Extraction of Silica from Cassava Periderm using Modified Sol-Gel Method

J. A. Adebisi^{1*}, J. O. Agunsoye², S. A. Bello³, M. Haris⁴, M. M. Ramakokovhu⁵, M. O. Daramola⁶, S. B. Hassan²

¹Department of Materials and Metallurgical Engineering, University of Ilorin, Ilorin, Nigeria.

²Department of Metallurgical and Materials Engineering, University of Lagos, Akoka, Nigeria.

³Department of Materials Science and Engineering, Kwara State University, Malete, Nigeria.

⁴Department of Physics, Karunya University, Karunya Nagar, Coimbatore 641 114, India.

⁵Department of Chemical, Metallurgical and Materials Engineering, Tshwane University of Technology, Pretoria, South Africa.

⁶School of Chemical and Metallurgical Engineering, Faculty of Engineering and the Built Environment, University of the Witwatersrand, Wits 2050, Johannesburg, South Africa.

ABSTRACT: Cassava peridem wastes are generated and disposed indiscriminately or burnt due to zero economic value. In this study, modified sol-gel synthesis of amorphous silica nanoparticles from cassava periderm (CP) was investigated. The wastes were pretreated with HCl to remove soluble metallic impurities. Both treated and untreated CP flakes were calcined at 700 °C to obtain their respective ashes. The ashes were leached with HCl to remove soluble metallic oxides and thereafter sol-gelled to obtain silica gel modified with ethylene glycol. The gel was dried at 80 °C for 18 hours and then characterized using SEM, EDX, XRD, FT-IR, Raman, TEM and PSA. The SEM micrographs showed that silica modified with ethylene glycol is less agglomerated with higher silica yield and lower particle sizes. All the silica produced showed similar functional groups and non-crystallinity. The silica nanoparticles could be used as starting materials for synthesis of silicon nanoparticles.

KEYWORDS: agricultural waste, cassava periderm, amorphous silica, agglomeration, sol-gel.

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I. INTRODUCTION

Agricultural wastes have been utilized for production of several useful materials to alleviate challenges associated with their disposal. Agricultural wastes have been used for production of solid fuel briquettes, adsorbents, cellulose, activated carbon, silica, silicon, refectory and ceramic products, livestock feed, inhibitors, biofuels, construction materials, lignin, and composite reinforcement (Agunsoye et al., 2017, Agbenyeku and Aneke, 2014, Heuzé et al., 2014, Aigbodion et al., 2010, Kalderis et al., 2008, Kurniawan et al., 2011, Ki et al., 2013, Liu et al., 2014, Odusote et al., 2016, Achinivu et al., 2014, Adepoju et al., 2016, Venkateswaran et al., 2012, Le Normand et al., 2014). Most often, these wastes are generated and disposed indiscriminately or burnt, resulting in environmental pollution with adverse effects on climate (IEA, 2016). Interestingly, Nigeria is the largest producer of cassava, and thus generates cassava periderm as a complete waste with no known value (Adebisi et al., 2017a).

Silica occurs naturally as quartz with largest percentage in the earth crust (Zulehner et al., 1997). It is crystalline in nature with low reactivity. Amorphous silica on the other

*Corresponding author's e-mail address: adebisijeleel@gmail.com

hand finds applications in many areas owing to its advantage of being highly reactive. Silica has been utilized for several applications across many industries such as pharmaceuticals, archaeology, biomedical, electronics and feedstock in silicon production (birth of semiconductor revolution) (Wikipedia, 2018). It has been used as intensive blue light emitter, fining agent in food industry, powder flow agent in pharmaceutical industry, extra-terrestrial particles collectors, DNA and RNA extractors, hard abrasives in toothpaste, inhibitor of human breast cancer, desiccant, oil absorption, capacitors, hydrogen storage and silicon production (Yu et al., 1998, Castellari et al., 2001, Forny et al., 2007, Chattoraj et al., 2011, Westphal et al., 2014, Tan and Yiap, 2009, Joiner, 2009, Joshi et al., 2008, Negre et al., 2016, Mohammad-Rezaei and Razmi, 2016, Yuvakkumar et al., 2014b, Konwar and De, 2014, Villareal et al., 2013).

The non-crystalline type has been synthesized from sodium silicate (Sudirmana et al., 2012), water glass (Lee et al., 2007) and tetraethoxylsilane (TEOS) (Liu et al., 2006). It has also been produced from agricultural wastes (Zemnukhova et al., 2012, Mohanraj et al., 2012, Adepoju et al., 2016, Vaibhav et al., 2015). The methods used for the synthesis from agricultural wastes involve combination of processes like acid treatment, partial burning (charring), burning, calcination, leaching, enzymatic treatment (Decker et al., 2007), pyrolysis, hydrolysis and sol-gel (Nayak and Bera, 2012, Mansha et al., 2011). Extensive review on production of silica from agricultural wastes has been reported by the authors (Adebisi et al., 2017a). Adepoju et al. (2016) synthesized silica nanoparticle from cassava periderm with average grain size of 62.69 nm using conventional solgel method. The particles were reported to be lumpy and observed to be highly agglomerated. This work investigated the effect of acid pre-treatment and addition of ethylene glycol as capping agent in modified sol-gel method for control of particle size and agglomeration.

II. MATERIALS AND METHODS

A. Materials

Cassava periderm (CP) used was extracted from cassava purchased from traders at Better Life market, Isale-Osun, Osogbo, Osun State. Reagents used include sodium hydroxide (NaOH), hydrochloric acid (HCl) and ethylene glycol (CH2OH.CH2OH). All reagents used were analytical grades from NICE Chemicals ltd., Kerala, India except NaOH purchased from Rankem Laboratory, Mumbai, India. Double distilled water (DDW) was used during all processes. Munkcell Ahlstrom filter paper (Grade 389, Ø125 mm; Whatman No 41 equivalent) was used with vacuum filter for all filtration processes.

B. Methods

Cassava periderm was prepared as described in Adebisi et al. (2017b). The cassava periderm was treated in 0.1 M HCl at 45 ± 5 °C for 90 min to remove some soluble compounds. The flakes were then rinsed with DDW five times to remove the leachate. The untreated and treated CP flakes were oven dried at 105 °C for 5 hours and then labelled untreated cassava periderm (UCP) and treated cassava periderm (TCP), respectively. UCP and TCP were calcined separately at 700 °C for 4 hours to obtain their respective ashes, labelled untreated cassava periderm ash (UCPA) and treated cassava periderm ash (TCPA).

500 ml of 0.54 M HCl was measured in two 1000 ml Erlenmeyer flasks and heated to 42.5 ± 5 °C on magnetic stirrer hot plates. 25 g each of UCPA and TCPA were separately leached in the warm HCl solution for 75 min, stirred at 600 rpm. The mixtures were filtered and residues rinsed with DDW five times to remove traces of the acid. The residues were then oven dried at 80 °C for 18 hours and labelled leached untreated cassava periderm ash (LUCPA) and leached treated cassava periderm ash (LTCPA).

500 ml of 2.0 M NaOH was heated to boiling point before introduction of 15 g of each LUCPA and LTCPA, in different 1000 ml Erlenmeyer flasks. The mixture was magnetically stirred for 2 hours at 700 rpm. The concentration was kept constant by addition of DDW to maintain the volume. The mixture was allowed to cool before filtration with Whatman No 41-filter paper to get sodium silicate (see Equation 1). LUCPA and LTCPA sodium silicates were separately stirred at 900 rpm at room temperature and titrated with 1.0 M HCl to precipitate silica. The mixtures were allowed to age for 18 hours before filtration to obtain silica gels. The gels were rinsed five times to remove NaCl (see Equation 2). For the modified sol-gel approach, 20 v/v% ethylene glycol (EG) was added to freshly prepared 500 ml LTCPA sodium silicate prior to the titration. Rinsing was repeated five times before 50 % EG was stirred with the rinsed gel before final filtration.

The three gels were oven-dried at 80 °C for 24 hours. Modified gel was thereafter dried in two stages at 130 °C for 60 min to remove water of hydration and then 250 °C for another 60 min to vapourize EG. All the silicas were labelled LUCPA silica, LTCPA silica and silica (EG). Silica (EG) was finally thermally treated at 1000 °C for 2 hours to examine the crystalline phases present.

$$\begin{split} SiO_{2(s,impure \ ash)} &+ 2NaOH_{(aq)} \rightarrow Na_2SiO_{3(aq)} + H_2O_{(l)} \quad (1) \\ Na_2SiO_{3(aq)} &+ 2HCl_{(aq)} \rightarrow SiO_{2(s,gel)} + 2NaCl_{(aq)} + H_2O_{(l)} \quad (2) \end{split}$$

C. Characterization

Thermogravimetric analysis (TGA) was carried out to investigate the effect of acid pre-treatment on thermal stability of CP. Scanning electron microscopy (SEM), energy dispersive X-ray spectroscopy (EDS) and X-ray diffractometry (XRD) were used to investigate morphology, elemental composition and phases of compounds, respectively, in all the samples. Silica (EG) was finally characterized with Fourier transform infrared spectroscopy (FT-IR), Raman spectroscopy, transmission electron microscopy (TEM) and particle size analysis (PSA) for its surface chemistry, functionality, morphology and particles sizes (and distribution), respectively.

III. RESULTS AND DISCUSSION

A. Thermogravimetric Analysis (TGA)

Figure 1 shows the TGA curves for UCP and TCP thermally decomposed in Nitrogen atmosphere. Both samples showed significant mass loss as the operating temperature increased from room temperature to 945 °C. TCP is thermally more stable due to removal of some phases during the acid treatment (Carmona et al., 2013). The first stage of thermal degradation that occurred at 105 °C could be attributed to dehydration of the flakes and mass loss of 6.7 and 4.3 % for UCP and TCP, respectively. Further loss at temperatures between 105 °C and 223 °C could be associated with removal of water of crystallization and some light volatiles (Adebisi et al., 2017b).

The second stage of significant mass loss of about 40 % (between 362 °C and 453 °C) is the same for both UCP and TCP. This shows that the hemicelluloses are unaffected by the acid treatment. TCP gave a distinctive cellulose removal between 362 and 453 °C, followed by carbonization stage that indicate decomposition of lignin. There exists overlap in

cellulose and lignin decompositions in UCP up to 783 °C. The remaining chars left are 16.8 and 3.1 % for TCP and UCP, respectively. The higher residue in TCP could be due to loss of some lignocellulosic phases during acid treatment (Rahman et al., 1997, Carmona et al., 2013). Oxidative thermal decomposition of cassava periderm has been reported with optimum calcination temperature of 700 °C against higher conditions observed in the Nitrogen environment (Adebisi et al., 2017b).



Periderm (UCP) and treated Cassava Periderm (TCP).

B. XRD Analysis

X-ray diffraction patterns for UCP and TCP with further processing conditions are shown in Figure 2. The patterns for UCP and TCP indicate amorphous phases with few crystalline phases at 24.7, 26.9, 35.2, 44, 64.3 and 77.5 diffraction angles with significant difference at 24.7 and 26.9. Calcination of both UCP and TCP suppressed their amorphic natures with identifiable crystalline phases. The effect of acid treatment is prominent at this stage since fewer peaks shown for TCPA confirm removal of some phases during acid pretreatment. Leaching of UCPA and TCPA with HCl further purified the residues. The XRD peaks detected correlate with Joint Committee on Powder Diffraction Standards (JCPDS) reference numbers of SiO2 (78-1254), KAlSi3O8 (83-1604), NaAl(SiO3)2•3H2O (03-0234) for LUCPA while only SiO2 and CaAlAlSiO6 (70-2129) are observed for LTCPA. Also, crystallinity of the residues is improved by the postcalcination acid treatments.

Silicas obtained from LUCPA, LTCPA and silica (EG) after several rinsing with DDW and drying (up to 250 °C for silica with EG) showed similar amorphous natures as shown in Figure 3. Amorphous nature of silicas obtained by chemical methods has been reported (Pijarn et al., 2010, Adebisi et al., 2017c). Ethylene glycol retained some NaCl, even after several rinsing, as observed from the XRD peaks before vapourization of the EG. Annealing of the silica at 1273 K transformed the amorphous silica to crystalline silica with JCPDS reference code 82-0512 which is found to be cristobalite with tetragonal crystal system (Yuvakkumar et al., 2012, Venkateswaran et al., 2012, Haastrup et al., 2016).



Figure 2: XRD of untreated and treated Cassava Periderm (UCP and TCP), Cassava Periderms calcined at 700 °C (UCPA and TCPA) and Acid Leached Chars (LUCPA and LTCPA).

This enables easy calculation of crystal size of amorphous powders using Scherrer equation. The crystallite size of the silica produced is given in Table 1 with average crystal size of 16.98 nm.



Figure 3: Silicas obtained without/with Ethylene Glycol (EG) and after annealing at 1273 K.

C. Scanning Electron Microscopy (SEM)

Figure 4 shows the SEM micrographs of cleaned CP (a1), treated CP (b1), calcined UCP (a2) and TCP (b2), leached UCPA (a3) and TCPA (b3). It is observed that UCP and TCP have nonporous and bulky tissues. Calcination of both UCP and TCP produced fine particles of UCPA and TCPA but TCPA is less fragmented due to its higher thermal stability. This is more pronounced after leaching of UCPA and TCPA as could be observed in Figure 4 (a3 and b3).



Figure 4: SEM Micrographs of (a) untreated CP, (b) HCl treated CP for (1) as received/treated, (2) calcined at 700 °C, and (3) HCl leached calcines.

The SEM micrographs of silicas produced are shown in Figure 5. Degree of agglomeration is highly influenced by processing conditions of the silicas. LUCPA silica is extremely agglomerated Figure 5(a) but reduced in the silica produced from pre-treated CP (Figure 5(b)). Drying silica (EG) at 105 °C doesn't remove the ethylene glycol that seems to bind the silica particles together as shown in Figure 5(c). Vapourization of ethylene glycol above its boiling point, at 250 °C left dispersed small agglomerates (Figure 5(d)). The morphology of the amorphous silicas produced revealed no definite shape in all the processing routes. This implies that particles formed are still agglomerated irrespective of methods of production.

Table 1: Crystal Size of Silica produced using Scherrer Equation.

No.	Peak pos. [2θ (degree)]	FWHM [2θ (degree)]	Rel. Int. [%]	Crystallite size (nm)
1	21.7779	0.2952	100.00	27.4
2	28.2853	0.5904	2.41	14.6
3	31.2592	0.5904	2.62	14.8
4	35.8209	0.3936	17.39	21.2
5	44.3416	0.5904	2.65	14.5
6	56.5653	0.9600	3.36	9.4



Figure 5: SEM Micrographs of Silica obtained from (a) asreceived CP, (b) HCl treated CP, (c) Ethylene Glycol Modified Silica, and (d) Silica after Vaporization of EG

D. Energy Dispersive Spectroscopy

Figure 6 shows the elemental composition of CP through the various processes such as acid pre-treatment, calcination and post calcination acid leaching. Silicon was not detected in UCP (Figure 6(a1)) which improved by acid leaching to a slight detectable level in TCP (Figure 6(b1)). Calcination produced residues composed mainly of elements Na, Mg, Al, Si, K, Ca and Fe. The high oxygen proportions indicate presence of oxides of the elements. Figure 6(a2 and b2) revealed higher silicon percent in both UCPA and TCPA.

Silicon percent increased after post calcination treatment and removal of soluble impurities. Both acid pre-treatment and post calcination leaching are effective in removing Na completely while post calcination treatment is effective for Mg removal. The acid treatments reduced Mg, K and Ca while Al and Fe are unaffected. Acid treatments have been reported to improve quality of silica produced from agricultural wastes (Chandrasekhar et al., 2005). Highest silicon proportions are obtained after post calcination treatments of with/without acid pre-treatments [Figure 6(a3 and b3)].



Figure 6: Energy Dispersive X-Ray Spectra and Composition of (a) untreated CP, (b) HCl treated CP for (1) as received/treated, (2) calcined at 700 °C, and (3) HCl leached calcines.

Table 2 shows the chemical compositions of silica obtained using three different routes. Presence of Na could be attributed to those retained from unrinsed NaCl from precipitation process. Al present in the modified silica is very low compared to other silicas produced. Both acid pre-treatment and introduction of ethylene glycol improved the purity of silica. Chakraverty et al. (1988) has reported effectiveness of acid treatments on purity of silica.

Table 2: Chemical Composition of Silica Produced from asreceived/treated CP and Ethylene Glycol (EG) Modified Sol-Gel Precipitated Silica.

Elements	LUCPA silica	LTCPA silica	Silica (EG)
0	54.70	54.95	56.48
Na	10.03	5.69	2.52
Al	10.97	9.15	3.01
Si	24.30	29.75	36.89
К	ND	0.47	0.09
Ca	ND	ND	0.08
Fe	ND	ND	0.65
Cu	ND	ND	0.29

E. Functional groups analysis in modified silica by FT-IR

The FT-IR spectrum of silica produced by morphological modification with ethylene glycol is shown in Figure 7. Silica is characterized by band between 400 and 1300 cm⁻¹ (Carmona et al., 2013, Espíndola-Gonzalez et al., 2010, Liou, 2004, Liou and Yang, 2011, Rafiee et al., 2012).



Figure 7: FT-IR Spectrum of Silica (EG) after drying at 250 °C.

The spectra from prepared silica exhibits bands at 448, 796 and 1085 cm^{-1} , corresponding to characteristics rocking of the Si-O bond, its bending and symmetric and asymmetric stretching of Si-O-Si bond, respectively (Espíndola-Gonzalez et al., 2010).

Drying the samples at 65 °C left the samples with OH and H-O-H bonds for silanol and adsorbed water (Carmona et al., 2013). The band at 1642 cm⁻¹ corresponds to H-O-H twisting while band spreading from 2600 to 3800 cm⁻¹ represent stretching vibrations of the surface silanols (Si-OH) groups excited by hydrogen bonds of either water adsorbed or those between Si-O molecules (Carmona et al., 2013, Rafiee et al., 2012).

F. Raman Spectroscopy

Figure 8 presents the Raman spectra of silica modified with ethylene glycol. Raman bands between 487 and 600 cm⁻¹ presents Raman spectra of SiO2 (Gottardi et al., 1984). The stretching band of silanol (Si-OH) is at 980 cm⁻¹ but spreads across 800 and 1100 cm⁻¹ (Gottardi et al., 1984, Yuvakkumar et al., 2014a, McMillan and Remmele, 1986). Hydroxyl (OH) bending and stretching band peaks are at 1630 cm⁻¹ and near 3450 cm⁻¹. The weak band near 2350 cm⁻¹ may correspond to silanol groups intra-tetrahedrally hinged by hydrogen bonding across SiO4 (McMillan and Remmele, 1986). The absorption bands for Si-H is also in the range from 2100 to 2400 cm⁻¹ (Vella et al., 2011).



G. Analysis of particle size and distribution (TEM and PSA)

Figure 9 shows TEM image of silica (EG) produced and dried at 250 °C. All the particles were found to be below 100 nm but with some agglomerated spherical morphology apparently in the upper part of the image. Figure 10 presents the particle size distribution of the silica nanoparticles produced. The maximum distribution was found between 3.12 to 50.75 nm. These values are in agreement with those obtained from Scherrer crystallite sizes.



Figure 9: TEM Image of Nano Silica (EG) produced after drying at 250 °C.



Figure 10: Particle Size Distribution of Nano Silica (EG) after drying at $~250\ ^{\rm o}{\rm C}.$

IV. CONCLUSION

Silica has been produced from cassava periderm using three different routes. Acid treatments before and after calcination were found to be effective for reducing or eliminating soluble metallic impurities. Silica obtained from acid pre-treatment, post-calcination treatment and ethylene glycol modified sol-gel methods are found to yield higher purity silica particles that are in nano range from 3.12 to 50.75 nm.

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