Variation of Physico-Chemical and Textural Properties of Laboratory Prepared Red Mud Through Acid and Thermal Activations

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To cite this article:

Received: March 28, 2017; Accepted: April 18, 2017; Published: June 1, 2017

Abstract: Red mud prepared in the laboratory was activated using hydrochloric acid with concentrations from 2-4 mol/L, heat at 900°C and combined acid and heat at 900°C, and the respective yields evaluated. The physico-chemical properties of these samples were determined using pH, EC, XRF, XRD, SEM, TGA-DTA analyses and textural properties (isotherm type, BET surface area, pore size distribution, total pore volume, average pore size, external surface area, micropore volume, micropore area) determined by nitrogen gas adsorption-desorption. Acid activation reduces red mud quantity than by thermal means. Raw red mud showed a pH of 11.0, EC of 2.50 x 10⁹ µS/cm, BET surface area of 13 m²/g, total pore volume of 0.063 cm³/g with the major oxides being Fe₂O₃, Al₂O₃, TiO₂ and SiO₂. It contain mesopores > macropores > micropores. All these properties are improved variably by acid and thermal activation. Acid modification increase alumina content but decreases that of iron. Conversely, heat treatment increase iron content but reduces alumina content. Combined acid and heat treated red mud shows high thermal stability than untreated and acid treated red muds. These results show that possible different applications of red mud can be achieved depending on modification conditions.

Keywords: Bauxite, Red Mud, Activation, Minim-Martap, Properties

1. Introduction

Red mud is the insoluble residue resulting from caustic alumina production via the Bayer process, often colloquially referred to as red mud or bauxite residue or Bayer process tailings [1-2]. About 90% of raw bauxite ore goes into the waste as alkaline red mud slurry during processing [3] as for every ton of alumina produced, between 1-2 tons (dry weight) are produced depending on the bauxite source and alumina extraction efficiency [4-5]. Globally, between 60 - 120 million tons are produced annually [6-7]. This may lead to serious pollution of the surrounding soil, air and groundwater due to its high pH (10-13) [8-9].

However, red mud contains a number of valuable metals and minerals from parent bauxite consisting of gibbsite Al(OH)₃, diasporic (AlO(OH)), hematite (Fe₂O₃), goethite (FeO(OH)) and those introduced during the Bayer process such as sodalite, cancrinite etc. [2, 7, 10]. The typical constituents of red mud (% w/w) are: Fe₂O₃ (30–60%), Al₂O₃ (10–20%), SiO₂ (3–5%), Na₂O (2–10%), CaO (2–8%), TiO₂ (trace–10%) [11-12]. In general, red mud is a very fine material in terms of particle size distribution, having an average particle size <10µm. Typical values would account for 90% volume below 75 µm. The specific surface area (BET) is a round 10-25 m²/g. Red mud contains various particles with different size and shape. Some crystals can also be found in the sample. In addition, it also presents porous surface [10].

Due to the alkaline nature and the chemical and mineralogical species present in red mud, this solid waste causes a significant impact on the environment and proper disposal of waste red mud presents a huge challenge where alumina industries are installed. The disposal cost is high, accounting for about 5% of alumina production [10, 13]. Up to now, red mud has found limited applications in brick, glass, aerated concrete block, road base material, filling
material in mining and plastic, as adsorbent to adsorb heavy metal ions and nonmetallic ions from wastewater and soil and SO$_2$ in the waste gas [7, 10]. Unfortunately, none of these applications has been economically applied on an industrial scale [10].

Because of the environmental impact of red mud and its enormous volume, its treatment to change the alkalinity is extremely urgent issues with the aim of converting the harmful red mud to useful products with improved properties for different small and industrial applications. Many physical, chemical and mechanical methods are available for the modification of structural, textural and surface properties of solids. They include: acid activation, heat treatment, pillaring, crushing etc. [14]. The appropriate method depends strongly on the conditions of the parameters, origin of the ore and subsequent use of treated material. Hydrochloric acid (HCl) has been the most used acid in red mud activation with concentrations of up to 2M. Sketchy results are reported on the behaviour of chemical composition (as oxides) of red mud resulting from different activation methods. Also very few studies report properties of red mud treated with combined acid and thermal treatment in particular and the physico-chemical and textural properties of red mud activated with varying concentration of HCl in general.

The characteristics of red mud depend on the bauxite source extraction efficiency and modification conditions [15]. Thus, in this work, red mud (RM) prepared from Minim-Martap (Cameroon) bauxite in the laboratory was activated with HCl (concentrations 2.0-4 mol/L), noted nRMA (n = acid concentration used), combined acid and heat treatment at 900°C, noted RMA and calcination of raw red mud at 900°C, noted RMC. The pH, EC and oxide composition of RM, nRMA, RMAC and RMC samples as well as the XRD, SEM, TG-DTA and nitrogen gas adsorption-desorption (isotherm type, BET surface area, pore size distribution, total pore volume, average pore size, external surface area, micropore volume, micropore area) of RM, 2.25M RMA and RMAC samples were investigated. This project constitute the long-term objective on the utilization of red mud and further studies as published information on red mud is owner and/or refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16]. This because each refinery has unique operating details and refinery specific and not consistent in either form or content [16].

The pH and electrical conductivity (EC) of all the samples (RM, nRMA, RMAC and RMC) were measured using a Hach HQ 40D multi pH meter within two decimal places. Also, the quantitative elemental composition (expressed as oxide) of all these red mud samples was determined by X-Ray Fluorescence (XRF) while the mineralogy of the raw red mud (RM), 2.25M acid activated (RMA) and combined acid and heat activated samples (RMAC) was determined by X-ray Diffraction (XRD) using a Brucker D2 phase in Bragg Brentano geometry with a Lynxeye detector using Cu-Kα radiation at 30 kV and 10 mA. The diffraction peaks were then compared with those of red mud samples reported in literature for identification. The different phase transformations of RM, RMA and RMAC samples were investigated using the differential scanning calorimetry-thermogravimetric (DSC-TGA) techniques on a perkin-Elmer apparatus while surface morphology of RM and RMA was determined by Scanning Electron Microscopy (SEM).

2.3. Determination of Physico-Chemical Properties of Red Mud

The pH and electrical conductivity (EC) of all the samples (RM, nRMA, RMAC and RMC) were measured using a Hach HQ 40D multi pH meter within two decimal places. Also, the quantitative elemental composition (expressed as oxide) of all these red mud samples was determined by X-Ray Fluorescence (XRF) while the mineralogy of the raw red mud (RM), 2.25M acid activated (RMA) and combined acid and heat activated samples (RMAC) was determined by X-ray Diffraction (XRD) using a Brucker D2 phase in Bragg Brentano geometry with a Lynxeye detector using Cu-Kα radiation at 30 kV and 10 mA. The diffraction peaks were then compared with those of red mud samples reported in literature for identification. The different phase transformations of RM, RMA and RMAC samples were investigated using the differential scanning calorimetry-thermogravimetric (DSC-TGA) techniques on a perkin-Elmer apparatus while surface morphology of RM and RMA was determined by Scanning Electron Microscopy (SEM).

2.4. Determination of Textural Properties of Red Mud

These properties for the RM, RMA and RMAC samples were determined by gas adsorption using nitrogen adsorption–desorption at the liquid nitrogen temperature (~195, 800°C) using a micromeritics tristar 3000 apparatus. The isotherm type was established by plotting quantity of gas adsorbed against...
experimental P/P₀ values. Samples isotherms obtained were compared with established models [18]. The BET plot was used to determine the total surface area, total pore volume and average pore size. The t-Plot was used to determine the external surface area, micropore volume and micropore area while the pore distribution was calculated using BJH (Barrett-Joyner-Halenda) method [19-20].

3. Results and Discussion

3.1. Yield of Red Mud After Activation

Results are presented in Table 1 which shows that simple acid activation reduces considerably the quantity of red mud than thermal activation. The percentage of the reduction of red mud increases as the acid concentration increases. This shows that acid treatment induced the dissolution of the oxides of red mud namely: Fe₂O₃, Al₂O₃. This is due to the fact that acid activation often replaces exchangeable cations with H⁺ ions and Al³⁺ and other cations escape out of the main structure. This increases the acidity of the material, along with the elimination of several mineral impurities and partial dissolution of the external layers [14]. Thermal treatment leads to little reduction of the initial amount of red mud. This could be attributed to dehydroxylation of adsorbed water at low temperature and dehydroxylation of hydroxides at higher temperature. It is also observed that acid-thermal treatment shows only 4% weight loss. This is probably due to the fact that this material contains only oxides. Of course most hydroxides have been dissolved during acid treatment.

<table>
<thead>
<tr>
<th>Yield (%)</th>
<th>2M RMA</th>
<th>2.25M RMA</th>
<th>2.5M RMA</th>
<th>3M RMA</th>
<th>4M RMA</th>
<th>2M RMAC</th>
<th>RMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>65</td>
<td>60</td>
<td>35</td>
<td>27</td>
<td>23.5</td>
<td>96</td>
<td>85</td>
<td></td>
</tr>
</tbody>
</table>

3.2. Physico-Chemical Characteristics

3.2.1. pH and EC

From Table 2 it is seen that, a pH of 11.0 was found for the red mud prepared. Generally the pH of red mud varies from 10-13 [5, 21]. This is due to the presence OH⁻ resulting from high amount of NaOH used in the Bayer process [22]. Acid activation leads to a drop in pH but with increased acid concentration the pH increases. These variations are due to the different buffering and dissolution reactions [2, 16] that occur in red mud. It should be noted that red mud particles contain two types of hydroxyl groups; free hydroxide ions (OH⁻) and ionized surface hydroxyl groups (S-O-H). Initially red mud particles consume H⁺ without a change in pH due to the presence of free hydroxide ions (OH⁻) reacting with protons (OH⁻ + H⁺ → H₂O) more readily than the ionized surface hydroxyl groups. When all the free hydroxide ions are consumed (at neutral pH) and protons added to the slurry, they are then consumed by surface hydroxyl groups (S-OH + H⁺ → S-OH₂⁺), causing a drop in the pH [23].

Our untreated red mud had an electrical conductivity (EC) of 2.5 x 10⁹ µS/cm (Table 2). This value falls within the range of EC usually reported for red mud, (1.4-28.4) x 10⁹ µS/cm [24]. This low EC value of our caustic red mud indicates low mobility of different ions within the red mud, thus, showing the compact nature of the residue. In the absence of detailed solution data, it is the only feasible quantity to measure total cation and anion concentrations and to estimate ionic strength. In red mud, ionic strength determines the double-layer thickness of charged particles and consequently the physical behavior such as dispersion and coagulation. The EC is due to high Na⁺ concentrations in solution and in the solid phases. Calcium, magnesium and other cations do not contribute significantly to the EC as their concentrations are negligible in solution at pH above 10 [2]. Furthermore, the total amount of CaO and MgO is lower than 1% in our sample. The drop in EC values from 2.5 x 10⁹ µS/cm in RM to 1432 µS/cm in 2M RMA, 1410 µS/cm in 2.25M RMA, 542 µS/cm in RMC and 16.7µS/cm in 2M RMAC is expected. This results from the fact that in caustic red mud, there are still several impurities and many free hydroxyl ions and sodium ions giving high EC, but when red mud is acid treated; the acid removes most of the impurities and these mobile ions (Na⁺, Al³⁺ and other cations escape out of the main structure. But it is noticed that EC increases again, from 1410 µS/cm in 2.25M RMA to 2.28 x 10⁹ µS/cm in 2.5M RMA followed by a drop again. According to Palmer et al. [21], this increase indicates the increase in soluble salt content.

<table>
<thead>
<tr>
<th>pH</th>
<th>RM</th>
<th>2M RMA</th>
<th>2.25M RMA</th>
<th>2.5M RMA</th>
<th>3M RMA</th>
<th>4M RMA</th>
<th>2M RMAC</th>
<th>RMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>11.00</td>
<td>11.00</td>
<td>11.00</td>
<td>11.00</td>
<td>11.00</td>
<td>11.00</td>
<td>11.00</td>
<td>11.00</td>
<td></td>
</tr>
<tr>
<td>2.5 x 10⁹</td>
<td>1432</td>
<td>1410</td>
<td>2.28 x 10⁹</td>
<td>978</td>
<td>681</td>
<td>16.7</td>
<td>525</td>
<td>10.37</td>
</tr>
</tbody>
</table>

3.2.2. Chemical Composition

Table 3 present the oxide composition of untreated red mud (RM) obtained as well as the different samples of this red mud activated under different conditions. It can be seen from these results that: The types of elements present and the trends in content are similar to those reported by other authors [2, 16, 24-25 etc.] with the major oxides present being those of Fe, Al, Ti and Si. Acid activation increases the quantities of oxides of Al, Si and Ti, with Al₂O₃ increasing from 13% in untreated red mud to 30.78% in 3M HCl activated sample, SiO₂ increasing from 2.76% in untreated red mud to 6.53% in 3M HCl activated sample and TiO₂ increasing from 2.49% in untreated red mud to 5.26% in 3M HCl activated sample.
increasing from 3.76% in untreated red mud to 17.65% in 4M HCl activated sample. However, acid activation reduces significantly the quantity of Fe$_2$O$_3$ from 48.59% in untreated red mud to 20.34% in the 4M HCl sample.

Heat treated red mud shows different behaviour with the quantity of Fe$_2$O$_3$ increasing from 48.59% in untreated red mud to 59.27% in the heat treated sample (RMC) and 51.62% in the combined acid and heat treated sample (2M RMAC). Thus, the reduction in the yield of red mud as it contains the same minerals with same intensities and 20 values. However, heat treated sample Figure 1c shows the disappearance of aluminium (except goethite) and calcite minerals. This result confirms the increase in iron content of red mud when it was heat treated at 900°C. The different peaks show high intensities. This is definitely due to the fact that minerals such as hematite, dicalcium silicate, perovskite, and CaO are not affected by temperature through the process contrary to Al and Ca bearing minerals. For example, gibbsite decomposes into Al$_2$O$_3$ and H$_2$O from 300- 550°C; calcite to CaO and CO$_2$ from 600- 800°C on heating [7].

### Table 3. Chemical composition of untreated and different activated samples of red mud in% (w/w).

<table>
<thead>
<tr>
<th>Sample</th>
<th>SiO$_2$</th>
<th>Al$_2$O$_3$</th>
<th>Fe$_2$O$_3$</th>
<th>CaO</th>
<th>MgO</th>
<th>K$_2$O</th>
<th>Na$_2$O</th>
<th>SO$_3$</th>
<th>TiO$_2$</th>
<th>MnO$_2$</th>
<th>P$_2$O$_5$</th>
<th>LOI</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>RM</td>
<td>2.76</td>
<td>16.41</td>
<td>48.59</td>
<td>0.53</td>
<td>0.08</td>
<td>0.07</td>
<td>4.23</td>
<td>0.12</td>
<td>3.76</td>
<td>0.03</td>
<td>0.11</td>
<td>9.02</td>
<td>82.33</td>
</tr>
<tr>
<td>2M RMA</td>
<td>3.44</td>
<td>24.77</td>
<td>44.90</td>
<td>0.39</td>
<td>0.10</td>
<td>0.07</td>
<td>0.36</td>
<td>0.07</td>
<td>9.99</td>
<td>0.06</td>
<td>0.21</td>
<td>16.18</td>
<td>100.54</td>
</tr>
<tr>
<td>2.25M RMA</td>
<td>2.21</td>
<td>16.10</td>
<td>52.29</td>
<td>0.05</td>
<td>0.06</td>
<td>0.04</td>
<td>0.18</td>
<td>0.06</td>
<td>5.79</td>
<td>0.03</td>
<td>0.14</td>
<td>15.46</td>
<td>92.41</td>
</tr>
<tr>
<td>2.5M RMA</td>
<td>2.99</td>
<td>21.86</td>
<td>41.36</td>
<td>0.32</td>
<td>0.09</td>
<td>0.06</td>
<td>0.06</td>
<td>0.06</td>
<td>10.04</td>
<td>0.04</td>
<td>0.20</td>
<td>22.05</td>
<td>99.15</td>
</tr>
<tr>
<td>3M RMA</td>
<td>6.53</td>
<td>30.78</td>
<td>27.11</td>
<td>0.34</td>
<td>0.13</td>
<td>0.12</td>
<td>0.19</td>
<td>0.11</td>
<td>10.58</td>
<td>0.08</td>
<td>0.16</td>
<td>23.04</td>
<td>99.17</td>
</tr>
<tr>
<td>4M RMA</td>
<td>5.79</td>
<td>29.57</td>
<td>20.34</td>
<td>0.98</td>
<td>0.13</td>
<td>0.16</td>
<td>0.24</td>
<td>0.07</td>
<td>17.65</td>
<td>0.18</td>
<td>0.18</td>
<td>23.31</td>
<td>98.55</td>
</tr>
<tr>
<td>2M RMAC</td>
<td>4.43</td>
<td>2.29</td>
<td>51.62</td>
<td>0.06</td>
<td>0.07</td>
<td>0.07</td>
<td>0.19</td>
<td>0.08</td>
<td>5.94</td>
<td>0.04</td>
<td>0.16</td>
<td>20.32</td>
<td>85.27</td>
</tr>
<tr>
<td>RMC</td>
<td>2.87</td>
<td>18.01</td>
<td>59.27</td>
<td>0.36</td>
<td>0.08</td>
<td>0.04</td>
<td>5.85</td>
<td>0.14</td>
<td>4.95</td>
<td>0.04</td>
<td>0.13</td>
<td>7.60</td>
<td>99.34</td>
</tr>
</tbody>
</table>

### 3.2.3. Mineral Composition

The XRD pattern of red mud (untreated) is shown in Figure 1a, which shows that Minim-Martap red mud is mainly composed of Hematite, goethite, quartz, gibbsite, calcite, diaspora and anatase. This composition is similar to the findings of [2]. It can also be seen from Figure 1b that acid activation thus not modifies the mineral structure of red mud as it contains the same minerals with same intensities and 20 values. However, heat treated sample Figure 1c
3.2.4. Surface Morphology (SEM)

Figure 2 shows the SEM images of untreated red mud and acid activated sample. As seen from these images, the crude red mud particles are polycrystalline and poorly-crystallized or amorphous while there is a clear visual evidence, for the new pores and fine particles generated by strong acid treatment. The different porosity observed results from different physical and chemical processes mainly removal of acid soluble salts and dissolution of iron oxide.
3.2.5. Thermal Analysis

Figure 3 shows the TGA-DTA plots of untreated (a), acid activated (b) and (c) combined acid and heat activated red mud samples. From these plots it can be seen that, the untreated (Figure 3a) and acid activated red muds (Figure 3b) show similar weight loss portions for TGA and similar phase transition for DTA. They all have three continuous weight loss portions.

From Figures 3a and 3b, it can be seen the same quantity of material is being lost in the untreated and acid treated red muds from 150-350°C (about 10%) and from 350-650°C (about 2%). However, the acid treated red mud shows a higher weight loss from 90-150°C than untreated sample. This is probably due to increase in surface water during acid treatment. In each case the first weight loss is attributed to the evaporation of physically adsorbed water [29-30] while the rapid second and third weight losses corresponds to the loss of chemically adsorbed water due principally to the decomposition of gibbsite (Al(OH)$_3$) to alumina plus H$_2$O and to decomposition of CaCO$_3$ to CaO with the release of CO$_2$ (1). Alumina then combines with the CaO to form tricalcium aluminate (2) or Gehlenite (3) in the 800–900ºC temperature range [7, 29, 31-32]. This is why a constant mass is observed for the two samples as from 800 ºC.

\[
\begin{align*}
\text{CaCO}_3 & \rightarrow \text{CaO} + \text{CO}_2 \\
3\text{CaO} + \text{Al}_2\text{O}_3 & \rightarrow \text{Ca}_3\text{Al}_2\text{O}_6 \text{ (tricalcium aluminate)} \\
4\text{CaO} + 2\text{Al}_2\text{O}_3 + 2\text{SiO}_2 & \rightarrow 2\text{Ca}_2\text{Al}_2\text{SiO}_7 \text{ (Gehlenite)}
\end{align*}
\]

The DTA curves of untreated (Figure 3a,) and acid activated (Figure 3b) Minim-Martap red mud shows three endothermic peaks. The first with a maximum at 90°C corresponds to the desorption of physically adsorbed water as already stated. It is observed that the peak is more intense for raw red mud than acid treated red mud. This shows that the latter contain less water than the former. Thus acid treatment makes the surface of red mud more hydrophobic. The second peak (at 300°C) of DTA curve corresponds to the phenomena described for TGA i.e. dehydration of gibbsite to alumina. The third peak at 600°C corresponds mostly to phase transition. At this temperature alumina combines with CaO to form aluminate or gehlenite as shown in (2) and (3). The DTA curves exhibited many incidents. This is a manifestation of many phase transitions that occur in the material. For example, the formation of Ca$_3$Al$_2$O$_6$, Ca$_2$Al$_2$SiO$_7$, Fe$_2$O$_3$ etc. For the combined acid and heat activated red mud (Figure 3c), there is only one weight loss portion which is rapid, continuous and very small (about 3%). This 3% loss can be
attributed to the loss of physically adsorbed water and decomposition of CaCO₃. Nonetheless, its DTA curve shows that is very stable on heating.

3.3. Textural Properties

3.3.1. Type of Isotherm

From the plots in Figure 4, it is evident that the isotherms of the three samples are of type IV isotherm model according to IUPAC classification [18, 33-34] characterized by monolayer formation (presence of micropores), presence of hysteresis (H1 or H2), indicating meso and macro pores for a and b samples (characteristic of many oxide gels) and formation of few multilayer. Hysteresis is absent in sample c as material is highly degraded.

3.3.2. Porosity and Surface Area Analyses

The BJH pore size distribution for untreated, acid activated and combined acid and heat treated Minim-Martap red mud is shown in Figure 5. The other textural properties determined notably, BET total surface area, external surface area, micropore area, micropore volume, diameter range, total pore volume and average pore width are presented in Table 4.

Figure 4. Isotherm plots for (a) untreated (b) acid activated and (c) combined acid and heat treated red muds.

Figure 5. Pore size distribution curves of BJH adsorption for RM, RMA and RMAC Minim-Martap red mud samples. D = Pore diameter.

The BJH curve Figure 5 shows that, red mud from Minim-Martap bauxite contains: macropores (pores > 50 nm in diameter), mesopores pores (2-50 nm in diameter), and micropores (pores < 2 nm in diameter) [19]. From Figure 5 the calculated composition in percentage of the pore size distribution shows 66.67% of mesopores in RM, 67.50% in RMA and 67.50% in RMAC. It also shows 30.77% of macropores in RM, 30% in RMA and 30% in RMAC while micropore composition shows 2.56% in RM, 2.50% in RMA and 2.50% in RMAC. The mesopores are thus dominant in all the samples with a slight increase in the activated samples. Acid and thermal activation do not significantly alter the pore distribution but significantly leads to an increase in pore volume (Figure 5) and other textural properties (Table 4). BJH results confirm the results of isotherm type shown in Figure 4.

This dominance of mesopores and macropores to a lesser extent is also shown by higher values of external surface area; 14.88 m²/g and 25.21 m²/g respectively for raw and acid treated samples Table 4. The corresponding BET total surface area is 13.1548 m²/g and 23.7978 m²/g for raw and the acid activated sample respectively. This analysis is based on the fact that external area is the area of those pores which are not micropores. Usually BET theory is applied to obtain the specific surface area of micro porous materials, although from a scientific point of view the assumptions made in the BET theory do not take into account micropore filling [20]. Mesopores, macropores and the outside surface is able to form a multiplayer, whereas micropores which have already been filled cannot contribute further to the adsorption process. The presence of micropores was confirmed using y-
intercept values from the t-plot (an intercept different from zero, indicate the presence of micropores [19]). These values in magnitude were: 0.595, 0.510 and 0.0784 cm$^3$/g for the untreated, acid activated and heat activated red muds. This also confirms the BJH results.

Table 4. Summary of textural properties of raw and modified red mud samples.

<table>
<thead>
<tr>
<th>RM</th>
<th>BET surface area (m$^2$/g)</th>
<th>2.25M RMA</th>
<th>2M RMAC</th>
</tr>
</thead>
<tbody>
<tr>
<td>RM</td>
<td>13.15</td>
<td>23.80</td>
<td>26.46</td>
</tr>
<tr>
<td>External surface area (m$^2$/g)</td>
<td>14.88</td>
<td>25.21</td>
<td>26.05</td>
</tr>
<tr>
<td>Micropore surface area (m$^2$/g)</td>
<td>*</td>
<td>*</td>
<td>0.4166</td>
</tr>
<tr>
<td>Micropore volume (cm$^3$/g)</td>
<td>0.000920</td>
<td>0.000789</td>
<td>0.000121</td>
</tr>
<tr>
<td>Diameter range (nm)</td>
<td>1.7-300.0</td>
<td>1.7-300.0</td>
<td>1.7-300.0</td>
</tr>
<tr>
<td>Total pore volume (cm$^3$/g)</td>
<td>0.063</td>
<td>0.109</td>
<td>0.111</td>
</tr>
<tr>
<td>Average pore width (nm)</td>
<td>19.04</td>
<td>18.38</td>
<td>16.85</td>
</tr>
</tbody>
</table>

*The micropore area is not reported because the calculated external surface area is larger than the total surface area.

The porosity and surface area analyses show that the textural properties of raw and activated Minim-Martap red mud increases with the activation methods used. These increases are due the dissolution of some red mud components and elimination of several impurities. Huang et al. [35] and Wang et al. [36], reported similar trends in their studies when they activated red mud by acid, combined acid and heat, and heat treatment.

4. Conclusions

Results of this study shows that red mud can be modified depending on its subsequent use. The choice of the modification method however, depends strongly on the conditions of the parameter(s). For example modifying red mud with acid decreases its quantity more than when modified by heat treatment. However, acid modification increase alumina content but decreases that of iron. Conversely, heat treatment increase iron content but reduces alumina content. Under both conditions (acid and heat), the Si and Ti content show increases with significant increase when acid is used than the case of heat (Ti increasing from about 3% to over 17% with 4 mol/L acid). Also acid treatment does not modify the mineral composition of red mud but combined acid and heat treatment of red mud at 900°C significantly modifies the mineral composition with the elimination of most aluminum minerals and leaving a mainly iron mineral structure. The acid treated red mud and combined acid and heat treated red mud show increase porosity and surface area with combined acid and heat treated red mud having higher porosity and surface area. Modified and non-modified red mud contain mesopores > macropores > micropores. Combined acid and heat treated red mud shows high thermal stability than untreated and acid treated red muds which are thermally very not stable up to about 700°C.

These results show that the harmful red mud can be converted to useful products with improved properties for different technological applications. This can contribute in an increase in utilization of red mud, thereby reducing its negative environmental impact and creating economic benefits.

Competing Interests

The authors declare that they have no competing interests.

Acknowledgments

This work was supported by the Department of Applied Chemistry, National Advanced School of Agro-Industrial Sciences, University of Ngaoundere, Cameroon.

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