Removal of Phosphate Using Red Mud: An Environmentally Hazardous Waste By-Product of Alumina Industry

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The industrial waste, bauxite residue generated in the Bayer chemical process of alumina production, commonly known as red mud (RM) has been used as the adsorbent for selective removal of phosphate in aqueous solutions. RM collected from the storage area of alumina industry was characterized by chemical analysis and physical methods such as BET surface area, Scanning Electron Microscopy (SEM), particle size analysis, and X-ray diffraction (XRD) methods. Among the various red mud samples (0.2–200 μ) studied, the samples treated with 1 M HCl for 2 h were found better for the selective adsorption of phosphate in comparison with untreated and heat treated RM samples. The presence of phosphate in the aqueous samples collected after adsorption studies with red mud was determined by standard spectrophotometric procedure using ammonium molybdate and ascorbic acid in nitrate medium at λmax 880 nm. The studies reported significant adsorption of phosphate on acid treated red mud in comparison with adsorption of phosphate on untreated and heat treated red mud, respectively. The adsorption of phosphate on raw red mud and activated red mud was further investigated with respect to stirring time, pH of the solution, dose of adsorbent, and varying phosphate concentration. Acid treated RM is observed as an efficient and cost-effective adsorbent for selective removal of phosphate in aqueous solutions.

1. Introduction

Phosphorus as phosphate ions is one of the important soil elements for living organisms. It is an essential element for plant growth present in most of the biological tissues. However, an excess level of phosphate in water bodies helps the development of photosynthetic algae and toxic cyanobacteria. The industrial growth, household applications, and advances in agricultural research and crop development inadvertently cause the phosphate pollution in the aqueous bodies. In view of its potentially hazardous nature, several methods have been developed to remove excessive phosphate from natural and waste water sources [1–8]. That includes biological removal, precipitation method, ion exchange methods using zeolites, and metal oxides [9–12]. The biological method is of low cost but not effective due to variability in temperature of waste water that would make the execution of this process inaccessible for waste water treatment. Chemical treatments such as alum, lime, and ferric chloride are common precipitants used for phosphate removal but this is cost-ineffective and sludge formation makes this as a tedious method for waste water treatments. Most commercial systems currently use activated carbon as sorbent for removal of dyes in waste water because of excellent adsorption ability. However, the involved high cost restricts its widespread use. In the recent years, considerable attention has been paid based on economic and environmental concern to the study of using different types of low cost sorbents such as alum sludge, red mud (RM), fly ash (FA) [8–14], and other waste materials as alternative adsorbents for sorption [15] of phosphate from water.

In the present study, we have used red mud as an effective adsorbent for the removal of phosphates from aqueous solution. Bauxite residue, red mud, is the major solid waste produced during alkaline leaching of bauxite in the Bayer process. It is known to be a potential toxic waste causing serious
environmental problems due to high alkalinity with pH 10–12 and bulk volume. Redness of red mud is mainly due to the presence of iron impurities and chemical analysis of red mud shows silica, aluminium, iron, calcium, titanium, and sodium as major components and other elements such as K, Cr, V, Ba, Cu, Mn, Pb, Zn, P, F, S, and As as minor components depending on origin of bauxite mines. For every tonne of alumina produced, approximately 1 to 2 tonnes (dry weight) of red mud residue are generated [8]. Because of the increasing demand for alumina during economic development, more and more RM is being produced. It is estimated that about 90 million tonnes of RM is produced annually on earth, while this is expected to increase by several times in the near future. Due to the alkaline nature and high volume of red mud, it is a serious ecological threat and proper disposal of the waste red mud presents a huge challenge for alumina industries. Many attempts have been made over past years to find some practical applications for red mud such as an additive pigment for mortar and concrete [16, 17] and surface treatment of carbon steel and many investigations have also been extended to develop red mud as an adsorbent to remove arsenate [18–20], toxic heavy metals [21–28], dyes [29, 30], methylene blue [31, 32], and phenols [31–35].

For phosphate removal, some investigators have used either acid or heat treated RM as adsorbent [2, 8]. Shiao and Akashi [1] reported adsorption of phosphate from aqueous solution using HCl activated red mud and achieved 80–90% removal efficiency. Studies have also been reported by Aitundogan and Tumen [4] in which the sample prepared by using red mud stirred with HCl for 2h as well as another sample prepared by heating RM at 700°C for 2h results in the maximum removal of phosphate (99% removal of phosphate). It is seen that acid or heat treatment could produce an effect on phosphate adsorption depending on the material and treatment conditions.

In this paper, we report the spectroscopic characterization and investigation of the adsorption studies on raw and activated red mud for phosphate removal from aqueous solution. We have investigated the effect on morphology and adsorptive behaviour of the modified red mud by heat and acid treatment. This is done with the help of BET surface analyser, particle size analyser, X-ray diffraction, and Scanning Electron Microscopy.

2. Experimental

2.1. Materials. The raw red mud sample used in the present study was provided by the National Aluminium Company Ltd., Damanjodi, Orissa, India. The chemical composition of raw material was determined by the wet chemical analysis method and chemical compositions of the red mud based on dry weight are Fe₂O₃ (54.6%), Al₂O₃ (16.44%), L.O.I (11.67%), TiO₂ (6.82%), SiO₂ (4.24%), Na₂O (2.76%), and CaO (1.99%). This raw red mud was washed several times with deionised water and hereafter referred to as red mud (RM). Initially these RM samples were calcined in a muffle furnace at 700°C in air for 2h. This is referred to as red mud heated (RMH) sample. The other RM samples were mixed with 1 M HCl for 24 h at liquid/solid ratio of 20 mL/g. These samples are referred to as acid treated RM-HCl (RMA). After acid treatment, the RMA was washed once again with deionised water and dried overnight at 105°C. The acid treated sample of RMA was then calcined in muffle furnace at 700°C in air for 2h. This is referred to as red mud acid heated (RMAH) sample. The resultant solid samples of RMAH were then allowed to cool down overnight in the furnace.

The chemicals and reagent used in this research including hydrochloric acid (HCl), sulphuric acid (H₂SO₄), ascorbic acid (C₆H₇O₆), sodium hydroxide (NaOH), potassium dihydrogen phosphate (KH₂PO₄), ammonium molybdate tetrahydrate [(NH₄)₆Mo₇O₂₄·4H₂O], and potassium antimonyl tartrate [K(SbO)C₆H₄O₆·1/2H₂O] were of analytical grade, and all the solutions were prepared by using deionised water as solvent.

2.2. Characterization. The prepared red mud samples, RM, RMH, RMA, and RMAH, were characterized using BET surface analyser, particle size analyser, X-ray diffraction, and Scanning Electron Microscopy. The surface area was obtained by nitrogen-desorption at the liquid nitrogen temperature (−196°C) using ASAP Micromeritics 2020. All samples were degassed at 200°C for 2 h before analysis. The BET equation was applied to determine the surface areas of RM, RMH, RMA, and RMAH.

X-ray diffraction (XRD) patterns of all samples were measured using an Xpert PRO, Pan Analytical Diffractometer with Cu Kα radiations generated at 40 kV, 30 mA. Scattering angles were ranged from 20 to 60°, with a scanning rate of 20° per minute.

Particle sizes of RM, RMH, RMA, and RMAH samples were analysed using Micromeritics Saturn DigSizer II 5205 particle size analyser. This data will help us in determining the percentage of particles belonging to particular size category that is attributed to surface area.

Microstructure and surface morphology of the adsorbent samples were characterized by a JSM IT-300 field emission Scanning Electron Microscopy (SEM) with an accelerating voltage of 15 kV at a magnification of 1 × 10000.

The thermogravimetric (TG) analysis of RM, RMH, RMA, and RMAH samples was carried out in N₂ atmosphere using NETSZ 402PC heat analyser at the heating rate of 20°C/min ranging from 30°C to 1000°C.

The results obtained from SEM analyses and TGA graphs suggest that some significant phase transformations take place on acid or heat treatment, which is discussed at later stages in the paper.

2.3. Adsorption Studies. Adsorption experiments were conducted by stirring the 20 mL of varying concentrations of phosphate solution ranging from 50 mg/L to 1000 mg/L (prepared by KH₂PO₄, phosphate concentration) with 0.1 g of adsorbent in the capped conical flasks at 30°C ± 1°C for 8 hours. After equilibrium, the clear filtered solutions were obtained using the pulp filtration method and we analysed the phosphate concentration by following the modified single solution method as given by Murphy and Riley [36] and in State Environmental Protection Administration [37]. A calibration curve was obtained by measuring the absorbance...
of phosphate concentration ranging from 0.02 to 0.2 mg of PO$_4^{2-}$/L as the basis to determine the concentration of unknown sample with the help of UV-Visible range Shimadzu UV-1800 spectrophotometer at 882 nm wavelengths. The amount of phosphate adsorbed per unit mass of adsorbent was calculated by

$$X = \frac{(C_i - C_f) \cdot V}{m},$$  \hspace{1cm} (1)

where $X$ is the concentration of the phosphate adsorbed; $C_i$ and $C_f$ are the initial and final concentrations of the phosphate in mg/L. $V$ is the volume in litre of solution taken in a conical flask for stirring whereas $m$ is the weight of adsorbent.

Now, in order to determine the mechanism of adsorption of phosphate on RM, RMH, RMA, and RMAH, Langmuir and Freundlich isotherms were applied to describe the equilibrium adsorption of solute from solution at constant temperature. Langmuir isotherm assumes the sorption process at specific homogeneous site for monolayer adsorption. The Langmuir isotherm can be expressed as

$$X = \frac{kX_mC_{eq}}{1 + kC_{eq}},$$ \hspace{1cm} (2)

where $X$ is the adsorbed amount of the solute in mg of PO$_4^{2-}$/g of adsorbent, $C_{eq}$ is the equilibrium concentration of the solution in mg of PO$_4^{2-}$/L, $X_m$ is the monolayer adsorption capacity (mg/g), and $k$ is a constant related to the free energy of adsorption. The Freundlich adsorption isotherm is an empirical relation used to describe heterogeneous system which can be expressed as

$$Q = KC_{eq}^{1/n},$$ \hspace{1cm} (3)

where $K$ is a constant that indicates the extent of the adsorption and $n$ is a constant that indicates the nonlinearity between concentration and the extent of adsorption at the experimental pressure as given by Dursun et al. [8].

### 3. Results and Discussion

#### 3.1. Chemical Analysis of Red Mud

Chemical analysis of raw and activated red mud shows the distinct variation in their chemical composition (Table 1). Variation in chemical composition of various samples of RM is attributed to their adsorption capacity. Specifically, Loss On Ignition (LOI) data show significant decrease in LOI in case of heat action that is reflected in increased concentration of iron oxides. This is attributed to the increased adsorption capacity of heat activated red mud.

The acid treated RMA shows higher adsorption compared to RMH and RMAH. It is also observed from experimental results (Figure 1) that the adsorption capacity of raw red mud can be improved significantly by acid treatment.

#### 3.2. Effect of Stirring Time

The effect of stirring time on the removal of phosphate was studied by varying the time of stirring from 2 hr to 10 hr for the optimum equilibrating time. Results (Figure 2) show that the equilibrium in adsorption of phosphate is attained at about 8 hr of stirring time in case of RM whereas, in case of activated red mud, variation in stirring time does not show any significant effect after 2 hr of stirring. Therefore, for the sake of self-sufficiency, the optimum stirring time of 8 hr has been chosen for all investigations throughout the study.

#### 3.3. Effect of Phosphate Concentration

Adsorption studies were carried out on a fixed weight of red mud with varying phosphate concentrations from 50 to 500 mg/L. It is observed from our studies that the phosphate removal efficiency increases with increasing phosphate concentrations and rises to maxima up to around 250 to 300 mg/L (Figure 1). Further,
it is observed that phosphate removal efficiency does not show any significant change after phosphate concentration of 300 mg/L. Zhao et al. [5] have shown that such behaviour is anticipated due to the buffering properties of phosphate compounds.

3.4. Effect of pH. The effect of variation in pH on the removal of phosphate has also been studied in the range from 1.0 to 10 pH of phosphate solution using 0.1% HCl and 0.1% NaOH solutions that gave the optimum value of pH for our investigations. It is apparent from Figure 3 that the phosphate removal shows maxima at pH 2.0 of solution and then decreases with further increase in pH from 3.0 to 10.0. This trend has no exceptions irrespective of RM, RMA, RMH, and RMAH. Significant enhancement of adsorption was achieved at about pH 2 (Figure 3). This is attributed to that fact that, at higher pH, surface of the red mud carries more negative charges which may serve to increase the repulsion of negatively charged phosphate species in solution and consequently results in lower RM adsorption capacity. These results are in good agreement with the available literature [6].

3.5. Thermal Analysis. Thermal gravimetric analysis of RM (Figure 4) shows a continuous weight loss distributed over the range of 30–900°C. Composition of RM mainly contains crystalline hematite (Fe$_2$O$_3$), boehmite (gamma-Al$_2$O$_3$·H$_2$O), quartz (SiO$_2$) with minor percentage of calcite (CaCO$_3$), gibbsite (Al(OH)$_3$), and other minor components like Na$_2$O, TiO$_2$, K$_2$O, and so forth [1]. It is observed from TGA (Figure 4) that the total weight loss is 26.79%, 19%, and about 2% in cases of RM, RMA, and RMH, respectively.

Further, the corresponding total weight loss in case of RM (Figure 5(a)) can be classified into three major stages, namely, the initial weight loss of 15.68% between 30 and 220°C. This is attributed to the evaporation of moisture adsorbed on the surface of powdered RM. The subsequent weight loss of 6.47% in the range from 220 to 400°C is due to the release of constituent water from crystals in the minerals, such as boehmite and gibbsite, and the final weight loss of 4.63% may be attributed to the decomposition of calcite and whewellite [2] and after that almost remains constant. In case of RMA (Figure 5(b)), the major loss appears to be in the range of 200 to 500°C that may be attributed to the loss of salts that might have formed during neutralization reaction of acid treatment along with hydrated water.

3.6. XRD Analysis. The crystalline phases of red mud samples were analysed by XRD patterns. In RM (Figure 6(a)), the major phases are gibbsite, hematite, goethite, sodalities, and calcite. The XRD patterns show remarkable changes between RM, RMA, RMH, and RMAH (Figures 6(a), 6(b), 6(c), and
Figure 5: (a) Derivative thermogravimetric (DTG) analysis of RM. (b) Derivative thermogravimetric (DTG) analysis of RMA.

Figure 6: (a) XRD pattern of RM. (b) XRD pattern of RMA. (c) XRD pattern of RMH. (d) XRD pattern of RMAH.

6(d)) which indicate the occurrence of phase transformations. After acid treatment, all the other phases disappear except gibbsite and hematite (Figure 6(b)). Also acid treatment shows a generation of new phase of rutilite which may be attributed to the decomposition of calcite, whereas heating effect on both samples (RMH and RMAH) cause significant enhancement in intensity of hematite (Figures 6(c) and 6(d)). This is in accordance with the fact that heating process destroys crystalline phase of RM and generates amorphous phase.

3.7. BET Surface Area Analysis. The BET surface area analyses (Table 2) reveal that the RM has a specific surface area of 20.82 m²/g whereas heating effect on RM (i.e., RMH) shows a greatest enhancement in the specific surface area up to more than double of its original surface area. On treating RM
Table 2: BET surface area analysis of various red mud samples.

<table>
<thead>
<tr>
<th>Sr. number</th>
<th>Samples</th>
<th>Area (m²/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>RM (raw red mud)</td>
<td>20.89</td>
</tr>
<tr>
<td>2</td>
<td>RMH (heated red mud at 700°C for two hours)</td>
<td>42.01</td>
</tr>
<tr>
<td>3</td>
<td>RMA (acid treated red mud)</td>
<td>28.63</td>
</tr>
<tr>
<td>4</td>
<td>RMAH (red mud treated with acid and then heated at 700°C for two hours)</td>
<td>37.34</td>
</tr>
</tbody>
</table>

Figure 7: (a) SEM image of RM. (b) SEM image of RMH. (c) SEM image of RMA. (d) SEM image of RMAH.

3.8. Morphological Analysis. Surface morphological studies of all the RM samples are carried out using SEM images (Figures 7(a), 7(b), 7(c), and 7(d)). Surface of the RM (Figure 7(a)) is relatively smooth and flat in contrast to that of RMH (Figure 7(b)) which gives a clear evidence of generation of new porous surface due to removal of moisture and constituent water from crystals in the minerals, such as boehmite and gibbsite, and due to the decomposition of calcite and whewellite. This is in strong agreement with the BET surface area analysis wherein surface area has been sharply increased almost more than double the original surface area. Acid treatment usually leads to formation of additional new cavities during the acidification process by strong acid which significantly enhances the surface area. The SEM image (Figure 7(c)) of acid treated samples shows many such new cavities along with coarse exterior due to some acid soluble salts. The generation of new surface area by acidification was observed from the difference between the SEM images of Figures 7(a) and 7(c). It is also interesting to see the effect of heat on acid treated sample (Figure 7(d)) because surface area increases more than 85% from the original surface area which may be attributed to the decomposition of deposited acid soluble fractions like gibbsite, sodalities, goethite, calcite, and some other acid soluble salts. Moreover, on comparing the surface morphology of RMH and RMAH, it is observed prominently that RMH shows...
higher percentage of cavities leading to higher surface area whereas RMAH shows higher porosity which, however, may not give maximum surface area.

3.9. Adsorption Studies. The sorption studies of phosphate ions on the surface of RM, RMH, RMA, and RMAH were carried out in the concentration range of 50 to 500 mg P/L. In order to study the mechanism of phosphate adsorption, Langmuir and Freundlich isotherms were applied. The Langmuir adsorption isotherm model represents one of the first theoretical treatments of nonlinear sorption and suggests that the uptake occurs on a homogenous surface by monolayer sorption without interaction between the adsorbed molecules. The Langmuir adsorption isotherm is often used to describe the maximum adsorption capacity of an adsorbent. The phosphate adsorption isotherms (Figure 1) suggest that phosphate adsorption increases with the increases in its concentration up to certain optimum value and reaches a plateau region where no significant enhancement in the adsorption can be obtained.

Among all the four samples of red mud, RMA, in spite of having lower surface area, has achieved adsorption up to 205.1 mg P/g of RMA, as compared to RMH and RMAH wherein the maximum adsorption is up to 190.8 mg of P/g RMH and 175.6 mg of P/g of RMAH, respectively, whereas it is up to 125.9 mg of P/g of RM in case of raw RM. This is due to the fact that phosphate can also form chemisorbed complexes by ligand exchange reactions with hydroxide as suggested by Wang et al. [14]. The hydrolysis of Fe$^{+3}$ forms Fe-OH on the goethite crystals and then phosphate replaces the hydroxyl group and forms a bridge between adjacent Fe$^{+3}$ as suggested by Atkinson et al. [38]. Also Kim et al. [39] have suggested that the phosphate ions can bind to Fe$^{+3}$ through two P-O-Fe linkages. Thus, chemisorption due to ion exchange is also an important aspect for the adsorption of phosphate on RMA.

The equilibrium data were analysed in light of the adsorption isotherm models. The data were then fitted to the Langmuir equation as

\[
\frac{C_{eq}}{X} = \frac{1}{X_m} + \frac{1}{kX_m} \frac{1}{C_{eq}},
\]

where \(X\) indicates the amount of phosphate adsorbed per unit mass of adsorbent, \(C_{eq}\) is the equilibrium phosphate concentration in solution, \(X_m\) is the monolayer adsorption, and \(k\) is the adsorption constant related to free surface energy. The experimental data of phosphate adsorption on RMA (Figure 8) is plotted using Langmuir equation (4).

The correlation coefficient is found to be 0.99. All the results show that the Langmuir equation for adsorption fits well. The Langmuir parameters for all the RM samples are summarised in Table 3.

<table>
<thead>
<tr>
<th>S. number</th>
<th>Sample</th>
<th>(X) (mg P/g RM)</th>
<th>(k)</th>
<th>(R^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>RM</td>
<td>127.64</td>
<td>9.45</td>
<td>0.95</td>
</tr>
<tr>
<td>2</td>
<td>RMA</td>
<td>205.13</td>
<td>12.65</td>
<td>0.99</td>
</tr>
<tr>
<td>3</td>
<td>RMH</td>
<td>190.79</td>
<td>9.20</td>
<td>0.97</td>
</tr>
<tr>
<td>4</td>
<td>RMAH</td>
<td>175.64</td>
<td>17.39</td>
<td>0.97</td>
</tr>
</tbody>
</table>

Figure 8: Langmuir plot of phosphate adsorption on acid treated activated red mud (RMA).

### Competing Interests

The authors declare that there is no conflict of interests regarding the publication of this article.

### Acknowledgments

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