

DEVELOPMENT AND CHARACTERIZATION OF HYBRID ALUMINUM MATRIX COMPOSITES REINFORCED WITH SILICON CARBIDE AND CARBONIZED COCONUT SHELL FOR HIGH STRENGTH/LIGHTWEIGHT APPLICATIONS

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Abstract: The production of hybrid aluminum matrix composites (HAMCs) using low-cost agro-waste materials as reinforcements is gaining popularity due to their excellent properties, wide industrial applications, and cost-effectiveness. In this study, a hybrid AA6061 matrix composite was synthesized using silicon carbide (SiCp) and carbonized coconut shell (CCSP) particles as reinforcements in a 1:1 ratio through stir casting. The effects of SiCp and CCSP particles of varying sizes and weight fractions on density, hardness, and strength properties were studied. The results indicated that the addition of CCSP and SiCp particles to the alloy decreased its density while enhancing the hardness and tensile strength properties. Mathematical models were developed to predict the hardness and tensile strength of the nano-composite and validated by physical experiments. The microstructure analysis revealed uniform distribution of reinforcing particles in the base matrix and excellent bonding between the base matrix and reinforcements after casting. The hybrid composite can be used in high strength/lightweight applications in the automobile and aerospace industries.

Keywords: Hybrid aluminum matrix composites, silicon carbide, carbonized coconut shell, stir casting, mechanical properties.

INTRODUCTION

The application of aluminium matrix composites (AMC) in industrial practice in recent times has grown tremendously because of their good combination of high strength, low density, durability, machinability, availability, and its comparatively low materials cost. Thus, they are very attractive for lightweight applications such as automobile, aerospace, military and transport sectors due to high specific strengths, good fatigue properties and wear resistance (Wang *et al.*, 2014; Karl, 2006). The scope of these properties can be extended by singly or multiply reinforced aluminium matrix with hard and light micro or nano particles leading to the formation of hybrid aluminium matrix composites (HAMCs), (Parswajinan *et al.*, 2018; Sudipt and Ananda, 2008; Michael *et al.*, 2015). The property of the AMCs depends on the property of the matrix and the reinforcement (Vijaya *et al.*, 2014). The reinforcement in AMCs may be in the structure of continuous



fibres, discontinuous fibres, whiskers or particulates. Widely used reinforcement ceramic materials in AMCs include silicon carbides (SiC), titanium borides (TiB₂), alumina (Al₂O₃), nitrides, boron and graphite.

Recently, reinforcements commonly used in aluminium matrix composites have been extended to include agrowastes such as rice waste ash, sugar cane bagasse, palm kernel ash, periwinkle shell ash and coconut shell ash (Senthilkumar *et al.*, 2016; Iyasele, 2018). These agro-wastes are cheap and readily available in high quantity. Ajibola and Fakeye (2015) reported the production and characterization of Zinc-Aluminium composite reinforced with silicon carbide and palm kernel shell ash (PKSA), in which samples prepared with compositions 5wt% SiC added with 0.2%, 0.4%, 0.6%, 0.8% and 1.0wt% PKSA were utilized to prepare the reinforcing phase with Zinc-Aluminium matrix composite using two step stir casting method. The agro waste material was preheated to 650°C before being introduced into the Zinc-Aluminium composite in molten state. Hardness and tensile test were used to characterize the composite produced. The result shows that increase in PKSA reinforcement increases the hardness and tensile strength of the composite, and the microstructure shows that SiC and PKSA were well dispersed in the alloy matrix.

Hima *et al.*, (2018) used aloe vera powder to reinforce the aluminum metal matrix in order to study the mechanical characteristics of the AMC. He compared the suitability of fly ash (reported by several researchers) and aloe vera as reinforcement particulates for the aluminum metal matrix, but his study reported that the use of aloe vera significantly reflected an improvement on the mechanical properties of the AMC with regards to tensile strength, impact strength, and hardness, compared to the popularly known and used fly ash. Nwobi-Okoye and Ochieze (2018) experimentally studied the behavior of aluminum alloy A356 reinforced with a particulate composite of cow horn, for application in a brake drum via the use of an artificial neural network (ANN), response surface methodology (RSM), and simulated annealing. The study was focused on modeling the age hardening process on the developed or reinforced AMMC using ANN and RSM. The result revealed that the modeling via ANN that generated the data for age hardening was improved by 0.9583 predictions compared with that of RSM of 0.9921 predictions of ANN.

Umunakwe *et al.*, (2017) investigated the density, porosity, microstructure and some mechanical properties of particulate periwinkle shell-aluminium 6063 metal matrix composite (PPS-AIMMC) produced by two-step casting and compared the properties of the composites with those of the aluminium 6063 (Al60603) alloy. Periwinkle shells were milled to particle sizes of 75µm and 150µm and used to produce PPS-AIMMC at 1, 5, 10 and 15wt% filler loadings using two-step casting technique. The addition of PPS to aluminium alloy reduced the density of the composite. It was observed that the filler distributes uniformly in the matrix due to the two-step casting technique. The porosities of the composites were within acceptable level of 0-5% except for the composite with 15wt% PPS of 150µm particle size. Improved strength, ductility, hardness and modulus were obtained when the filler was used to reinforce the alloy. However, using a filler of higher particle size resulted to higher porosity, reduced tensile strength, ductility and toughness.

In this work, attempt has been made to prepare and characterize hybrid AA6061 alloy composites by adding SiC and carbonized coconut shell reinforcement particles into the matrix using stir casting method

MATERIALS AND METHODS

The various materials used for this study includes aluminium alloy 6061, silicon carbide and coconut shell. Table 1 shows the chemical composition of Al6061 Alloy. The coconut shells were washed, dried and carbonized at 600°C for 5 hours. Both the carbonized coconut shell and silicon carbide particulates were ball milled for 60 hours and 50 hours respectively in accordance with the work of Bello (2017). After milling, the reinforcement particulates were classified for particle size. Samples of the AA6061/SiC/CCS composites were then fabricated by stir casting method at 700°C using a crucible (Figure 1e), with addition of 2wt% Magnesium to enhance the wettability between reinforcements and molten metal. The molten metal was stirred using a stainless steel stirrer and measured quantity of reinforcement (0, 3, 6, 9, 12 and 15%wt of equal mixtures of CCS and SiC respectively with different particle sizes) was added. The mixture was stirred for 5 minutes for uniform distribution. Then molten metal was poured in a sand mould and allowed to solidify.

Table 1: Composition of AA6061

Element	Si	Fe	Cu	Mn	Mg	Ni	Zn	Ti	Cr	Al
Weight (%)	0.514	0.230	0.161	0.071	0.960	0.010	0.015	0.031	0.103	Bal.



Figure 1: (a) Coconut shells (b) Silicon carbide (c) Cast AA6061 ingot (d) Crushed carbonized coconut shells (e) Carbonized coconut particulates (f) Stir casting furnace

Density Measurements

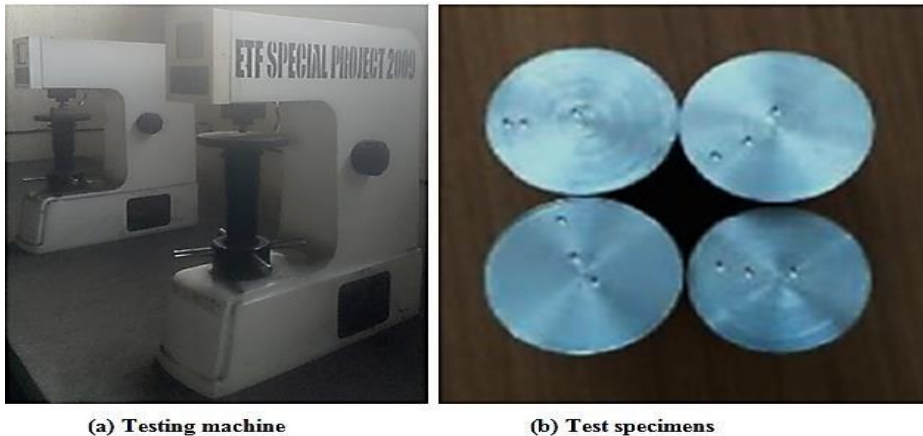
Density measurements were carried out on the developed composite samples using Archimedes' principle. The samples were first weighed in air and then weighed when fully immersed in water. The difference in mass of each sample was recorded and used to calculate the density of the sample. The experimental density of composite was thus calculated using Equation 1.

$$\rho_{cexp} = \frac{W_{air}}{W_{air} - W_{water}} \times \rho_{water} \quad (1)$$

where, W_{air} is the weight of composite in air, W_{water} the weight of composite in water and ρ_{water} the density of composite in water

Hardness Testing

The hardness of the aluminium alloy and composites were determined with Vicker's hardness tester (Figure 2a). The dimension of each specimen for hardness testing was diameter 30mm \times 15mm thick (Figure 1b) and each specimen was grinded and polished to obtain a flat smooth surface. During the testing, a load of 980.7N was applied for 10s on the specimen through square based pyramid indenter and the hardness reading taken in a standard manner. The readings were taken in two different points at the surface of the hardness specimen and the average computed as the hardness of the specimen.



(a) Testing machine

(b) Test specimens

Figure 2: Hardness test

Tensile testing

The tensile test was carried out on the composite samples to determine its strength and ductility, yield strength and other tensile properties. In the present investigation all the tensile tests were conducted using Monsanto Tensometer {type W, serial 11148} tensile test machine in the Department of Metallurgical and Materials Engineering, Federal University of Technology, Owerri. The standard tensile test sample of a dumbbell shape with a diameter of 6.5 mm and gauge length 28 mm was used for the research. An initial load of 5kN was applied on the samples, with the load increased until failure occurred. The specimen was axially loaded in tension; the distance between the gauge marks was monitored. The specimen was elongated by the moving crosshead, load cell and extensometer measured the magnitude of the load and the elongation. All tests were conducted at room temperature.

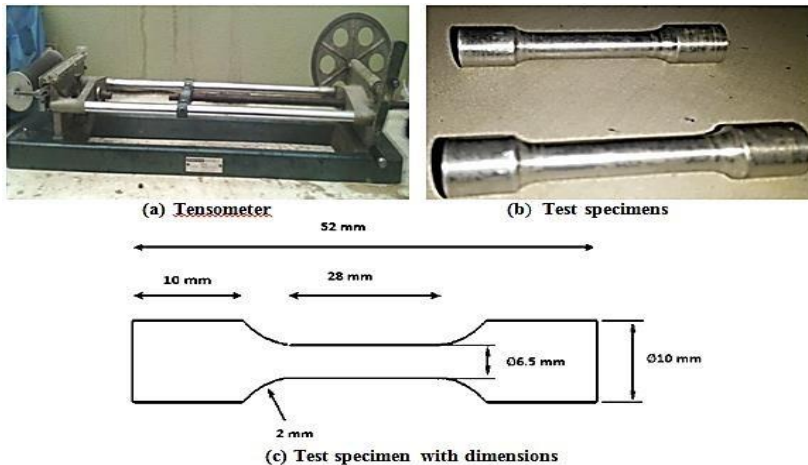


Figure 3: Tensile test set up

Modeling Tensile strength

In the current study the aluminum matrix is assumed to be an elastic-plastic material and the inclusions are approximated as being made of spherical and linear-elastic silicon carbide and carbonized coconut shell particulates. The interface between the matrix and the inclusions assumed to be ideal and particle cracking is not considered. Analytical model established by Punkanzky *et al.*, (1988) gives an empirical relationship between strength of composite (σ_c) and strength of matrix (σ_m) as given in Equation 2.

$$\sigma_c = \sigma_m \frac{1-V_p}{1+2.5V_p} e^{BV_p} \quad [(\quad)]$$

(2)

where, an empirical constant B is dependent on particle surface area, interfacial bonding strength, and particle density. The B value varies in between 3.49 to 3.87. The Punkanzky empirical relationship as given in Equation 1, gives the ultimate strength of the particulate metal matrix composite which depended on the strong particlematrix interfacial bonding. Also, considering adhesion, precipitate formation, particle size, agglomeration, voids/porosity, dislocation obstacles and the particle/matrix interfacial reaction, modified Voigt model $\bar{\sigma}_c = V_m \bar{\sigma}_m + \sum V_{rj} \bar{\sigma}_{rj} + \varepsilon_f$ considering these factors may be expressed as;

$$\bar{\varepsilon}_c = V_m \bar{\varepsilon}_m + \sum V_{rj} \bar{\varepsilon}_{rj} + \varepsilon_f \quad \{j = 1, 2, \dots, n\} \quad (3)$$

where $\bar{\sigma}_c$, $\bar{\sigma}_m$ and $\bar{\sigma}_{rj}$ are the composite, matrix and reinforcement strength respectively and $\bar{\varepsilon}_c$, $\bar{\varepsilon}_m$ and $\bar{\varepsilon}_{rj}$ are the corresponding strains. The volume fractions of matrix material and reinforcement are respectively, V_m and V_{rj} . The term ε_f , is a correction factor accounting for the effect of reinforcement particle size, porosity and interfacial interaction, and was determined from experimental data of the composite. In this study, the matrix material is reinforced with two different types of particulates, thus Equation 3 is rewritten as

$$\left. \begin{aligned} \bar{\sigma}_c &= V_m \bar{\sigma}_m + V_{r1} \bar{\sigma}_{r1} + V_{r2} \bar{\sigma}_{r2} + \varepsilon_f \\ \bar{\varepsilon}_c &= V_m \bar{\varepsilon}_m + V_{r1} \bar{\varepsilon}_{r1} + V_{r2} \bar{\varepsilon}_{r2} + \varepsilon_f \end{aligned} \right\} \quad (4)$$

$$\text{But } V_m + V_{r1} + V_{r2} = 1 \Rightarrow V_m = 1 - (V_{r1} + V_{r2})$$

where, V_m , V_{r1} and V_{r2} are the volume fractions of the matrix and the two different particulates 1 and 2 respectively.

The modified load transfer effect according to Sanaty-Zadeh (2012) is given as;

$$\Delta\sigma_{LT} = \frac{1}{2} V_p \sigma_m \quad (5)$$

and the Hall-Petch strengthening equation as modified by Hull and Bacon (2001) is;

$$\Delta\sigma_{H-P} = \frac{k_y}{\sqrt{d}} \quad (6)$$

√

Combining Equations 4 and 5, and neglecting the volume fractions of voids/porosity gives Equation 7.

$$k \quad \sigma_c = \frac{1}{2} V_p \sigma_m + \frac{y}{\sqrt{d_p}} \quad (7)$$

For matrix material reinforced with n different particulates, Equation 7 becomes,

$$\sigma_c = \frac{1}{2} (\sum V_{pi}) \sigma_m + \frac{k_y}{\sqrt{(\sum d_{pi})}} \quad \{i = 1, 2, n\} \quad (8)$$

For matrix material reinforced with two different particulates, Equation 8 becomes

$$\sigma_c = \frac{1}{2} (V_{p1} + V_{p2}) \sigma_m + \frac{k_y}{\sqrt{(d_{p1} + d_{p2})}} \quad (9)$$

Table 2 gives the material data for models prediction.

Table 2: Material and model parameters

Material	Elastic modulus (GPa)	Tensile strength (MPa)	Density (g/cm ³)	Poisson ratio
Silicon carbide particles	410	336	3.21	0.14
Carbonized coconut shell particles	512	18.03	1.68	0.92
AA6061	69	162.6	2.72	0.33

RESULTS AND DISCUSSION Chemical composition of Reinforcements

The results obtained from the XRF chemical compositional analysis of the carbonized coconut shell particles (CCSp) and SiC shown in Tables 3, revealed that the carbonized coconut particulates contained SiO₂, MgO, Al₂O₃ and Fe₂O₃ in great amount with small amounts of CaO, K₂O, Na₂O, MnO and ZnO. The presence of hard substances like SiO₂, Al₂O₃ and Fe₂O₃ suggest that the carbonized coconut shell particles can be used as particulate reinforcement in various metal matrix composites (Apasi *et al.*, 2016; Madakson *et al.*, 2012). This result is consistent with the results by Donald *et al.*, (2018), Apasi *et al.*, (2016) and Madakson *et al.*,

(2012), and the chemical composition has similarity with the XRF analysis of periwinkle shell ash, rice husk ash, fly ash and bagasse ash currently used in metal matrix composite as reported by Nwabufoh (2015), Hassan and Aigbodion (2010) and Rajan *et al.*, (2007). Therefore, the present work suggests the suitability of carbonized coconut shell as particulates reinforcement in metal matrix composites.

Table 3: Chemical composition of Reinforcements

CCS									
Element	Al ₂ O ₃	CaO	Fe ₂ O ₃	K ₂ O	MgO	Na ₂ O	SiO ₂	MnO	ZnO
%	16.50	0.58	14.50	0.51	16.51	0.50	46.70	0.42	0.35
SiC									
Element	SiC	Si	SiO ₂	Fe	Al	C			
%	98.6	0.30	0.6	0.09	0.10	0.3			

Density of composites

The results of the experimental density of the alloy and composites reinforced with micro and nano-particulates of SiC and CCS are shown in Table 4. Addition of reinforcement particles to the matrix alloy has significant effect on the composite density.

Table 4: Densities of composites

%wt. of reinforcement	Density (g/cm ³)								
	AA6061 with								
	μSiC	μCCS	n _f SiC	n _s SiC	n _f CCS	n _s CCS	μSiC/μCCS	n _f SiC/n _f CCS	n _s SiC/n _s CCS
0	2.72	2.72	2.72	2.72	2.72	2.72	2.72	2.72	2.72
3	2.75	2.67	2.74	2.75	2.56	2.60	2.70	2.65	2.67
6	2.80	2.54	2.77	2.79	2.47	2.50	2.68	2.63	2.65
9	2.84	2.50	2.80	2.82	2.43	2.47	2.65	2.61	2.62
12	2.87	2.42	2.83	2.85	2.30	2.38	2.64	2.57	2.60
15	2.91	2.37	2.86	2.90	2.09	2.21	2.62	2.49	2.53
n _f SiC: First nano SiCp (42.3nm) n _s SiC: Second nano SiCp (55.8nm) μSiC: SiC micro-particles (38μm) n _f CCS: First nano CCSp (50.01nm) n _s CCS: Second nano CCSp (60.3nm) μCCS: CCS micro-particles (63μm)									

Table 4 indicates that addition of micro and nano particles of SiC increased the composite density beyond that of the matrix alloy, whereas, the composite density decreased upon addition of micro and nano particulates of coconut shell. The observed increase in composite density may be caused by the higher density of SiC (3.21g/cm³) as compared with that of the base alloy. The decrease in density associated with the use of CCS particles is caused by the lower density of CCS (1.68g/cm³) compared to the density of aluminium matrix which is 2.72 g/cm³, oxidation of the base alloy, gas evolution and porosity in the composites. These results are consistent with earlier work of Prasad (2006) and Abba-Aji (2021). Table 4 also indicates that the

density of nano-composites decreased with both increasing weight fraction and decreasing particle size. Notably, the use of CCS nanoparticles further decreased the composite density. Peter *et al.* (2020) reported similar observation with the use of CCS in spark plasma sintered Ti-Ni based metal matrix composite. The density of the developed

AA6061/ n_f CCS and AA6061/ n_s CCS composites reduced by 23.2% and 18.75% on addition of 15wt.% each of n_f CCS and n_s CCS respectively, whereas that of the AA6061/ n SiC composites increased by 5.2% and 6.62% on addition of 15wt.% each of n_f SiC and n_s SiC, respectively. Note that two different categories of nanocomposites were presented for each reinforcement; first SiC nano-composites (AA6061/ n_f SiC), second SiC nano-composites (AA6061/ n_s SiC), first CCS nano-composites (AA6061/ n_f CCS) and second CCS (AA6061/ n_s CCS). The trend of experimental density values obtained therefore indicates that the first CCS nanocomposites (AA6061/ n_f CCS) acquired lowest densities. For the hybrid composite, densities were significantly reduced below that of the matrix alloy upon addition of combined micro and nano particulates of SiC and CCS. The composite density decreased by 0.74%, 2.6% and 1.84% at 3wt.% addition of μ SiC/ μ CCS, n_f SiC/ n_f CCS and n_s SiC/ n_s CCS, respectively. Figure 4 shows the comparative density plots of the different composites produced. It indicates that maximum and minimum densities were achieved with composites singly reinforced with μ SiC and nCCS respectively. The observed order of decrease in density were with; μ SiC/ μ CCS, n SiC/nCCS, μ CCS and nCCS, respectively.

The composite densities also increased in the order, μ SiC and n SiC. Hence it is evident that the CCS particulates reduced the composite density whereas the SiC particulates increased the composite density. Similar trend were reported by Donald *et al.*, (2018) and Padmavathi and Ramakrishnan (2019). This result trend therefore indicates that the hybrid AA6061/ n SiC/nCCS composites will be better suited for light weight applications. Note that n SiC, nCCS and n SiC/nCCS in Figure 4 represent average response for n_f SiC and n_s SiC; n_f CCS and n_s CCS; n_f SiC/ n_f CCS and n_s SiC/ n_s CCS respectively.

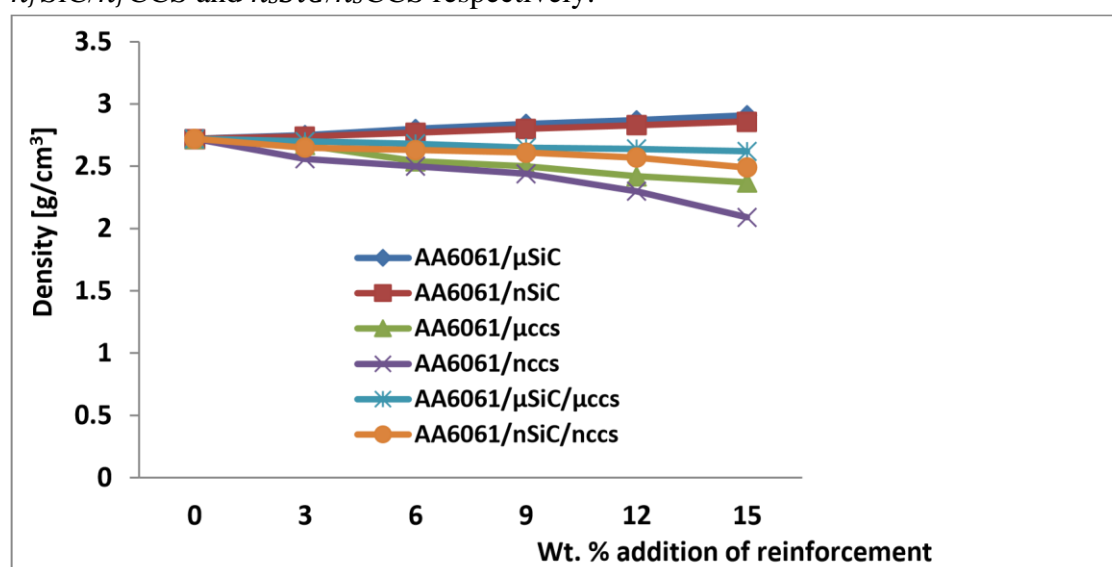


Figure 4: Comparative plots of the aluminium alloy 6061 composite densities Hardness of Composites

The average hardness values of the matrix alloy and composites reinforced with different weight percent and particle sizes of SiC and CCS are presented in Table 5. The composite hardness values were observed to increase with increasing weight percent and decreasing particle size of reinforcements.

Table 5: Average hardness of composites

%wt. of reinforce ment	Hardness (HV)								
	AA6061 with								
	μ SiC	μ CCS	n_f SiC	n_s SiC	n_f CCS	n_s CCS	μ SiC/ μ CCS	n_f SiC/ n_f CCS	n_s SiC/ n_s CCS
0	78.8	78.8	78.8	78.8	78.8	78.8	78.8	78.8	78.8
3	80.0	79.2	83.0	81.6	80.0	79.1	81.0	85.3	83.5
6	82.0	80.8	84.5	83.5	83.1	82.0	84.1	87.0	85.0
9	84.2	83.1	85.7	85.0	84.0	83.2	85.8	88.6	86.9
12	86.0	84.0	88.0	86.2	85.6	85.0	88.6	92.2	89.4
15	87.0	84.8	90.0	87.4	86.3	85.7	90.0	95.0	92.0
n_f SiC: First nano SiCp (42.3nm) n_s SiC: Second nano SiCp (55.8nm) μ SiC: SiC micro-particles (38 μ m) n_f CCS: First nano CCSp (50.01nm) n_s CCS: Second nano CCSp (60.3nm) μ CCS: CCS micro-particles (63 μ m)									

The composite hardness value increased by 2.79%, 8.25% and 5.97% with 3wt.% addition of μ SiC/ μ CCS, n_f SiC/ n_f CCS and n_s SiC/ n_s CCS, respectively. The composite hardness further increased by 14.21%, 20.55% and 16.75% with 15wt.% addition of μ SiC/ μ CCS, n_f SiC/ n_f CCS and n_s SiC/ n_s CCS, respectively. Maximum hardness values were achieved with addition of n_f SiC particulates. The observed increase in hardness value is attributed to the matrix phase refinement and the integration of hard and brittle phases of micro and nano-particles of silicon carbide (μ SiC and n SiC), SiO₂, Al₂O₃ and Fe₂O₃ (obtained from carbonized coconut shell particulates) in the matrix material. Similar trend has been reported in literature (Ajibola and Fakeye, 2015; Hima *et al.*, 2018; Bello, 2017). Umunakwe *et al.*, (2017) also reported the increase in hardness of composite reinforced with 75 μ m particulate size over those reinforced with and 150 μ m particulate size. It is believed that the decrease in particle size enhanced interfacial excellent bond, increased dislocation density and thus improved hardness and strength. Notably, SiC reinforced composites acquired superior hardness compared with those reinforced with CCS. This is because SiC has higher hardness value than CCS. Compared with the hybrid micro composites (AA6061/ μ SiC/ μ CCS), the hybrid nano composites (AA6061/ n SiC/ n CCS) possessed superior hardness. This is caused by further particle size reduction and higher bond strength between the nanoparticles and the matrix alloy.

Modeling of Hardness of Hybrid AA6061/SiC/CCS nano composites

The effects of reinforcement weight percent (Wt) on the hardness of AA6061/ n SiC/ n CCS composites was appraised using a linear model developed for theoretical prediction of the hardness (response) with percent weight, W_t as predictor. Therefore, this model involves 5 different levels and two predictors (5¹) equivalent

to 5 experimental runs using balanced two – factor factorial design of experiment. Table 6 reveals the experimental design, actual, predicted and residual hardness. Functions derived for the model revealed in Equations 10 and 11. The adequacy and validity of the developed model was tested through ANOVA as shown in Tables 7 and 8 The model has a p – value of 0.000 which is < 0.05 (i.e. $\alpha = 0.05$ or 95% confidence) and F – value of 241.938 indicating that the model is considered to be statistically significant. The significant model terms are considered to generate the model and are identified with p values < 0.05 . However, model terms with p value > 0.05 are regarded as insignificant. The difference between the predicted $R^2(0.984)$ and adjusted $R^2(0.980)$ is within the acceptable limit < 0.2 in order to establish a reasonable agreement.

$$\text{Hardness} = \beta_0 + \beta_1(\text{Wt})^n \quad (10)$$

with $\beta_0 = 77.9$, $\beta_1 = 178.67$ and $n = 1$, the model equation becomes

$$\text{Hardness} = 77.9 + 178.67(\text{Wt}\% \text{ nSiC/nCCSp}) \quad (11)$$

Table 6: Comparison of measured and modeled hardness of hybrid nano composites

Order	Predictor Wt.% (nSiC+nCCS)	Response Hardness (Hv)		
		Measured	Model	Residual
1	3	89.2	88.62	-0.58
2	6	93.0	99.34	6.34
3	9	117.3	110.06	-7.24
4	12	123.0	120.78	-2.22
5	15	127.8	131.50	3.70

Table 7: Test for significance of nSiC/nCCS composites Hardness

Order	Predictor	Unstandardized β	Standard error	Standardized β	t	Significance	Tolerance	VIF
1	Constant	77.90	1.719		44.669	0.000		
2	Wt %	178.67	0.189	0.992	15.554	0.000	1.000	1.000

CCSp/SiC

Dependent Variable: Hardness VIF: Variance inflation factor

Table 8: Test result of ANOVA for nSiC/nCCS composites Hardness

Source	Sum squares	of Degree freedom (df)	of Mean square	P value	(F-value)	Predicted R^2	Adjusted R^2
Model		1		0.000		0.984	0.980
	Regression	1364.897	1	0.000	241.938		
			4	5.642			
Residual			22.566				

Comparative plots of measured and model predicted composite hardness values are given in Figure 5. It indicates deviation of model predictions from measured data, with minimum and maximum deviations of 0.65% at 3wt. % addition of nSiC/nCCS and 6.17% at 9wt. % addition of nSiC/nCCS respectively. The observed deviations of model predicted values from experimental results are believed to be caused by non-inclusion of effect of other process parameters such as mould/melt interactions, agglomeration, porosity, stirring speed, pouring temperature in model development.

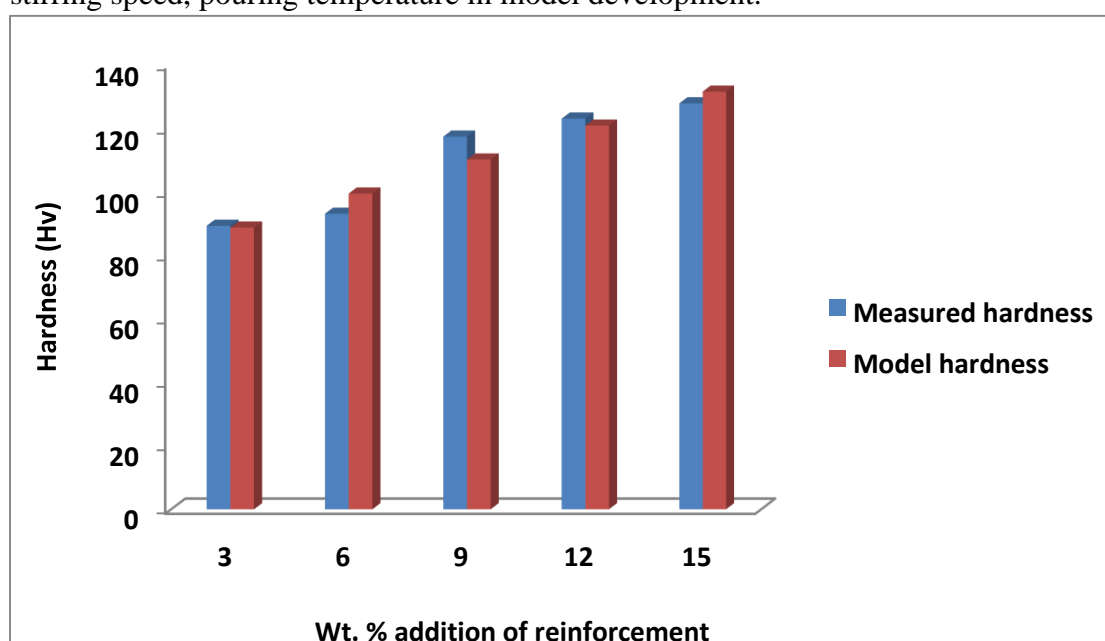


Figure 5: Hardness values of hybrid nano-composite

Tensile and Yield strength of Hybrid AA6061/SiC/CCS Composites

Figure 6 depicts progressive improvement in ultimate tensile and yield strength values for hybrid micro and nano composites. The ultimate tensile and yield strengths of the composites were observed to continuously improve as the percentage of the particles was increased. This improvement continues with decreasing reinforcement particulate size. This is evident as the hybrid nano-composites acquired superior ultimate tensile and yield strengths than the hybrid micro-composites. Similar trend has been reported in literature (Prasad and Krishna, 2010; Abba-Aji, 2021; Michael, 2022). Donald *et al.*, (2018) and Padmavathi and Ramakrishnan (2019) also reported improved tensile strength properties of AMC composite. The observed increase in strength value is attributed to the matrix phase refinement and the integration of hard and brittle phases of micro and nano- particles of silicon carbide (μ SiC and nSiC), SiO_2 , Al_2O_3 and Fe_2O_3 (obtained from micro and nano- particles of carbonized coconut shell; μ CCS and nCCS) in the matrix material. The hard and brittle phases constitute barriers to dislocation movement across the grain boundaries. This in effect strengthens the soft matrix and improves the composites hardness. Another factor that may contribute to the better interaction between the particles and the matrix is the clean and clear interface between them,

conferring the composite an increased load-carrying capacity. Composites with 15wt.% nSiC/nCCS and 15wt.% μ SiC/ μ /CCS presented maximum improvement in ultimate tensile strength value (53.4% and 41.5% respectively, for nSiC/nCCS and μ SiC/ μ /CCS) compared to the base alloy. Another possible reason for the improved ultimate tensile strength value could be the interaction between the particles and dislocations when the composites are under load; could also be due to the existence of several appending dislocations around the particles owing to the variation in the thermal expansion coefficient between the particles and the base matrix. The mismatch in thermal expansion coefficient between the matrix and the particles will also strengthen dislocation at the grain boundary based on the level of mismatch. The increase in yield strength was due to the decreased inter-particle spacing between the SiC and CCS particles as the particles are harder than the base alloy. Also, the addition of micro and nano particulates of SiC and CCS to the base alloy suppressed stress deformation, thereby improving the yield strength of the composites.

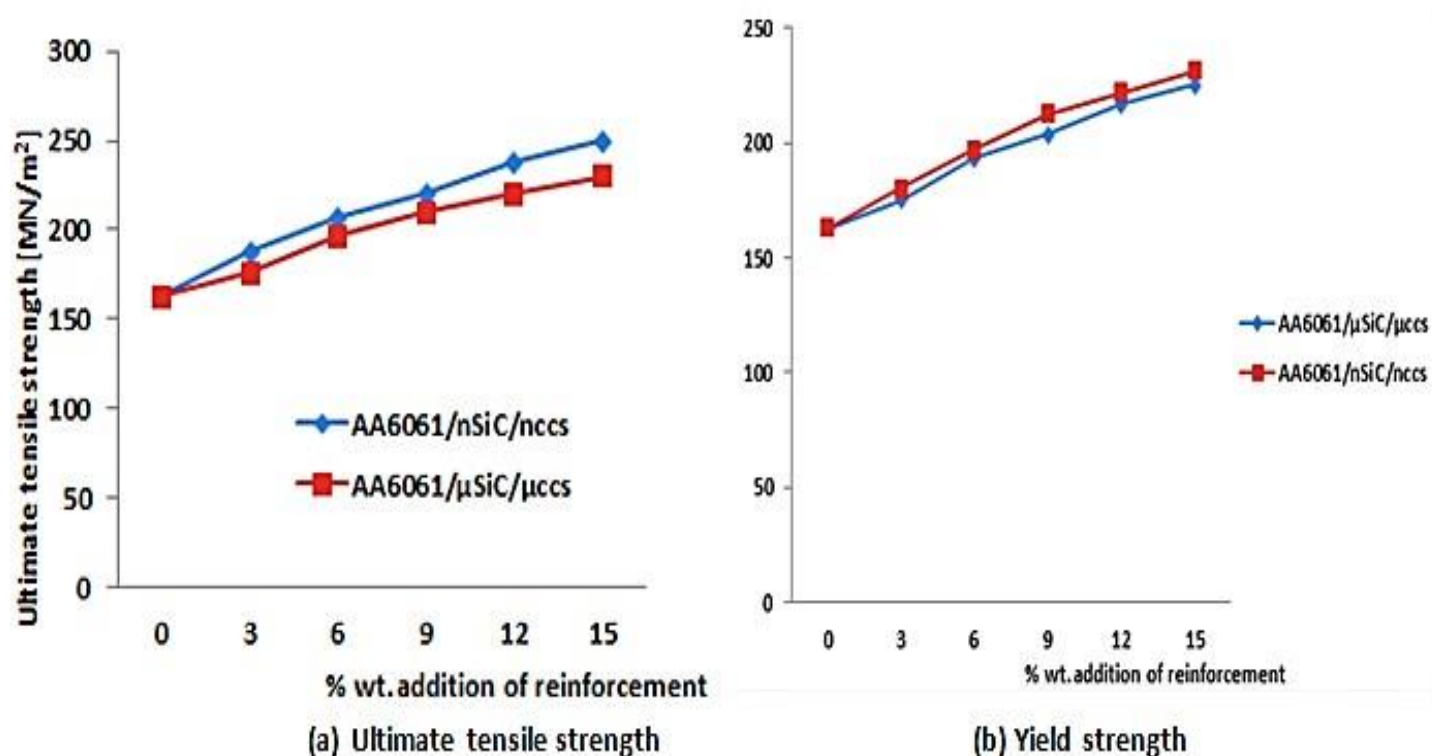


Figure 6: Variation of the strengths of Hybrid AA6061/SiC/CCS micro and nano composites with reinforcement additions

Modeling of Tensile Strength of hybrid AA6061/SiC/CCS nano composites

The experimental data for composite strength, accompanied by mathematical model predictions from earlier works and present study are shown in Table 9 and Figure 7. Widely used models for the prediction of composite strength and the proposed model are listed in Table 10. Table 9 and Figure 7 indicates that the results of the predictions by the proposed model in this work are in excellent agreement with those of the

experimental data, Voigt's model and Punkankzy's model and thus validates its suitability for tensile strength prediction of particle reinforced hybrid metal matrix composites. Further, it was observed that the popular Voigt's model gives significantly higher values than the measured values and those by Reddy and the present work. This is because the Voigt's model does not take into account the effect of reinforcement particle size, porosity and interfacial reactions in the composite. However, a modified Voigt model was developed by introducing a correction factor that accounts for the effect of particle size, porosity and interfacial interactions that was determined from the experimental data of the composite. Hence the results of the modified Voigt model given in Table 9 are consistent with those of the experimental data, Punkankzy's model as well as the proposed model.

Table 9: Comparison of Experimental and Model predicted composite strengths for the Al6061 composites

S/N	Wt.% of nSiC and nCCS	Measured σ_c (MPa)	Voigt's		Punkankzy's model		Proposed model	
			Predicted σ_c (MPa)	Residuals σ_c (MPa)	Predicted σ_c (MPa)	Residuals σ_c (MPa)	Predicted σ_c (MPa)	Residuals σ_c (MPa)
1	3	187.5	187.51	0.01	187.60	0.10	187.34	- 0.16
2	6	207.0	206.90	- 0.10	207.20	0.20	207.12	- 0.12
3	9	220.1	220.10	0.00	220.14	0.04	220.09	- 0.01
4	12	238.0	237.90	- 0.10	238.04	0.04	238.28	0.28
5	15	249.5	250.00	0.50	249.40	- 0.10	249.68	0.18

The value of strength constant k_y for the proposed model (see Table 10) as determined from experimental data ranges from 3.92×10^{-3} MPa (at 6wt.% addition of nSiC and nCCS) to 4.48×10^{-3} MPa (at 30wt.% addition of nSiC and nCCS) while the correction factor for the modified Voigt's model ranges from 24.04 to 82.58 MPa for reinforcement addition ranging from 6wt. % to 30 wt. % addition of nSiC and nCCS. Also the B value in Punkankzy's model varies in between 4.48 to 5.74 in decreasing amount of reinforcement addition in contrast with the stated range of 3.49 to 3.87 as given by Punkankzy. The variation in B values compared with those of Punkankzy was attributed to non-inclusion of effect of precipitate formation, agglomeration, voids/porosity and dislocation barriers in the response computation.

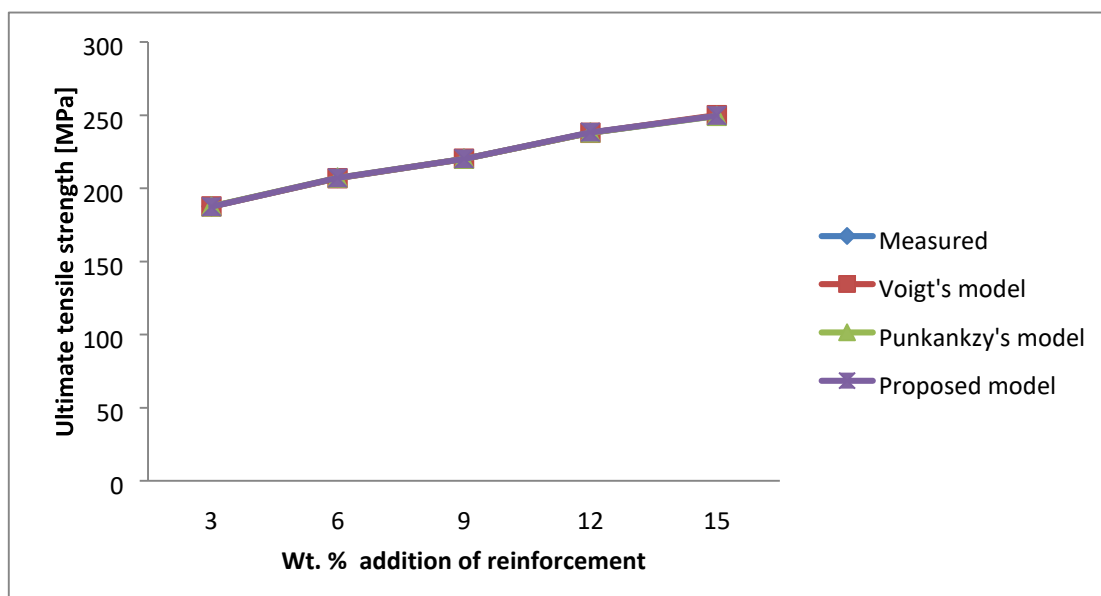


Figure 7: Comparative plots of hybrid nano-composite strength obtained from model prediction

Table 10: Models for prediction of composite strength

S/N	Author(s)	Model	Equation
1	Voigt	$\sigma_c = V_m \bar{\sigma}_m + \sum V_{rj} \bar{\sigma}_{rj} + \varepsilon_f \quad \{j = 1, 2, \dots, n\}$	3
2	Punkankzy	$\sigma_c = \left[\sigma_m \left(\frac{1 - (\sum V_{p_i})}{1 + 2.5(\sum V_{p_i})} \right) \right] e^{B(\sum V_{p_i})} \quad \{i = 1, 2, n\}$	2
3	Present work	$\sigma_c = \frac{1}{2} (\sum V_{p_i}) \sigma_m + \frac{k_y}{\sqrt{(\sum d_{p_i})}} \quad \{i = 1, 2, n\}$	7

However, comparative analysis of the composite strength precisely obtained from experiment and predictive models shows that the model- predicted values, deviated from experimental results as presented in Figure 5. The maximum tolerable deviations (residuals) of model predictions from experimental data were 0.2%, 0.1% and 0.12% for Voigt's model, Punkankzy's model and the proposed model respectively. The observed deviations are caused by the effect of other process parameters such as stirring speed, stirring temperature, particle agglomeration, melt/mould interactions (reactions) etc. which were ignored during the model development and evaluations. It can therefore be concluded that the proposed model is suitable for prediction of tensile strength of nano-particulate composites with some modifications. The modifications may be the introduction of error factor that will account for the various process parameters affecting the measured response (strength), which can be determined from experimental data for nanocomposite composition.

Microstructure of Hybrid AA6061/nSiC/nCCS Composites

The SEM micrographs of the AA6061 alloy and those of the composites are shown in Figure 8. Figure

8(a) shows the micrograph of the unreinforced alloy revealing the white α -phase with equiaxed grain shape (Boppana *et al.*, 2020). Fine grains are found along grain boundaries as far as possible in the structure of AA6061 alloy. The result of SEM for microstructural examination of the specimens revealed fairly uniform distribution of nSiC and nCCS in the matrix in all %wt addition of the reinforcements (Figure 8b-d).

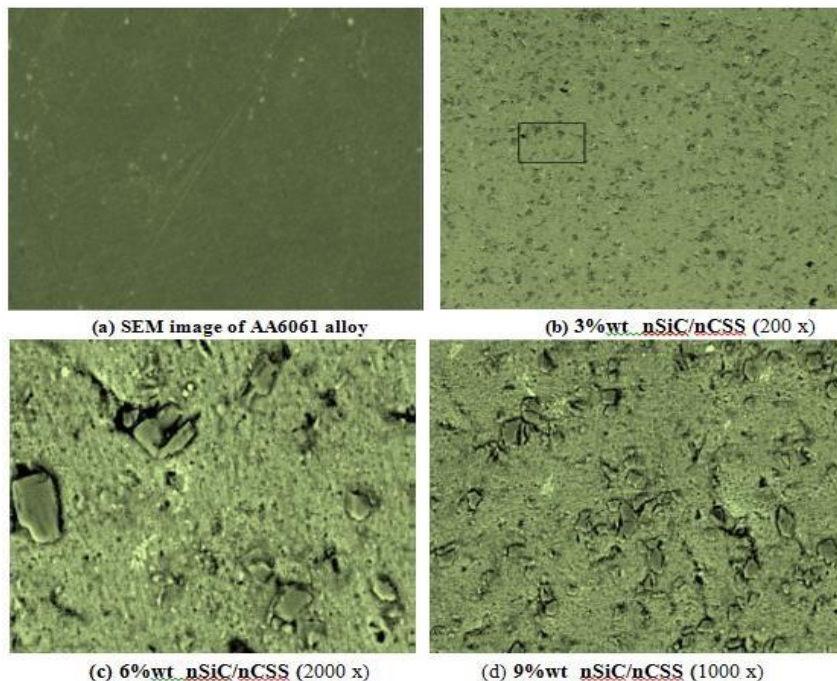


Figure 8: SEM images of hybrid AA 6061/nSiC/nCCS composites

The uniform distribution of nSiC and nCCS particles in the matrix can be attributed to a number of factors which includes, effective stirring of the melt, degassing tablet used and good wettability that resulted in improved interfacial bonding between the matrix and the coconut shell ash particles (Apasi *et al.*, 2016). No segregation of nSiC and nCCS particles were found along the grain boundaries. Distribution of particles was observed to be intra-granular, in which the majority of the particles locate inside the grains. This distribution is preferred in AMCs to have better mechanical and tribological properties. Silicon carbide and carbonized coconut nanoparticles were thermodynamically stable, and there were no pores or voids around them. The reinforcement particles resisted aluminium grain growth and resulted in nucleation sites growth, which led to the formation of finer grains. As the concentration of nSiC and nCCS particles increased in the matrix, composite properties such as hardness, UTS, and resistance to wear also showed improvement.



CONCLUSION

The hybrid AA6061/SiC/CCS composites have been successfully produced by stir casting method and characterized for analysis of density, hardness, strength and microstructural properties. The effect of reinforcement particulates size and volume fraction on the properties of the AA6061/SiC/CCS composites were also investigated. From the analysis of results obtained during this study, the following conclusions were made:

1. The incorporation of the silicon carbide and coconut shell particulates into the aluminium alloy matrix decreased the density but increased the hardness, ultimate tensile and yield strength values of the composites
2. The hybrid AA6061/SiC/CCS nano-composites acquired superior hardness and tensile strength properties with least density when compared with the hybrid micro-composite and the matrix alloy.
3. The nano-particulates reinforced composite presented maximum improvement in ultimate tensile strength value (53.4% of that the unreinforced matrix) at reinforcement level of 15wt.% nSiC/nCCS.
4. The microstructure analysis of the composites revealed a uniform distribution of the reinforcement particles in the matrix. This resulted in excellent bonding properties and grain refinement.
5. The composites can be used in areas where higher strength to weight ratio is required within the aerospace, automotive and electronic industries such as cylinder liners in engines, aluminium calipers and power electronic modules.

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