Ceramic Waste-Based Natural Rubber Composites: An Exciting Way for Improving Mechanical Properties

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ABSTRACT

Large amounts of fired ceramic waste produced in ceramic industry do not have a proper method to reuse and mainly thrown away into landfills since fired ceramics have already been sintered and thus their utilization as a raw material is limited. However, these solid wastes have a major environmental and economic concern. Thus, a proper management of such solid wastes is eminent. This research is dedicated to evaluating the possibility of using such ceramic waste as a low-cost filler material in the manufacture of natural rubber based composites. Ceramic particles smaller than 125 μm were selected initially for the preparation of natural rubber composites. For the latter part of the study, particles in sub-micrometer length scales were used. Elemental analyses and composition of the phases of the ceramic particles were determined by X-ray fluorescence and diffraction respectively. The average particle size was characterized by scanning electron microscope (SEM). The surfaces of the sub micrometer size ceramic particles were modified using Silane69 coupling agent. The surface modification was confirmed by Fourier-transform infrared spectroscopy, thermo gravimetric analysis and SEM coupled with energy-dispersive X-ray spectroscopy. Natural rubber based composites were prepared with different levels of ceramic filler loadings. The mechanical properties of the composites such as hardness, resilience, compression set, abrasion volume loss and tensile properties were improved. The reinforcing fillers significantly increase or maintain the strength of rubber composites, while the non-reinforcing fillers are helpful in reducing the cost. This work shows the composite nature of ceramic waste and its potential for rubber composites. The reinforcing fillers are helpful in reducing the cost [12, 13]. Fillers which contain chemically active surfaces from various chemical groups facilitate better interaction between rubber and filler by the formation of strong covalent bonding. The hydrophobic and hydrophilic nature of rubber and filler also influence the compatibility and dispersibility of filler in the rubber matrix.

Keywords: Ceramic Waste, Surface Modification, Rubber Composite, Reinforcement of Rubber

1. Introduction

An increase in the generation of different type of wastes can be observed due to the industrialization and rapid economic growth in recent decades [1]. Current practices in dumping or destroying waste in industries have badly affected in the natural environment including water, soil, air and noise pollution [12]. At the same time, these practices represent an economic cost. If waste is managed correctly it can be converted into a useful resource, which contributes to savings in raw materials, conserve action of natural resources and the climate, and promotes sustainable development [3].

According to the predicted statistics about 19 billion tons of solid wastes are expected to be generated annually by the year 2025 [2]. Annually, Asia alone generates 4.4 billion tons of solid wastes. A high demand, scarcity of raw materials, and high price of energy the cost of construction materials is increasing day by day [3]. So the use of alternative constituents in construction materials has now become a global concern [4-7]. If these ceramic wastes are utilized to re-use in industry, it will minimize the cost for raw materials and minimize the environmental pollution [3, 4].

The ceramics industry is mainly comprised of the following subsectors: wall and floor tiles, sanitary ware, bricks and roof tiles, refractory materials, technical ceramics and ceramic materials for domestic and ornamental use [4]. It was observed that ceramic wastes are the kind of major wastes in manufacturing sectors. Ceramic wastes are produced as a result of the ceramic processing. These wastes cause soil, air, and groundwater pollution. The main problem which ceramic waste inheriting is they are already sintered and cannot be easily reused due to their irreversible phase transition during the processing [8-10].

In this research work ceramic wastes from floor tiles and sanitary wares were planned to be composited with natural rubber latex.

Natural rubber has properties of plasticity or viscosity, elasticity, tensile strength. However, to improve the quality of the rubber several kinds of fillers are added to the natural rubber matrix [11]. The reinforcing fillers significantly increase or maintain the strength of rubber composites, while the non-reinforcing fillers are helpful in reducing the cost [12, 13]. Ceramic waste can be used to replace the fillers such like silica and carbon black. The filler compatibility in rubber composites depends on particle size, the structure of filler particles, and surface activity [12, 13]. Fillers which contain chemically active surfaces from various chemical groups facilitate better interaction between rubber and filler by the formation of strong covalent bonding. The hydrophobic and hydrophilic nature of rubber and filler also influence the compatibility and dispersibility of filler in the rubber matrix.

In this research waste ceramic was crushed and it was used as the filler material with natural rubber. Natural rubber so called cis-poly isoprene is a non-polar substance [14-16]. To overcome this polarity mismatch, the surface of the ceramic waste powder was modified using Silane69 (bis [3-triethoxysilylpropyl) tetrasulfane] coupling agent. In order to provide more available contact between the rubber molecule and the filler particle, the particle size of ceramic particles were greatly reduced. We have observed interesting results with different loadings of surface modified ceramic powders.
2. Experimental Methods

2.1 Preparation of Ceramic Waste Powder and Characterization

Ceramic floor tile waste and sanitary ware waste were collected from manufacturing sectors and domestically. Those waste were crushed and grinded using a mechanical method. Then the crushed ceramic wastes were sieved using mechanical sieve shaker (Aimil Ltd., India) and particles size less than 125 μm were obtained.

In the preparation of sub micrometer sized ceramic waste powder, initially prepared ceramic waste powder was ball milled for 6 hours at an 800 rpm speed by a laboratory planetary ball mill (YEMCAN, China) followed by dry milling [17-19].

Fourier transform infrared (FTIR) spectra of KBr pellets prepared from this initial ceramic powder were obtained using FTIR spectrophotometer (Bruker, USA). Elemental analyses and composition of the phases of these initial ceramic particles were determined by X-ray fluorescence spectrometer and X-ray diffractometer (Rigaku Ultima IV, USA) respectively. The average particle size was characterized by scanning electron microscope (EVOL S15, Germany).

2.2 Surface Modification of Ceramic Particles and Characterization

Ceramic powder surfaces were modified with Silane69 coupling agent with slight modifications of the methods followed by Wu et al. [16]. Firstly, 10 g of ceramic powder was added to an appropriate amount of ethanol in a round flask equipped with a magnetic stirrer rod and spherical condenser. After stirred for 30 min, the solution was heated to a predetermined temperature and a certain amount of Silane69 (commercial grade) was added (0.002 mol per 1 g of ceramic powder).

Then the reaction was maintained for 6 hours at 50 °C temperature. Finally, the product was purifed with ethanol, centrifuged and air dried.

The surface modification was confirmed by Fourier-transform infrared spectrometer (Bruker, USA), thermo gravimetric analyzer (SDFQ600, USA), x-ray fluorescence spectrometer and SEM coupled with energy-dispersive X-ray spectrometer (EVOL S15, Germany).

2.3 Preparation of Ceramic Filler Loaded Natural Rubber Composites

All the formulations were mixed using the two roll milling process and the vulcanization was done to prepare the final composites.

2.3.1 Preparation of Natural Rubber (NR) and Ceramic Waste (C125) (particle size ≤ 125 μm) Composites

Natural rubber based composite materials were prepared with different levels of ceramic filler loadings as shown in Table 1.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Compounding formulation for NR+C125 composites sample series</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ingredients</td>
<td>Units = parts per rubber</td>
</tr>
<tr>
<td>Sample ID</td>
<td>S0</td>
</tr>
<tr>
<td>NR</td>
<td>100</td>
</tr>
<tr>
<td>C125</td>
<td>0</td>
</tr>
<tr>
<td>ZnO</td>
<td>5</td>
</tr>
<tr>
<td>Stearic acid</td>
<td>2</td>
</tr>
<tr>
<td>TTBBS</td>
<td>2</td>
</tr>
<tr>
<td>TMTD</td>
<td>0.1</td>
</tr>
<tr>
<td>IPPD</td>
<td>0.5</td>
</tr>
<tr>
<td>Sulphur</td>
<td>2.5</td>
</tr>
<tr>
<td>Processing oil</td>
<td>7.5</td>
</tr>
</tbody>
</table>

2.3.2 Preparation of Natural Rubber (NR) and Surface Modified C125 Ceramic Powder with Silane (CS) Composite

Natural rubber based composite materials were prepared with different levels of ceramic filler loadings as shown in Table 2.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Compounding formulation for NR+CS composites sample series</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ingredients</td>
<td>Units = parts per rubber</td>
</tr>
<tr>
<td>Sample ID</td>
<td>S7</td>
</tr>
<tr>
<td>NR</td>
<td>100</td>
</tr>
<tr>
<td>CS</td>
<td>5</td>
</tr>
<tr>
<td>ZnO</td>
<td>5</td>
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<tr>
<td>Stearic acid</td>
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<td>2.5</td>
</tr>
<tr>
<td>Processing oil</td>
<td>7.5</td>
</tr>
</tbody>
</table>

2.4 Evaluation of Mechanical Properties of the Composites

All the prepared composites were vulcanized and test pieces were prepared using required standards.

2.4.1 Determination of Tensile Properties

The tensile properties of samples were determined by a Tensometer (Instron 3300, USA) material testing machine at a crosshead speed of 500 mm/min according to ISO 37: 2011 standards. All the tests were carried out at 27 ± 2 °C. Dumbbell shaped test pieces were cut from 2 mm thickness sheets. Then the specimen was placed on the grips of the testing machine and pulled at a rate of 50 mm/min until rupture. Four test pieces were used and average value was determined.

2.4.2 Determination of Tear Strength

The tear test was done using the Tensometer (Instron 3300, USA) according to ISO 34-1: 2011 standards. All the tests were carried out at 27 ± 2 °C. Crescent-shaped test pieces were cut from 2 mm thickness sheets. Then the specimen was placed on the grips of the testing machine and pulled at a rate of 50 mm/min until rupture. Four test pieces were used and average value was determined.

2.4.3 Determination of Hardness

The hardness of the molded samples was determined using a digital hardness tester (Elastoco, Sweden) according to the ISO 48: 2011(E) standards at 27 ± 2 °C. The tests were performed on unstrained samples of 38 mm diameter and 1 ± 0.2 mm thickness. Readings were taken after 10 seconds from the indentation at IRHD values.

2.4.4 Determination of Abrasion Volume Loss

Abrasion volume loss of the samples was measured using an abrasion tester (Hampden APH 40, England). Test specimens of 15 mm diameter and 8 ± 0.2 mm thickness were placed on a rotating holder and a 10 N load was applied. A pre run was given for conditioning the sample and the initial sample weight was taken. Weight after the test was also taken. The difference in weight is the weight loss of the test piece after its travel through 40 cm on a standard abrasive surface. All the tests were carried out at 27 ± 2 °C. Three test pieces were used and average value was calculated.

\[
\text{Abrasion volume loss} = \frac{(\text{Initial weight} - \text{Final weight})}{\text{Density}} \times 1000
\]

2.4.5 Determination of Resilience

The horizontal rebound method according to ISO 4662: 2009R was used to measure the rebound resilience of the samples. According to the method a plunger suspended from a given height above the specimen was released and the rebound height was measured. The resilience scale was marked in 100 equally spaced divisions and hence the rebound height was equal to the resilience percentage. Three test pieces were used and average value was calculated. The rebound resilience test was carried out at room temperature (27 ± 2 °C).

2.4.6 Determination of Compression Set

Compression set of the test samples were measured by applying a constant load on test pieces for continuous 72 hours using compression set testing apparatus. Three test pieces were used and average value was calculated. The compression test was carried out at room temperature (27 ± 2 °C). Obtained data were interpreted and analyzed using Microsoft Excel 10.0 and Origin 6.0 software.

2.5 Preparation of ball Milled Ceramic Powder Filler Added Natural Rubber Composites

In order to investigate the effect of the incorporation of reduced particle sized filler in to the natural rubber, it was prepared three different composites with the ball milled sub micrometer sized ceramic powder filler loading of 15, 25 and 50 phr. Other ingredients were added as the exactly same compositions in section 2.3. The mechanical properties were tested for these composites as mentioned in section 2.4.
3. Results and Discussion

3.1 Preparation of Ceramic Waste Powder

Fourier transform infra-red (FTIR) spectra of KBr pellets prepared from this initial ceramic powder (Fig. 1) were obtained using Alpha FTIR spectrophotometer (Bruker, USA).

![Fig. 1 FTIR spectra of the ceramic powder before modification](image1)

Fig. 1 FTIR spectra of the ceramic powder before modification

Fig. 1 shows the FTIR spectra of the initial ceramic waste powder and it is shown that major absorption bands are in good agreement with the reported values of SiO$_2$ (Table 3) [20].

<table>
<thead>
<tr>
<th>Table 3 FTIR absorption bands of initial ceramic powder with band assignments and reported bands of SiO$_2$ pristine sample [20]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bonding vibration modes</td>
</tr>
<tr>
<td>Si-O Asymmetric bending</td>
</tr>
<tr>
<td>Si-O-Si Symmetric stretching</td>
</tr>
<tr>
<td>Si-O-Si Asymmetric stretching</td>
</tr>
<tr>
<td>Combined vibrations of the SiO$_2$ network</td>
</tr>
<tr>
<td>OH stretching - due to surface OH</td>
</tr>
</tbody>
</table>

These FTIR results indicate that the composition of initial ceramic powder is much similar to the SiO$_2$ as SiO$_2$ is one of the major ingredients in the ceramic manufacturing process. Elemental analyses and composition of the phases of these initial ceramic particles were determined by X-ray fluorescence spectroscopy (Fig. 2, Table 3) and X-ray diffraction techniques (Fig. 3) respectively.

![Fig. 2 X-ray fluorescence spectroscopy of initial ceramic powder](image2)

Fig. 2 X-ray fluorescence spectroscopy of initial ceramic powder

<table>
<thead>
<tr>
<th>Table 4 Elemental data obtained form XRF for initial ceramic sample</th>
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</thead>
<tbody>
<tr>
<td>Element</td>
</tr>
<tr>
<td>Si</td>
</tr>
<tr>
<td>Al</td>
</tr>
<tr>
<td>Na</td>
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<tr>
<td>K</td>
</tr>
<tr>
<td>Fe</td>
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<tr>
<td>Ti</td>
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</table>

According to the XRF spectrum (Fig. 2) Si-Ka and Al-Ka lines represent Si and Al as the major constituents in the initial ceramic powder. According to the Table 4, Si present as SiO$_2$ is in the mass percentage of 60.3% and Al present as Al$_2$O$_3$ is in the mass percentage of 20.2%. Elements like Na, K, Fe and Ti are present as minor constituents in the initial ceramic powder.

Fig. 3 shows the X-ray diffractograms (XRD) of the initial ceramic sample. The peaks corresponding to the (111), (220), and (311) planes could indicate the presence of clusters of silicon. The result reveals that a high percentage of these particles are amorphous, but a few of them are crystalline.

The average particle size and the surface morphology of the initial ceramic powder were investigated using scanning electron microscopic (SEM) techniques (Fig. 4).

![Fig. 3 X-ray diffractograms of initial ceramic powder](image3)

Fig. 3 X-ray diffractograms of initial ceramic powder

![Fig. 4 Scanning electron microscope images of initial ceramic powder (a) under 10000x magnification (b) under 50000x magnification](image4)

Fig. 4 Scanning electron microscope images of initial ceramic powder (a) under 10000x magnification (b) under 50000x magnification

The SEM micrograph indicates the prepared initial ceramic powder is irregular in shape and size. The morphology is rough in all the samples. The particle sizes are between 125 μm and below.

The elemental composition present in the initial ceramic powder was further identified by the energy-dispersive X-ray spectroscopy (Fig. 5, Table 5) coupled with SEM.

![Fig. 5 Energy-dispersive X-ray spectrum of the initial ceramic powder](image5)

Fig. 5 Energy-dispersive X-ray spectrum of the initial ceramic powder

Energy-dispersive X-ray spectroscopy results suggest the presence of Si and oxygen elements, confirming the presence of SiO$_2$ particles. The SiO$_2$ compound percentage is found as much higher 77.15% which is according to the Table 5. Also Al also present in the form of Al$_2$O$_3$ and having the compound percentage of 18.47%. Elements like Al, Na, K, Fe and Ti are found as minor constituents.

<table>
<thead>
<tr>
<th>Table 5 Elemental composition of the initial ceramic powder obtained from the energy-dispersive x-ray spectroscopy</th>
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</thead>
<tbody>
<tr>
<td>Element</td>
</tr>
<tr>
<td>Si</td>
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<tr>
<td>O</td>
</tr>
<tr>
<td>Al</td>
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<tr>
<td>K</td>
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<tr>
<td>Na</td>
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<tr>
<td>Fe</td>
</tr>
<tr>
<td>Ti</td>
</tr>
</tbody>
</table>

3.2 Surface Modification of Ceramic Particles

The surface modification was carried out by Silane69 coupling agent. After the chemical procedure of the surface modification by Silane69 coupling agent, the modification was confirmed using Fourier-transform infrared spectroscopic results as shown in the Fig. 6.

![Fig. 6 FTIR spectra of initial ceramic powder, Silane69 and surface modified ceramic powder using silane](image6)

Fig. 6 FTIR spectra of initial ceramic powder, Silane69 and surface modified ceramic powder using silane
In the surface modification process a grafting reaction occurs between the hydroxyl groups from ceramic powder and the silanol groups formed by hydrolysis of the Silane69 coupling agent. Therefore, the grafting reaction between Silane69 and ceramic powder can be identified by the appearance of characteristic bands in FTIR spectra, such as characteristic band of \( \text{v}_{\text{O-H}} \) appearing at 2976-2886 cm\(^{-1}\). Fig. 6 shows FTIR spectra of ceramic powder, Silane69 grafted ceramic powder and Silane69. It can be seen that the \( \text{v}_{\text{O-H}} \) band exists in Silane69 grafted ceramic powder, which is the evidence that Silane69 molecules are grafted on the ceramic powder surface and the surface has been modified.

The surface modification caused by Silane69 coupling agent was further confirmed by X-ray fluorescence spectroscopy (Fig. 7, Table 6).

According to the XRF spectrum (Fig. 7) Si-Ka and Al-Ka lines represent Si and Al as the major constituents as in the initial ceramic powder (Fig. 2). Instead of them S-Ka line is present in the spectrum that indicates S as a major constituent in the surface modified ceramic powder. According to the Table 6, Si present as \( \text{SiO}_2 \) is in the mass percentage of 43.1%, Al present as \( \text{Al}_2\text{O}_3 \) is in the mass percentage of 14.8% and S present in the mass percentage of 9.18%. Na, K, Fe, Ti are present as minor constituents in the surface modified ceramic powder.

The surface modification caused by Silane69 coupling agent was further confirmed by thermo gravimetric analysis (TGA) (Fig. 8).

The surface modification of ceramic particles with the excess presence of C and S of modified ceramic particles looks smooth and sleek compare to the pristine initial ceramic particle morphology.

![Fig. 7 X-ray fluorescence spectroscopy of surface modified ceramic powder using Silane69](image)

The surface modification of ceramic particles with the excess presence of C and S of modified ceramic particles looks smooth and sleek compare to the pristine initial ceramic particle morphology.

![Fig. 8 Thermo gravimetric curve of surface modified ceramic powder using Silane69](image)

TGA of surface modified ceramic powder using Silane69 (Fig. 8) shows gradual weight loss at 246.44 °C (10.49%). This weight loss may be due to removal of free water absorbed from atmosphere and presence of any structural water. Gradual weight loss from 250 °C to 550 °C (1.36-4%) is due to dihydroxylation for structural OH. This confirms the surface modification of ceramic powder by Silane69.

The surface modified ceramic powder were subsequently analyzed by scanning electron microscopy (SEM). The SEM micrographs of surface modified ceramic powder (Fig. 9) confirmed our speculation. From SEM, pristine initial ceramic particles (Fig. 4) presented irregular shaped and rough morphology and small aggregates could be observed due to hydrogen bonds between the surface hydroxyl groups of silica particles. After Silane grafting, the surface modified particles show regular spherical structure. The modified particles have stickied to each other and formed large aggregations after silane modification. Also the surface morphology of modified ceramic particles looks smooth and sleek compare to the pristine initial ceramic particle morphology.

![Fig. 9 Scanning electron microscope images of surface modified ceramic powder (a) under 25000 × magnification (b) under 50000× magnification](image)

The Silane69 coupling agent which was used for the surface modification is composed of C, O, H, S and Si in its chemical structure (Fig. 10).

![Fig. 10 Chemical structure of Silane69 coupling agent](image)

Energy-dispersive x-ray spectroscopy results suggest the surface modification of ceramic particles with the excess presence of C and S elements, compare to the pristine initial ceramic powder (Table 5).

![Fig. 11 Energy-dispersive x-ray spectrum of the surface modified ceramic powder](image)

Energy-dispersive X-ray spectroscopy results suggest the surface modification of ceramic particles with the excess presence of C and S elements, compare to the pristine initial ceramic powder (Table 5).

The surface modification caused by Silane69 coupling agent was further confirmed by examining the elemental composition present in the surface modified ceramic powder using energy-dispersive x-ray spectroscopy (Fig. 11, Table 7) coupled with SEM.

![Fig. 12 Scanning electron microscope images of ball milled ceramic powder (a) under 25000 × magnification (b) under 50000× magnification](image)
3.3 Preparation of Sub Micrometer Sized Ceramic Waste Powder

Initially prepared ceramic waste powder was ball milled for 6 hours at an 800 rpm speed by a laboratory planetary ball mill followed by dry milling. The average particle size and the surface morphology was characterized by scanning electron micrographs (Fig. 12).

The SEM micrograph indicates the ball milled ceramic powder is irregular in shape and size. The particle sizes are between few nm to few hundred nanometers. The particle size has become smaller compared to the pristine initial ceramic powder sample with the effect of ball milling. Particle aggregations also can be observed up to some extent.

X-ray diffraction pattern shows the crystalline structure the ball milled ceramic powder (Fig. 13). Typical silica characteristic is observed with an intense sharp peak centered at $2\theta = 26^\circ$. Reduction in crystallite size is observed with the effect of ball milling.

![Fig. 13 X-ray diffractograms of ball milled ceramic powder](image)

3.4 Mechanical Property Comparison of Surface Modified and Non-Modified Ceramic Powder Filler Added Natural Rubber Composites

3.4.1 Tensile Strength

Tensile strength of ceramic filler containing composites is higher than that of the pristine rubber composite (control) up to 50 phr filler loading. At higher filler loadings (75 phr), tensile strength is lower than that of the control and could be attributed to agglomeration of filler particles. Surface modified ceramic filler loaded composites have higher tensile strength than non-modified ceramic filler loaded composites at some of the filler loadings (Fig. 14).

![Fig. 14 Tensile strength comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)

3.4.2 Modulus at 100% Elongation

Modulus at 100% elongation is related to hardness or stiffness of the composites. As in the case of hardness, modulus at 100% elongation of ceramic filler loaded composites is higher than that of the control. However, no significant difference is observed between the modulus at 100% elongation of surface modified ceramic filler loaded composites and non-modified ceramic filler loaded composites (Fig. 15).

![Fig. 15 Modulus at 100% elongation comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)

3.4.3 Elongation at Break

As expected, elongation of the control is higher than that of the ceramic filler containing composites. Elongation at break of the surface modified ceramic filler loaded composites is higher than that of non-modified ceramic filler loaded composites at filler loadings 50 phr and above (Fig. 16).

![Fig. 16 Elongation at break comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)

3.4.4 Tear Strength

Tear strength of ceramic filler loaded composites is lower than that of the control. As in the case of elongation at break, tear strength of surface modified ceramic filler loaded composites is higher than that of non-modified ceramic filler loaded composites at filler loadings 50 phr and above (Fig. 17).

![Fig. 17 Tear strength comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)

3.4.5 Hardness

As expected, hardness increases with the increase of ceramic filler loading. Surface modified ceramic filler containing composites have lower hardness than non-modified ceramic filler containing composites at 25 phr loading and above (Fig. 18).

![Fig. 18 Hardness comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)

![Fig. 19 Abrasion volume loss comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)
3.4.6 Abrasion Volume Loss

Abrasion volume loss of the composites varies with the ceramic filler loading according to a cyclic pattern. At most of the ceramic filler loadings, surface modification has resulted in a decrease in the abrasion volume loss (Fig. 19).

3.4.7 Resilience

Resilience of ceramic filler loaded composites is lower than that of the control except at 5 phr and 10 phr non-modified filler added composites. Resilience gets decrease with the increasing amount of filler added. Resilience of non-modified filler added composites is larger than modified filler added composites except at 50 phr filler loading (Fig. 20).

![Fig. 20 Resilience (%) comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)

3.4.8 Compression Set

Compression set of the composites decrease with the ceramic filler loadings compared to the control. Surface modified composites have little lower compression set compared to non-modified composites. Compression set is increasing after 15 phr filler loadings and this may cause due to agglomerations (Fig. 21).

![Fig. 21 Compression set (%) comparison of surface modified and non-modified ceramic powder filler added natural rubber composites](image)

Due to the addition of fillers, molecular chain mobility is decreasing with time. Up to some extend ceramic fills the void spaces between natural rubber. So that hardness, compression resistance, 100%modulus, tensile strength like properties gets increased. With the higher filler adding insufficient sites of natural rubber is increasing. So that above properties get decreased (Fig. 22).

![Fig. 22 Explanation of obtained mechanical properties](image)

3.5 Mechanical Properties of Ball Milled Ceramic Powder Filler Added Natural Rubber Composites

In this experiment ceramic filler added natural rubber composites were made in 0, 5, 10, 15, 25, 50 and 75 phr compositions. It was found to have these composites show the optimal properties at moderate compositions. In order to investigate the effect of the incorporation of reduced particle sized filler in to the natural rubber, it was prepared three different composites with the ball milled sub micrometer sized ceramic powder filler loading of 15, 25 and 50 phr. When it compares the modulus at 100% elongation of the ball milled sub micrometer sized ceramic filler added composites with the initial non-ball milled ceramic filler added composites, it was found to have much improved modulus can be achieved by reduced particle size filler loadings due to the dense packing. Optimal mechanical properties can achieve by adding less loading of ball milled ceramic particles compared to non-ball milled ceramic particles (Fig. 23). This may due to the ability of ball milled ceramic particles that they tend to fill the void spaces of natural rubber matrix more than that of the non-ball milled ceramic particles.

![Fig. 23 Modulus at 100% elongation comparison of ball milled and non-ball milled ceramic powder filler added natural rubber composites](image)

4. Conclusion

Ceramic waste are the kind of major wastes in manufacturing sectors. The aim of this investigation was the utilization of these ceramic waste as a low cost filler material in the manufacture of natural rubber based composites. Mechanical properties of ceramic waste-based natural rubber composites have been investigated as a function of fiber loading, ratio and treatment. The surface modification was it found that the incorporation of ceramic waste filler enhanced the mechanical properties of the natural rubber. As the filler concentration increases, the modulus at the 100% elongation and tensile strength slightly increased, but the elongation at break decreased may due to reinforcing effects. The hardness was also found to increase with the loading level of ceramic waste in the composites. This is due to the increased material stiffness. These composites also found to have better compression resistant compared to the pristine natural rubber. The filler surface modification with Silane69 coupling agent further improved the mechanical properties of composites. Furthermore, optimal mechanical properties can achieve by adding less loading of ball milled ceramic particles compared to non-ball milled ceramic particles. Overall, the mechanical performance of the ceramic waste based natural rubber composites was better than that of the control. It was observed that the optimal mechanical properties of the composites are present at moderate loading of fillers. Pristine natural rubber compound shows low mechanical properties such as; low hardness, low modulus at 100% elongation and low tensile strength. Compared to that these novel ceramic waste based natural rubber composites shows improved mechanical properties such like compression resistant, high hardness, high modulus and high tensile strengths. Finally, it can be concluded that, within the limited scope of the experiments carried out in this investigation, these ceramic waste based natural rubber composites can be utilized as a proper flooring material.

Acknowledgments

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