, even though they were also assigned to the structures shown in Fig. 11.53,54 The formation of episulfides and epipolysulfides as heteroalkene rings implies that a large number of carbon-sulfur linkages may not contribute to

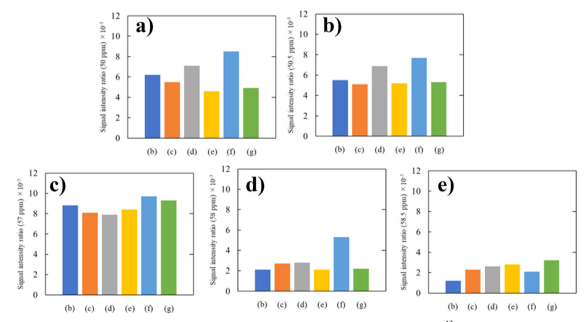


Fig. 12 Intensity ratio of signals at (a) 50.0, (b) 50.5, (c) 57.3, (d) 58.0, and (e) 58.5 ppm in the rubber-state ¹³C-NMR spectra

crosslinking as crosslinking junctions, suggesting that the vulcanization is not performed efficiently. This corresponds to the fact that the tensile strength, tear strength, and crosslink density of the silica-filled NR vulcanizates were lower than those of the silica-glucose filled NR vulcanizates. These results show that the outstanding mechanical properties of silicaglucose filled NR vulcanizates were found to be due to the ability of glucose to promote the vulcanization of natural rubber efficiently. 52-54

3.4. Morphology analysis

Fig. 13 shows FIB-SEM images of the 10 phr silica-filled NR vulcanizate and 10 phr silica-1 phr glucose-filled NR vulcanizate, for which elemental analysis was performed from the surface to inside after making a cross-section through FIB processing with the Ga ion. The red domains represent silicon, the yellow domains represent oxygen, and the green domains represent sulfur. As for the 10 phr silica-filled NR vulcanizate, concentrations of silicon, oxygen, and sulfur were higher on the vulcanizate's surface than inside, reflecting surface segregation of silica and sulfur-containing compounds due to their heterogeneous distributions. Furthermore, silicon, oxygen, and sulfur appeared at the same position in the FIB-SEM image. This may be explained to be due to adsorption of TBBS, a sulfur-containing compound, on the surface of SiO2, as reported by Choi and collaborators. 55 By contrast, for the 10 phr silica-1 phr glucosefilled NR vulcanizate, silicon and oxygen existed sparsely on the surface of the vulcanizate as particles of approximately 0.1 µm in diameter at the same position, indicating that they are SiO2, but sulfur appeared evenly on the surface due to a good mixing. Comparing the FIB-SEM image of the 10 phr silica-filled NR vulcanizate with that of the 10 phr silica-1 phr glucose filled NR vulcanizate, we found that glucose played an important role in

the dispersion of not only silica but also TBBS onto natural rubber. In particular, glucose prevented the adsorption of TBBS onto the surface of SiO₂. 55,56

Fig. 14 shows FIB-SEM images of the 30 phr silica filled NR vulcanizate and the 30 phr silica-3 phr glucose filled NR vulcanizate. For the 30 phr silica-filled NR vulcanizate, silica and TBBS adsorbed on silica aggregated to cover the surface of the vulcanizate. By contrast, for the 30 phr silica-3 phr glucose filled NR vulcanizate, silica particles with a diameter of about 0.2 µm existed sparsely on the surface of the vulcanizate, whereas sulfur was finely dispersed homogeneously.

Fig. 15 shows FIB-SEM images of the 50 phr silica filled NR vulcanizate and the 50 phr silica-5 phr glucose filled NR vulcanizate. For the 50 phr silica-filled NR vulcanizate, silica with TBBS (silica-TBBS) was densely aggregated to cover the surface of the vulcanizate, and the silica and TBBS were interconnected to each other. This corresponds to the fact that the hardness of the 50 phr silica-filled NR vulcanizate was the highest among the vulcanizates. By contrast, for the 50 phr silica-5 phr glucose filled NR vulcanizate, silica particles with a diameter of about 0.2 µm existed sparsely on the surface of the vulcanizate, whereas sulfur was finely dispersed as in the case of the 30 phr silica-3 phr glucose filled NR vulcanizate.⁵⁶

These results imply that glucose was adsorbed on the surface of silica to prevent the adsorption of TBBS and to enable the homogeneous dispersion of silica in the NR vulcanizates. The vulcanization was, thus, promoted to form crosslinking junctions by suppressing side reactions that form episulfides and epipolysulfides as heteroalkene rings and dangling polysulfides, since TBBS efficiently formed a complex with Zn. The higher crosslink density and outstanding mechanical properties of silica-glucose filled NR vulcanizates were attributed to the ability of glucose to cover the surface of silica, which

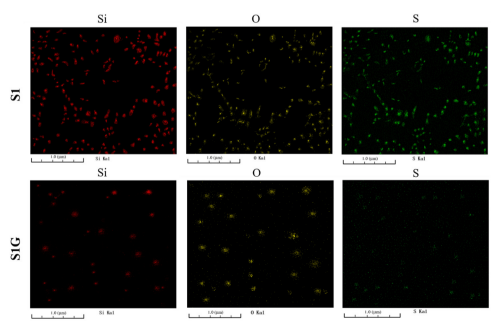


Fig. 13 FIB-SEM images of the 10 phr silica filled NR vulcanizate (S1) and the 10 phr silica-1 phr glucose filled NR vulcanizate (S1G).

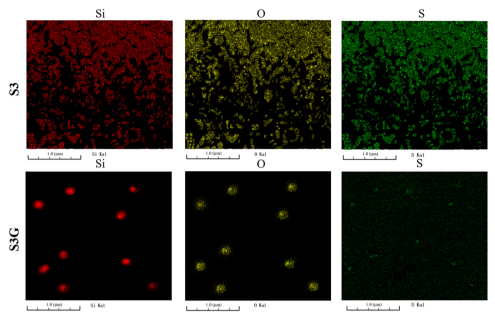


Fig. 14 FIB-SEM images of the 30 phr silica filled NR vulcanizate (S3) and the 30 phr silica-3 phr glucose filled NR vulcanizate (S3G).

resulted in free TBBS to form a Zn complex, thus efficiently promoting the accelerated sulfur vulcanization of natural rubber. Furthermore, glucose contributed to the good dispersion of silica in the NR vulcanizates. The hardness of the silica-filled NR vulcanizates increased without glucose even though the amounts of sulfur were the same in the compounded NR.⁵⁵

Fig. 16 depicts the AFM photomicrographs (left-height images and right-phase images) of the treated and untreated NR vulcanizates for a scan size of 15 μ m \times 15 μ m. In 3D (height) images the light brown colour represents the hard silica particles, and the dark brown colour indicates the elastomer matrix region.⁵⁷ However, a broader range of light colour regions can be clearly observed in the case of 3D photomicrographs of the untreated NR vulcanizates. This signifies a higher number of agglomerated silica particles in the untreated NR matrix, resulting in higher filler-filler interaction during dynamic testing, indeed supporting the results of strain sweep (Fig. 7). 58,59 Concurrently, the shorter ranges of the light colour region in glucose-treated samples exhibited comparatively smoother surfaces compared to the untreated samples.

This is attributed to a homogeneous dispersion of silica particles and a better polymer-filler interaction in the presence of glucose in the silica-filled NR matrix. 60,61 Therefore, a significant improvement in dynamic and mechanical properties

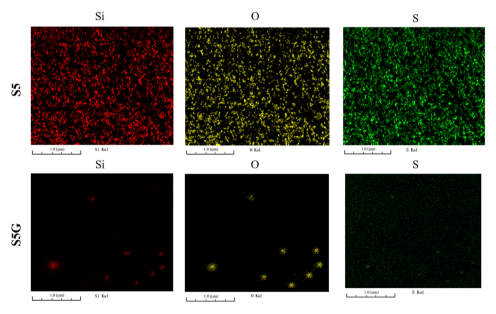


Fig. 15 FIB-SEM images of the 50 phr silica filled NR vulcanizate (S5) and the 50 phr silica-5 phr glucose filled NR vulcanizate (S5G).

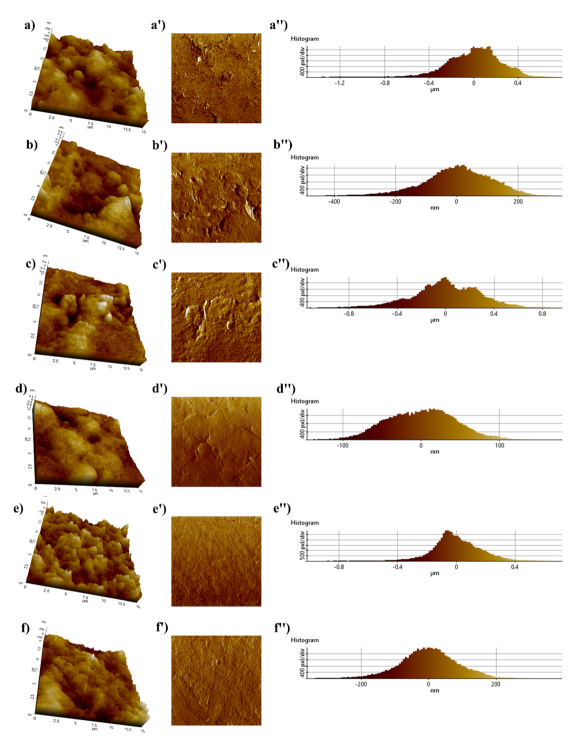


Fig. 16 AFM photomicrographs – (a), (a') and (a") 3D view, phase image and histogram of the 10 phr silica loaded sample, (b), (b') and (b") 3D view, phase image and histogram of the 30 phr silica loaded sample, (c), (c') and (c") 3D view, phase image and histogram of the 50 phr silica loaded sample, (d), (d') and (d") 3D view, phase image and histogram of the 10 phr silica loaded glucose treated sample, (e), (e') and (e") 3D view, phase image and histogram of the 30 phr silica loaded glucose treated sample, and (f), (f') and (f") 3D view, phase image and histogram of the 50 phr silica loaded glucose treated sample.

was witnessed. A similar kind of trend can be observed in the phase images and histograms, which indeed support the trend of 3D images of AFM photomicrographs.

This work has significant potential for industrial applications. The compound can primarily be used as a tyre tread compound to produce green tyres, as it exhibits low rolling resistance along with excellent mechanical properties even at high silica loading. It is well known that glucose is an inexpensive carbohydrate, readily available in nature. Therefore, incorporating small amounts of glucose in silica-filled natural rubber results in a more homogeneous dispersion of the silica filler, compared to using costly chemicals or coupling agents.

This formulation can be used for production trials by preparing a 70 kg batch-weight compound and mixing it in a large internal mixer. The only additional step is the inclusion of glucose during the mastication of rubber at the initial stage of mixing, which is a simple and easy process that significantly enhances the properties. The production process should be highly compatible with existing methods, as the presence of glucose also facilitates

better processing due to its plasticizing nature.

4. Conclusions

Paper

The incorporation of glucose in a silica-filled natural rubber compound discloses a hidden way to improve silica dispersion in a natural rubber matrix. The investigation of natural rubber bio-chemistry revealed the microstructure of NR, where the functional groups like proteins and phospholipids are located at two opposite ends of the natural rubber microstructure. Now, the incorporation of glucose in a silica-filled NR compound results in the interaction between carbonyl groups of glucose and amine groups of proteins, which is known as the Maillard reaction. This is the primary reason for the reduction of T_{c90} in the presence of glucose in NR vulcanizates. Additionally, a gradual enhancement in hardness and M300 (modulus at 300%) can be observed along with the increment of silica loading. Moreover, the incorporation of glucose exhibited comparatively higher values of both the hardness and modulus except for the hardness and M100 at 50 phr loading of silica. This is attributed to the plasticization effect of glucose at higher loading. A significant improvement in mechanical properties, like tensile strength, tear strength, and crosslink density, could be observed in the presence of glucose in every loading of silica due to the Maillard reaction. This assisted in a better silica dispersion in the NR matrix, resulting in a superior polymerfiller interaction. The Payne effect study showed the same in the presence of glucose, especially at higher loading. Moreover, the strain sweep analysis at high temperature (70 °C) exhibited a similar kind of improvement in polymer-filler interaction to that at room temperature, as confirmed by the Payne effect analysis. This confirms the effectiveness of glucose in NR vulcanizates even at high temperatures. DMA temperature sweep, another vital dynamic property, is an essential test for tyre applications, where the $\tan \delta$ at 70 °C indicates that the rolling resistance of a heavy vehicle tyre is found to be decreased for the glucose-incorporated compounds, without any deterioration in the storage modulus at high temperature. Moreover, a detailed rubber state 13C NMR study unveiled inefficient sulfur vulcanization of silica-filled NR compounds in the absence of glucose. This is attributed to the formation of episulfides and epipolysulfides as heteroalkene rings, implying that a higher number of C-S linkages may not contribute to crosslink or crosslinking junctions. On the other hand, an exceptional improvement in both mechanical and dynamic properties can be observed in the presence of glucose, as it promoted the vulcanization of the NR matrix efficiently. The homogeneous dispersion of the silica filler in the NR matrix is

distinctly evident from the in-depth morphological analyses, *i.e.*, FIB-SEM and AFM analyses.

This work unveils the significant contribution of glucose to the uniform dispersion of the silica filler. It also provides strong proof of the Maillard reaction between the amine groups of NR and carbonyl groups of glucose. Therefore, this simple and sustainable step of adding glucose in silica-filled natural rubber compounds opens a new door towards preparing green and sustainable tyre and non-tyre products.

Author contributions

Abhijit Bera: conceptualization, writing – original draft preparation, writing – reviewing and editing; Masaki Yamano: conceptualization, writing – original draft preparation, writing – reviewing and editing; Seiichi Kawahara: conceptualization, writing – reviewing and editing, and overall supervision; and Santanu Chattopadhyay: conceptualization, writing – reviewing and editing, and overall supervision.

Data availability

Data will be available on special request.

Conflicts of interest

The authors declare that there are no conflicts of interest to declare.

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References

- 1 Y. Zhou, K. Kosugi, Y. Yamamoto and S. Kawahara, *Polym. Adv. Technol.*, 2017, 28, 159–165.
- 2 A. Nimpaiboon, M. Sriring, S. Kumarn and J. Sakdapipanich, *J. Appl. Polym. Sci.*, 2020, **137**, 1–11.
- 3 A. G. Thomas and J. M. Whittle, *Rubber Chem. Technol.*, 1970, 43, 222-228.

4 A. H. Eng, S. Kawahara and Y. Tanaka, Rubber Chem. Technol., 1994, 67, 159-168.

Materials Advances

- 5 S. Kawahara, T. Kakubo, J. T. Sakdapipanich, Y. Isono and Y. Tanaka, Polymer, 2000, 41, 7483-7488.
- 6 A. Bera, D. Ganguly, J. P. Rath, S. Ramakrishnan, J. Kuriakose, S. K. P. Amarnath and S. Chattopadhyay, Mater. Chem. Phys., 2023, 295, 127151.
- 7 A. Bera, D. Ganguly, S. K. Ghorai, J. P. Rath, S. Ramakrishnan, J. Kuriakose, S. K. P. Amarnath and S. Chattopadhyay, Chem. Eng. J. Adv., 2022, 11, 100349.
- 8 A. Bera, B. Manna, D. Ganguly, S. K. P. Amarnath, S. Nanda, A. Ghosh and S. Chattopadhyay, ACS Appl. Polym. Mater., 2022, 5, 451.
- 9 A. Bera, K. Sarkar, D. Ganguly, S. K. Ghorai, R. Hore, N. Kumar, S. K. P. Amarnath and S. Chattopadhyay, J. Polym. Res., 2024, 31, 1-21.
- 10 Y. Xiao, Z. Xu, Z. Gong, B. Li, Y. Huang, H. Bian and C. Wang, J. Appl. Polym. Sci., 2023, 140, e54049.
- 11 P. Nuinu, C. Sirisinha, K. Suchiva, P. Daniel and P. Phinyocheep, J. Mater. Res. Technol., 2023, 24, 2155-2168.
- 12 A. Guchait, D. Ganguly, C. Sengupta, S. Chattopadhyay and T. Mondal, ACS Sustainable Chem. Eng., 2022, 10, 16780-16792.
- 13 A. Bera, M. Goswami, D. Ganguly, J. P. Rath, S. Ramakrishnan, J. Kuriakose, S. K. P. Amarnath and S. Chattopadhyay, J. Mater. Sci., 2023, 58, 996-1011.
- 14 W. Kaewsakul, K. Sahakaro, W. K. Dierkes and J. W. M. Noordermeer, *Polym. Eng. Sci.*, 2015, 836–842.
- 15 T. Jiang, T. Kuila, N. H. Kim, B. C. Ku and J. H. Lee, Compos. Sci. Technol., 2013, 79, 115-125.
- 16 C. Hayichelaeh, L. A. E. M. Reuvekamp, W. K. Dierkes, A. Blume, J. W. M. Noordermeer and K. Sahakaro, Rubber Chem. Technol., 2020, 93, 195-207.
- 17 C. P. Tripp and M. L. Hair, J. Phys. Chem., 1993, 97, 5693-5698.
- 18 J. Zheng, D. Han, X. Ye, X. Wu, Y. Wu, Y. Wang and L. Zhang, Polymer, 2018, 135, 200-210.
- 19 S. Das, S. Chattopadhyay, S. Dhanania and A. K. Bhowmick, Polym. Eng. Sci., 2020, 60, 3115-3134.
- 20 W. Kaewsakul, K. Sahakaro, W. K. Dierkes and J. W. M. Noordermeer, Rubber Chem. Technol., 2013, 86, 313-329.
- 21 S. O. Movahed, A. Arsarifar and M. Song, Polym. Int., 2009, 58, 209-217.
- 22 T. Xu, Z. Jia, J. Li, Y. Luo, D. Jia and Z. Peng, Polym. Compos., 2018, 39, 377-385.
- 23 P. Manoharan and K. Naskar, Polym. Compos., 2019, 40, 871-883.
- 24 P. K. Chattopadhyay, U. Basuli and S. Chattopadhyay, Polym. Compos., 2010, 31, 835-846.
- 25 K. Katueangngan, T. Tulyapitak and A. Saetung, Procedia Chem., 2016, 19, 447-454.
- 26 S. S. Sarkawi, W. K. Dierkes and J. W. M. Noordermeer, Rubber World, 2012, 247, 26-31.
- 27 D. Mahata, K. Sarkar, P. Mondal, O. Prabhavale, S. Dhanania, G. B. Nando and S. Chattopadhyay, Iran. Polym. J., 2020, 29, 393-401.

- 28 S. S. Sarkawi, W. K. Dierkes and J. W. M. Noordermeer, Eur. Polym. I., 2013, 49, 3199-3209.
- 29 A. Nimpaiboon and J. Sakdapipanich, Polym. Test., 2013, 32, 1408-1416.
- 30 X. Wang, Z. Luo, Y. Xu, J. Zhong, H. Zhang and J. Liang, J. Appl. Polym. Sci., 2023, 140, e53966.
- 31 Y. Wang, L. Liao, J. Zhong, D. He, K. Xu, C. Yang, Y. Luo and Z. Peng, J. Appl. Polym. Sci., 2016, 133, 1-9.
- 32 N. Rattanasom, T. Saowapark and C. Deeprasertkul, Polym. Test., 2007, 26, 369-377.
- 33 S. Prasertsri and N. Rattanasom, Polym. Test., 2011, 30, 515-526.
- 34 A. Kraibut, W. Kaewsakul, K. Sahakaro, S. Saiwari, J. W. M. Noordermeer and W. K. Dierkes, Mater., 2024,
- 35 Y. Jiang, S. Wang and Y. Zhang, Polym. Bull., 2023, 80, 12373-12392.
- 36 M. Sedlačík, B. You and S. Jin, *Mater.*, 2024, 17, 3131.
- 37 S. Prasertsri and N. Rattanasom, Polym. Test., 2012, 31, 593-605.
- 38 J. Y. Lee, N. Park, S. Lim, B. Ahn, W. Kim, H. Moon, H. J. Paik and W. Kim, Adv. Mater. Sci. Eng., 2016, 24, 711-727.
- 39 S. Sattayanurak, J. W. M. Noordermeer, K. Sahakaro, W. Kaewsakul, W. K. Dierkes and A. Blume, Adv. Mater. Sci. Eng., 2019, 2019, 1-8.
- 40 H. Yao, G. Weng, Y. Liu, K. Fu, A. Chang and Z. R. Chen, J. Appl. Polym. Sci., 2015, 132(20), DOI: 10.1002/APP.41980.
- 41 Y. Li, B. Han, S. Wen, Y. Lu, H. Yang, L. Zhang and L. Liu, Composites, Part A, 2014, 62, 52-59.
- 42 C. M. Roland, Viscoelastic Behav. Rubbery Mater., 2011, 298-318.
- 43 V. Krikorian and D. J. Pochan, Macromolecules, 2004, 37, 6480-6491.
- 44 P. Bindu and S. Thomas, J. Phys. Chem. B, 2013, 117, 12632-12648
- 45 L. Huang, F. Yu, Y. Liu, A. Lu, Z. Song, W. Liu, Y. Xiong, H. He, S. Li, X. Zhao, S. Cui and C. Zhu, Compos. Sci. Technol., 2023, 233, 109905.
- 46 T. Saito, M. Yamano, K. Nakayama and S. Kawahara, Polym. Test., 2021, 96, 107130.
- 47 NMR methods for characterization of synthetic and natural polymers, ed. R. Zhang, T. Miyoshi and P. Sun, Royal Society of Chemistry, 2019.
- 48 A. K. Mishra, S. Chattopadhyay, P. R. Rajamohanan and G. B. Nando, Polymer, 2011, 52, 1071-1083.
- 49 H. T. Luu and S. Kawahara, ACS Sustainable Chem. Eng., 2024, 12(8), 3279-3288.
- 50 Y. Iizuka, Y. Yamamoto and S. Kawahara, Colloid Polym. Sci., 2019, 297, 133-139.
- 51 Y. Akahori, M. Hiza, S. Yamaguchi and S. Kawahara, Rubber Chem. Technol., 2021, 94, 657-668.
- 52 M. Yamano, Y. Yamamoto, T. Saito and S. Kawahara, Polymer, 2021, 235, 124271.
- 53 N. T. Thuong, P. T. Nghia and S. Kawahara, J. Sci. Technol. -Eng. Technol. Sustain. Dev., 2022, 32(2), 008-015.

Paper

54 B. Kim, K. Boonkerd and S. Kawahara, Polym. Adv. Technol., 2023, 34, 3003-3010.

- 55 L. Fukuhara, K. Kosugi, Y. Yamamoto, H. Jinnai, H. Nishioka, H. Ishii, M. Fukuda and S. Kawahara, Colloid Polym. Sci., 2015, 293, 2555-2563.
- 56 L. Fukuhara, K. Kosugi, Y. Yamamoto, H. Jinnai, H. Nishioka, H. Ishii and S. Kawahara, Polymer, 2015, 57, 143-149.
- 57 Y. Xiao, H. Zou, L. Zhang, X. Ye and D. Han, Polym. Test., 2020, 81, 106195.
- 58 L. L. Johnson, Rubber Chem. Technol., 2008, 81 359-383.
- 59 A. Bera, D. Ganguly, R. Hore, J. P. Rath, S. Ramakrishnan, J. Kuriakose, S. K. P. Amarnath and S. Chattopadhyay, J. Polym. Res., 2023, 30, 1-16.
- 60 T. Karino, Y. Ikeda, Y. Yasuda, S. Kohjiya and M. Shibayama, Biomacromolecules, 2007, 8, 693-699.
- 61 S. Salina Sarkawi, W. K. Dierkes and J. W. M. Noordermeer, Rubber Chem. Technol., 2015, 88, 359-372.