Iron-loaded Sphagnum moss extract residue for phosphate removal

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Abstract

Sphagnum moss extract residue (SMER), obtained after pressurized hot water extraction, was modified with Fe(III) and investigated for phosphate sorption. Although moss extract contains value-added compounds, SMER is considered to be waste until suitable uses can be developed. The effect of modification conditions were investigated, i.e. different initial Fe(III) concentrations (0.024, 0.048 and 0.072 mol/L Fe\textsuperscript{3+}) and modification pH values (5, 7 and 9). A modification pH of 5 and the highest initial Fe(III) concentration (0.072 mol/L Fe\textsuperscript{3+}) resulted in the highest phosphate removal efficiency, and thus was selected for further study. The removal efficiency was found to decreased with increasing pH in the range of 3-9. Maximum removal efficiency (82\%) for phosphate sorption was observed at pH 3 after 24 h contact time (dosage 2 g/L, initial concentration 15 mg P/L). With increased contact time, the phosphate removal efficiency improved and reached equilibrium within 48 h. The Elovich model was found to provide the best fit to the kinetic data. A capacity of 9–13 mg P/g was obtained with a 24-hour contact time at pH 4. A
good fit was achieved with the Redlich–Peterson equation. FTIR analysis confirmed that carboxylic acid groups were involved in the modification process. X-ray diffraction analyses showed that amorphous two-line ferrihydrite was precipitated onto SMER, which was supported by X-ray photoelectron spectroscopy analyses.

Keywords: Hot water extraction, Iron(III) modification, Surface composition, Sorption, Ferrihydrite

1. Introduction

Phosphorus is a very important nutrient for living creatures. Generally, the content of phosphorus in natural water bodies is assumed to be extremely low. However, over the past decade, the discharge of phosphorus has increased significantly through agricultural wastes, municipal wastewater, expansion of fossil fuel combustion and the use of fertilisers. Phosphate in wastewaters has drawn a lot of attention in recent years. Various conventional technologies are available for phosphate removal such as membrane filtration, coagulation and flocculation, and biological assimilation. Nevertheless, problems of high cost, sludge and waste disposal, and operation difficulties have always posed a major challenge for these conventional methods (Gupta and Ali, 2012; Xu et al., 2016).

As one of the wastewater treatment technologies with the most potential, sorption has received a lot of attention due to its low cost and ease of operation (Ali et al., 2012; Adegoke et al., 2013; Ali et al., 2016a; Ali et al., 2016b; Dehghani et al., 2016; Xu et al., 2016). Numerous virgin or activated low-cost sorbents such as agricultural products (wheat straw, rice husk, cotton seed hulls and sugar beet pulp), natural residues (sawdust,
bark, peat and peat moss), industrial wastes (fly ash, red mud, petroleum wastes, scrap tyres and blast furnace slag), marine biomasses (sea weed and algae) and some other biological materials (fungi, yeast, chitosan and bacterial biomass) have been investigated (Ali et al., 2012; Ali et al., 2014; Bhatnagar and Minocha, 2006; Crini, 2006; Khan et al., 2011).

Sphagnum moss is a plant that grows in moist and wet areas, whereas peat moss is partially decomposed plant material (e.g. from Sphagnum moss). These materials are complex biomaterials comprising lignin-like substances and cellulose as the major components, as well as humic substances. Sphagnum moss is widely available due to its abundance; according to Charman (2002), globally peatlands cover an area of some four million square metres, with a renewing rate of 50,000 ha/a. The high content of oxygen-containing functional groups, large specific surface area and high porosity enable Sphagnum moss and peat moss to uptake various pollutants (mainly metal cations) through ion exchange, complexation and other reactions (Babel and Kurniawan, 2003; Gardea-Torresdey et al., 1996).

As Sphagnum moss contains various high-value bioactive constituents with antioxidative (Tarnawski et al., 2006), antimicrobial (Baas et al., 2000) or otherwise medically interesting properties (Gamenara et al., 2001; Yamada et al., 2007), it is feasible to employ a cascade approach starting with extraction, after which the biomass residue remains useful for the production of biosorbent for wastewater treatment. This study focuses on the utilization of Sphagnum moss extract residue (SMER) as a novel support material, onto which iron compounds would be introduced to improve the phosphate
sorption capacity. A further benefit of using the residue in water treatment is its low leaching characteristics (e.g. low humic content).

In general, the sorption ability of biomaterials as such is inadequate for anionic pollutants (Ali et al., 2015; Xu et al., 2016). Therefore, a variety of modification methods have been developed, among which metal-impregnated biomass has attracted a great deal of interest. Commonly used metal ions include Zr(IV), Al(III) and Fe(III). Iron-modified biosorbents have been used to remove various anionic species such as As(III) and As(V) (Aryal et al., 2010), Cr(VI) (Li et al., 2004) and phosphate. Some studies on Fe(III)-loaded materials for phosphate removal are listed in Table 1. As can be seen from the table, good sorption capacities were obtained. However, very little is known about the optimal iron modifications for biosorbents and only few articles have studied the type of iron compounds loaded on the biosorbents. Modification pH, for example, has a significant effect on the composition and structure of iron oxide (Adegoke et al., 2013). Further research on iron modification is thus of great importance.

Table 1. Phosphate sorption capacities of some iron-loaded materials

<table>
<thead>
<tr>
<th>Biomaterial</th>
<th>Sorption Capacity (mg P/g)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe(III)-loaded orange waste</td>
<td>13.9</td>
<td>Biswas et al. (2007)</td>
</tr>
<tr>
<td>Fe(III)-loaded collagen fibre</td>
<td>34.1</td>
<td>Liao et al. (2006)</td>
</tr>
<tr>
<td>Fe(III)-loaded cellulose nanofibres</td>
<td>46.6</td>
<td>Cui et al. (2016)</td>
</tr>
<tr>
<td>Fe(III)-loaded okara</td>
<td>4.8</td>
<td>Nguyen et al. (2013)</td>
</tr>
<tr>
<td>Fe(III)-loaded coir pith</td>
<td>23.1</td>
<td>Krishnan &amp; Haridas (2008)</td>
</tr>
<tr>
<td>Fe(III)-loaded wood particles</td>
<td>2.8</td>
<td>Eberhardt and Min (2008)</td>
</tr>
<tr>
<td>Anionic polymer and Fe(III)-loaded wood particles</td>
<td>4.1</td>
<td>Eberhardt and Min (2008)</td>
</tr>
</tbody>
</table>
In view of the above, the present work investigated the optimum modification process for iron-modified SMER. The modification pH and initial concentration of the Fe(III) solution were studied. Subsequently, phosphate sorption tests were conducted to determine the capacity of the product, the effect of contact time and the optimal pH. X-ray diffraction (XRD), Fourier transform infrared spectroscopy (FTIR) and X-ray photoelectron spectroscopy (XPS) were used to study the characteristics of Fe(III)-loaded SMER, providing new knowledge about both the iron-binding and the phosphate-binding mechanisms.

2. Materials and methods

2.1 Raw materials and chemicals

SMER was provided by the Natural Resources Institute of Finland. *Sphagnum magellanicum* moss was collected from Pehkeensuo (a small sedge bog) in Utajärvi, Finland. Sphagnum moss was treated by pressurized hot water extraction (15 bar) using a 300-litre extraction unit (Kilpeläinen et al., 2014). Sphagnum moss (18 kg, 7.8% dry matter content) was loaded into the extraction chamber. Preheated water (180°C) was pumped through the extraction chamber at a flow rate of 20 kg/min for one hour. Then the extract and residue (SMER) were cooled down and the SMER was frozen and stored. The extract was used for other purposes. In the modification experiment, melted SMER was dried at 80°C for 24 h, then ground and sieved to retain the 250–500 µm fraction.

Na$_3$PO$_4$ (Sigma-Aldrich) was used to prepare different concentrations of phosphate solution. Diverse iron concentration solutions were prepared by dissolving FeCl$_3$·6H$_2$O (Merck) in pure Milli-Q water (Merck Millipore). HCl (Merck) and NaOH (Sigma-Aldrich) were used for pH adjustment. LCK-350 phosphate cuvettes (Hach-Lange,
Germany) were used for the analysis of phosphate after the sorption test. All the chemicals used were of analytical quality.

2.2 Biosorbent characterization

Five samples were prepared for characterization: virgin SMER, iron-modified SMER at different modification pH values (5, 7 and 9) with the highest iron concentration (0.072 mol/L Fe$^{3+}$), and phosphate-treated iron-modified SMER (modification pH 5 with the highest iron concentration). Samples were characterized using XRD, FTIR and XPS. One gram of each sample was ground to fine powder with a hand mortar and then collected for XRD and FTIR analyses. A small fraction of the samples (as such, without grinding) were used for XPS.

XRD analyses were conducted using a Rigaku Smartlab diffractometer and a Co lamp (40 kV, 135 mA). The samples were measured at room temperature using Bragg-Brentano para-focusing geometry (300 mm goniometer). The speed of acquisition was 3 deg/min at 0.02 deg/step. FTIR spectra were measured using a Bruker Vertex V80 vacuum FTIR spectrometer, which was connected to a computer containing the OPUS program to display the recorded spectra. The FTIR spectra were recorded in the 400-4000 cm$^{-1}$ wave number region. XPS was carried out with a Thermo Fisher Scientific ESCALAB 250Xi and using a monochromatic Al K$\alpha$ source (1486.6 eV). Wide-scan spectra were recorded in steps of 1 eV and a pass energy of 150 eV, and high-resolution spectra with steps of 0.1 eV and a pass energy of 20 eV. The charge correction was performed by setting the binding energy of adventitious carbon to 284.8 eV.

2.3 Biomass modification and batch sorption test
A full factorial design was chosen to investigate the biomass modification. The two factors investigated were initial iron concentration (0.024, 0.048 and 0.072 mol/L Fe\(^{3+}\)) and modification pH (5, 7 and 9). Phosphate removal efficiency was selected as the response. The results were evaluated by computing the coefficients using multiple linear regression (Modde 12.0 software, Umetrics AB). The coefficient was considered significant if the p value was below 0.05.

The experiment was conducted at room temperature. Two repeats were conducted for each iron concentration under the same modification pH. The modification process was conducted as follows: 1 g SMER was treated with 50 mL Fe(III) solution. The modification pH was adjusted to the desired value (5, 7 or 9). Next, the mixture was agitated by magnetic stirring at room temperature (20 ± 3 °C) for 6 h. After that, the modified SMER was separated from the solution by centrifugation (Jouan C4.12, 10 minutes, 2500 rpm). The sample was washed several times with water until the pH reached neutral and the solution became clear and colourless. Then the final product was dried at 80°C for 26 h.

Fe(III)-treated and virgin SMER were then used for the removal of phosphate from aqueous solutions. Samples were weighed (0.1 g) and transferred into polypropylene bottles, followed by the addition of 50 mL of phosphate solution with a concentration of 15 mg P/L. The pH of the phosphate solution was pre-adjusted to 5.4–5.6. The batch experiment for each product was done in duplicate. The bottles were then shaken for 24 h at room temperature at a speed of 20 rpm/min. After two hours, pH readjustment was performed for each sample by adding HCl or NaOH to prevent pH change, thus ensuring the accuracy of the experiments. After the sorption test, the sample was centrifuged (10
min, 2500 rpm) and filtered with a 0.45 µm membrane, then analysed for residual phosphate content. On the basis of the results, the SMER modified at pH 5 and with the highest Fe concentration was selected for further study (capacity, contact time and pH).

2.4 Phosphate removal capacity

The phosphate sorption capacity was determined for two batches (batches 1 and 2) of modified SMER (pH 5, Fe 0.072 mol/L). Sorption tests were performed with a similar procedure as that mentioned in section 2.3. Different initial concentrations of phosphate solution (10–175 mg P/L) were prepared and the pH of the solution was pre-adjusted to 4. Other parameters, i.e. the contact time (24 h) and biosorbent dosage (2 g/L), were kept constant. All the experiments were done in triplicate. Sorption capacity was calculated using the following equation (1):

$$q_e = \frac{(C_0-C_e)V}{m} \quad (1)$$

where $q_e$ is the phosphate sorption capacity (mg/g); $C_0$ is the concentration of phosphate ions in the initial solution (mg/L); $C_e$ is the residual phosphate concentration in solution after the sorption test (mg/L); $V$ is the volume of the solution (L); and $m$ is the weight of the biosorbent (g). The non-linear Langmuir, Freundlich and Redlich-Peterson isotherms were used to evaluate the experimental adsorption data (supplementary material).

2.5 Effect of pH of aqueous media

Batch experiments to test the effect of pH were conducted at seven different pH levels: 3, 4, 5, 6, 7, 8 and 9. Other parameters, i.e. the initial concentration of the phosphate solution (15 mg P/L), contact time (24 h) and biosorbent dosage (2 g/L), were kept constant. All
the tests were performed in duplicate, following a similar batch procedure as that described in section 2.3.

2.6 Effect of contact time

The effect of different contact times (10 mins, 30 mins, 1 h, 6 h, 24 h, 48 h and 72 h) was investigated by using 15 mg P/L phosphate solution as the initial phosphate concentration with a pH value of 4 at room temperature. The biosorbent dosage was 2 g/L. pH readjustment was performed after two hours of shaking for the samples with contact times of 6, 24, 48 and 72 h. All the experiments were done in duplicate, following a similar batch procedure as that described in section 2.3. Three kinetic models (pseudo-first-order, pseudo-second-order and Elovich) were used to analyse the kinetic data (in the supplementary material). The non-linear method was applied to determine the parameters of the kinetic models.

3. Results and discussion

3.1 Biosorbent characterization

3.1.1 XRD analysis

The XRD profiles of virgin SMER, iron-modified SMER (0.072 mol/L Fe\textsuperscript{3+} concentration) at three different modification pH values (5, 7 and 9), and phosphate-treated modified SMER (modification pH 5) are shown in Fig. 1. Small peaks were observed in the 2θ range of 30–40° in the XRD pattern of virgin SMER, which disappeared after iron modification. In addition, the loaded iron was in the amorphous phase as indicated by the XRD pattern, hence amorphous iron oxyhydroxide existed in the modified SMER. Two broad peaks with a d-value of 0.26 nm (2θ ≈ 40°) and 0.15 nm
(2θ ≈75°) are characteristic of two-line ferrihydrite (Cornell and Schwertmann, 2003; Mathew et al., 2011). XRD analysis thus indicated that two-line ferrihydrite was loaded on the SMER and the iron was in trivalent state. Several formulae of ferrihydrite have been proposed in the past, but $5\text{Fe}_2\text{O}_3\cdot9\text{H}_2\text{O}$ is the most commonly used formula in the literature (Mallet et al., 2013).

Fig. 1 a single-column fitting image

Two-line ferrihydrite and six-line ferrihydrite are two commonly distinguished types of ferrihydrite. Lewis and Cardile (1989) demonstrated that acid hydrolysis of dilute Fe(NO$_3$)$_3$ solutions at room temperature resulted in six-line ferrihydrite, whereas two-line ferrihydrite was produced by quick basic hydrolysis. The modification conditions used in this study thus favour the formation of two-line ferrihydrite. The strong adsorptive capacity and large surface area of two-line ferrihydrite enable it to take up various water pollutants (Das et al., 2010). Ferrihydrite is the unstable phase of iron oxyhydroxide. Under certain conditions, it will transform slowly over time into more stable and more crystalline phases, i.e. hematite and goethite (Schwertmann et al., 1999). Hematite formation is enhanced close to the point of zero charge (p$\text{H}_{\text{PZC}}$) of two-line ferrihydrite or at higher temperature, whereas goethite is formed at pH values well above or below the p$\text{H}_{\text{PZC}}$.

3.1.2 FTIR

Fig. 2 shows the FTIR spectra of virgin and iron-modified SMER (pH 5 with 0.072 mol/L Fe$^{3+}$ concentration). Bands at 1730 and 1610 cm$^{-1}$ may be associated with C=O stretching vibrations. The band from 1650 to 1600 cm$^{-1}$ was attributed to the carbonyl bond from carboxylic groups (COO$^-$), while the band at 1730 cm$^{-1}$ refer to the carbonyl bond from
carboxylic acid compounds (–COOH) and esters (Bulgariu et al., 2011). After iron modification, a significant change was observed. The band from 1730 cm\(^{-1}\) became very weak and shifted to a lower wave number (1722 cm\(^{-1}\)). Conversely, the band at 1610 cm\(^{-1}\) was enhanced, which indicated that carboxylic acid groups had reacted with the iron compounds. This supports previous findings that carboxylic acid groups are responsible for the binding of iron on biomass (Aryal et al., 2010). The binding of Fe(III) ions to carboxylic acid groups is based on the ion exchange mechanism. Gardea-Torresdey et al. (1996) demonstrated that other functional groups such as phenol also bound Cu(II) to Sphagnum peat moss. After treating with phosphate, no obvious changes were observed in the intensities of carboxylic acid groups.

Fig. 2. a single-column fitting image

3.1.3 XPS analysis

The elemental compositions of virgin SMER, iron-modified SMER (0.072 mol/L Fe\(^{3+}\)) at three different modification pH values (5, 7 and 9), and phosphate-treated modified SMER (modification pH 5) are listed in Table 2. The corresponding Fe:O and O:C ratios were also calculated and included. Iron was successfully impregnated on SMER during the modification process (Table 2). The Fe:O ratio was 1:2.6 and 1:2.5 for modification pH values of 5 and 7, respectively. However, the ratio changed to 1:3.3 when the highest modification pH of 9 was used. The atomic percentage of iron on the surface of iron-modified SMER was 17.6% for pH 5 and 18.5% for pH 7, which were higher values than for iron-modified SMER in alkaline conditions (11.7%). This indicated that more iron could be loaded on SMER under lower pH conditions. As the modification process dramatically changed the chemical composition of the SMER surface, the ratio of O:C
also increased. The carbon content decreased while the oxygen content increased, but the changes were not so significant when alkaline modification (pH 9) was applied in comparison with modification pH values of 5 or 7.

Table 2. Surface composition (at.%) of virgin, iron-modified (0.072 mol/L Fe\(^{3+}\) concentration) and phosphate-treated iron-modified SMER (initial concentration of phosphate=15 mg P/L)

<table>
<thead>
<tr>
<th>Biosorbent type</th>
<th>pH</th>
<th>Atomic percentage of the elements (%)</th>
<th>C</th>
<th>Fe</th>
<th>O</th>
<th>N</th>
<th>P</th>
<th>Fe:O</th>
<th>O:C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Virgin SMER</td>
<td>-</td>
<td></td>
<td>74.0</td>
<td>-</td>
<td>25.0</td>
<td>1.1</td>
<td>-</td>
<td>-</td>
<td>1:3.0</td>
</tr>
<tr>
<td>Modified SMER</td>
<td>5</td>
<td></td>
<td>36.8</td>
<td>17.6</td>
<td>44.9</td>
<td>0.7</td>
<td>-</td>
<td>1:2.6</td>
<td>1:1.3</td>
</tr>
<tr>
<td>Modified SMER</td>
<td>7</td>
<td></td>
<td>34.2</td>
<td>18.5</td>
<td>46.4</td>
<td>0.9</td>
<td>-</td>
<td>1:2.5</td>
<td>1:1.4</td>
</tr>
<tr>
<td>Modified SMER</td>
<td>9</td>
<td></td>
<td>48.9</td>
<td>11.7</td>
<td>38.5</td>
<td>1.0</td>
<td>-</td>
<td>1:3.3</td>
<td>1:0.8</td>
</tr>
<tr>
<td>P-treated modified SMER</td>
<td>5</td>
<td></td>
<td>35.9</td>
<td>15.0</td>
<td>46.3</td>
<td>0.9</td>
<td>2.0</td>
<td>1:3.1</td>
<td>1:1.3</td>
</tr>
</tbody>
</table>

The O 1s spectra of virgin SMER, iron-modified SMER at three different modification pH values (5, 7 and 9), and phosphate-treated iron-modified SMER (modification pH 5) are shown in Fig. 3. The virgin SMER sample displayed a peak centred at 532.8 eV, which was assumed to be mainly the organic C–O in the SMER but there should also be C=O bonding on the lower BE side of that peak (Nguila Inari et al., 2006; Pereira et al., 2014). The O 1s spectra of iron-loaded SMER were fitted with three peaks: organic C–O bonding (532.9 eV ± 0.1 eV), Fe–OH bonding and C=O bonding (531.5 ± 0.2 eV), and Fe–O bonding (530.1 ± 0.2 eV) (Cui et al., 2016; Nguila Inari et al., 2006; Mallet et al., 2013; Popescu et al., 2009). After phosphate treatment, the O–P bonding obtained is assumed to be included in the middle peak along with Fe–OH and C=O bonding (531.5 eV, Fig. 3e) (Mallet et al., 2013).

Fig. 3. a 2-column fitting image
The shape of the Fe 2p spectra under different modification pH values was almost the same (Fig. S1), which supported the XRD data that the same iron compound, i.e. two-line ferrihydrite, was loaded on the SMER. The Fe 2p\textsubscript{3/2} and Fe 2p\textsubscript{1/2} peaks of the iron-loaded SMER samples had binding energies at 711.1±0.2 eV and 724.3±0.3 eV, which correspond to the binding energy of ferrihydrite (Frau et al., 2010; Mallet et al., 2013). However, it should be noted that some other iron compounds such as akaganéite (β-FeOOH) and goethite (α-FeOOH) also have similar binding energies (Ding et al., 2000; Missana et al., 2003; Zhang and Jia, 2014). After phosphate adsorption, slight shifts of the Fe 2p\textsubscript{3/2} and Fe 2p\textsubscript{1/2} peaks towards a higher binding energy (0.1−0.4 eV) were observed. The P 2p spectrum of phosphate-treated modified SMER (modification pH 5) displayed a peak at 133.6 eV (data not shown), which was assumed to be Fe−O−P bonding (Mallet et al., 2013). Therefore, XPS gave evidence of the formation of an iron-phosphorus complex.

C 1s spectra (Fig 4) were fitted with four peaks for all SMER samples at 284.8 eV, 286.5±0.1 eV, 288.1±0.2 eV and 289.2±0.4 eV, corresponding to C1: carbon atoms bond only with carbon and/or hydrogen(C−C/C−H); C2: carbon atoms bond with a single oxygen atom C−O); C3: carbon atoms bond with two non-carbonyl oxygen atoms or a single carbonyl oxygen atom (O=C−O/C=O); C4 carbon atoms bond with a carbonyl and a non-carbonyl group (O−C=O) (Nguiila Inari et al., 2006; Pereira et al., 2014; Popescu et al., 2009). After iron modification, the binding energy of C3 had slightly increased and the intensity of C2 increased when modified in acid (pH 5) and neutral (pH 7) conditions, which suggested that the structure of SMER had changed due to iron modification. The bond area of C4 (O−C=O) became larger after iron modification, whereas the area decreased after phosphate sorption, indicating that the modification process may activate
some functional groups such as carboxylic groups and, once treated with phosphate, these functional groups become part of a more complicated iron-phosphate complex.

Fig. 4. a 2-column fitting image

3.2 Biosorbent modification and sorption test

The virgin SMER displayed an inadequate capacity (0.14 mg P/g) for phosphate sorption, therefore modification of the material was required. Phosphate removal efficiency was found to increase with increasing iron concentration (Fig. 5). The initial concentration of the Fe$^{3+}$ in the system influences the mass transfer of Fe$^{3+}$ ions between the solid and aqueous phase (Aryal et al., 2010). Additionally, when more Fe$^{3+}$ ions were introduced to the system, more iron could be attached to the biosorbent.

Fig. 5. a 1.5-column fitting image

The lowest studied modification pH (5) was found to be the optimal pH for the iron modification process. For each modified SMER treated with the same initial iron concentration, higher phosphate removal efficiency was observed at the lowest modification pH. The statistical analyses confirmed that the initial iron concentration and modification pH and also their interaction were significant in the modification stage (p value < 0.05) (Table S1).

The modification pH is considered to be one of the most critical variables affecting the binding of species. The solution pH has an effect on the speciation of metal ions in solution and on the ionization of the surface functional groups of biomass. At acidic pH
values, Fe\textsuperscript{3+} and positive hydroxide species of iron dominate, whereas insoluble iron hydroxide is formed in the near-neutral region. At higher pH values, negative hydroxide species of iron start to dominate. These results were supported by XPS; more iron was introduced onto the SMER in acidic or neutral solutions.

The optimal pH is strongly dependent on the properties of the biomass. For \textit{Staphylococcus xylosus} biomass, the optimal pH for Fe(III) uptake was 3 when the effect of pH was investigated in the pH range of 1–7 (Aryal et al., 2010). Pagnanelli et al. (2000) investigated the effect of pH (3, 4 and 5) on the sorption of iron(III) by \textit{Arthrobacter sp}. Their results showed that there was a strong increase in the uptake capacity for iron when the pH was 5, which might have been due to the presence of other mechanisms besides binding by functional groups, such as surface precipitation (taking place locally) and co-precipitation. In the present study, as higher modification pH values were studied, surface precipitation played an important role in the binding of Fe(III) on SMER.

3.3 Sorption capacity

The capacity curve for phosphate sorption by iron-modified SMER at different initial phosphate concentrations is presented in Fig. 6. Two batches of iron-modified SMER were compared. The achieved phosphate sorption capacity of batch 1 and batch 2 with a 24 h contact time was approx. 9 mg P/g and 13 mg P/g, respectively. The Redlich-Peterson model was found to provide the best fit to the experimental data (the lowest chi-squared ($\chi^2$) value and the highest coefficient of determination ($R^2$)) (Table S2, Fig. S2). Batch 2 showed better sorption capacity. The difference may arise from experimental uncertainty, for example the kinetics of pH adjustment in the modification. Nevertheless, iron-modified SMER showed great potential for phosphate removal. The capacity
obtained is at the same level as that obtained for other Fe(III)-loaded biomaterials, as shown in Table 1. Further optimization of the procedure (pH and iron concentration) could increase the capacity still further. In addition, pretreatment of biomass has been suggested in order to increase the binding sites for iron. For example, carboxymethyl cellulose treatment of wood fibres and particles enhanced phosphate removal from 0.65–2.78 mg P/g to 2.32–5.67 mg P/g (Eberhardt and Min, 2008).

Fig. 6. a single-column fitting image

3.4 Effect of pH of aqueous media

The pH of the aqueous solution is a critical variable that influences the uptake of anions and cations at solid-liquid interfaces. The sorption of phosphate is highly dependent on the solution pH. Thermodynamic calculations indicate that phosphate exists in aqueous solution in the forms of H₃PO₄, H₂PO₄⁻, HPO₄²⁻ and PO₄³⁻ in different ratios, depending on the solution pH (Krishnan & Haridas, 2008):

\[
\begin{align*}
H_3PO_4 & \leftrightarrow H_2PO_4^- + H^+ & pK_1 = 2.23 \\
H_2PO_4^- & \leftrightarrow HPO_4^{2-} + H^+ & pK_2 = 7.21 \\
HPO_4^{2-} & \leftrightarrow PO_4^{3-} + H^+ & pK_3 = 12.32
\end{align*}
\]

In the pH range of 3–7, H₂PO₄⁻ is the major species, whereas when the pH value is between 7.2 and 12.3, HPO₄²⁻ is the predominant species.

The effect of pH on phosphate uptake is shown in Fig. 7. A strongly acidic environment was avoided in the experiment since a strong acid solution might have led to the destabilization of the biosorbent and iron leaching would have become more likely. Hence, the effect of the solution pH value on phosphate sorption was studied in the range
of 3–9. As shown in Fig. 7, phosphate removal efficiency decreased as the pH value increased. The maximum phosphate removal efficiency reached 82% at pH 3 (initial concentration 15 mg P/L). In addition, more than 50% of phosphate was removed when the solution pH was in the range of 3–5. In contrast, inadequate phosphate uptake was observed in alkaline conditions. Considering that it is not always practical to apply low pH in real activities, a pH value of 4 was selected for subsequent experiments.

Fig. 7. a single-column fitting image

Several studies have revealed a similar effect of pH on phosphate sorption behaviour, i.e. that phosphate sorption was maximized at a pH value in the range of 3–6 (Cui et al., 2016; Liao et al., 2006). In addition, our results were consistent with a previous study conducted by Krishnan and Haridas (2008). They prepared natural and iron-modified coir pith for phosphate sorption, and it was reported that the pH_{PZC} (by potentiometric titration) of iron-modified coir pith was observed at pH 5.8. This means that the iron-modified biomass had a net positive surface charge when the pH was lower than 5.8. Thus, the attachment of negatively charged phosphate species (H$_2$PO$_4^-$) was favoured when the pH was lower than 5.8. However, the value of pH$_{PZC}$ can vary for different iron materials. For two-line ferrihydrite, which was found to be the iron form on the SMER according to XRD, the pH$_{IEP}$ (isoelectric point, by zeta potential measurements) is around pH 7 (Chernyshova et al., 2011).

The low phosphate sorption capacity at high pH values was due to the more negative surface above the pH$_{PZC}$ and thus increased electrostatic repulsion between the surface and negatively charged phosphate ions. In addition, the concentration of OH$^-$ became higher and hydroxides competed with phosphate species for binding sites at high pH.
values. The suggested mechanism for phosphate sorption on similar iron materials involves the ligand exchange for hydroxyl ions and the formation of an inner-sphere complex (Krishnan and Haridas, 2008).

3.5 Effect of contact time

Fig. 8 presents phosphate sorption on iron-modified SMER as a function of time (initial concentration of 15 mg P/L and pH 4). The phosphate removal efficiency slowly increased within the first hour, reaching 26% after 6 h. Subsequently, the removal efficiency continued to increase up to 70% after 48 h contact time. Three kinetic models (pseudo-first-order, pseudo-second-order and Elovich) were used to analyse the kinetic data (supplementary material). Considering both the correlation co-efficient ($R^2$) and the chi-squared ($\chi^2$) value, the Elovich model was found to provide the best fit to the kinetic data (Table S3, Fig. S3). The Elovich model is known to describe the kinetics of chemisorption well (Tran et al. 2017).

Reaching equilibrium is a slow process for amorphous iron materials due to the significant role of the pore diffusion mechanism (Swedlund et al., 2014; Sigdel et al., 2016). Sigdel et al. (2016) reported that alginate beads impregnated with hydrous iron oxide reached equilibrium for arsenic adsorption after around seven days. The hydrous iron oxide was confirmed to be two-line ferrihydrite and the As(V) adsorption was mainly governed by intra-particle diffusion.

Fig. 8. a single-column fitting image

4. Conclusions
The iron loading of Sphagnum moss extract residue (SMER), a waste material, resulted in an effective sorbent for phosphate removal. A modification pH of 5 and the highest initial Fe(III) concentration (0.072 mol/L Fe$^{3+}$) were observed to be the optimal modification conditions. According to XPS, more iron could be attached to the surface of SMER in an acidic solution. FTIR analysis proved that carboxylic acid functional groups (–COOH) were able to capture metal ions during the modification process, whereas XRD analysis confirmed that two-line ferrihydrite was precipitated on the SMER. Phosphate sorption was greatly affected by the pH (maximum removal at pH 3) and contact time. These results corroborate the fact that iron-modified SMER can be used effectively as a low-cost and eco-friendly biosorbent for phosphate removal from wastewaters such as agriculture run-offs. Used sorbent could be safely disposed of in soil as a phosphate fertiliser provided that no toxic components from the treated wastewater are attached. Moreover, the biosorbent material itself would improve the physical properties of the soil.

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**References**


FIGURE CAPTIONS

**Fig. 1.** X-ray diffraction profiles of virgin, iron-modified (Fe-SMER) and phosphate-treated iron-modified SMER (P-Fe-SMER) samples.

**Fig. 2.** FTIR spectra of virgin and iron-modified SMER (pH 5 with 0.072 mol/L Fe$^{3+}$ concentration).

**Fig. 3.** O 1s spectra of SMER: (a) virgin SMER; (b) iron-modified SMER at pH 5; (c) iron-modified SMER at pH 7; (d) iron-modified SMER at pH 9; (e) phosphate-treated iron-modified SMER at pH 5.

**Fig. 4.** C 1s spectra of (a) virgin SMER; (b) iron-modified SMER at pH 5; (c) iron-modified SMER at pH 7; (d) iron-modified SMER at pH 9; (e) phosphate-treated iron-modified SMER at pH 5.

**Fig. 5.** Effect of modification conditions on phosphate sorption (Biosorbent dose 2 g/L; Contact time 24 h; Sorption solution pH 6; Initial phosphate concentration 15 mg P/L; Temperature 20±3°C; Error bars represent the deviation of four repeats (two products made in same conditions and two shaking tests for each product).

**Fig. 6.** Phosphate-P sorption capacity by iron-modified SMER at pH 5 with 0.072 mol/L Fe$^{3+}$ concentration (Biosorbent dose 2 g/L; Contact time 24 h; Sorption solution pH 4; Temperature 20±3°C; Error bars represent the deviation of three repeats).

**Fig. 7.** Effect of pH on phosphate removal by iron-modified SMER at pH 5 with 0.072 mol/L Fe$^{3+}$ concentration (Biosorbent dose 2 g/L; Contact time 24 h; Initial phosphate concentration 15 mg P/L; Temperature 20±3°C; Error bars represent the deviation of two repeats).

**Fig. 8.** Effect of contact time on phosphate removal by iron-modified SMER at pH 5 with 0.072 mol/L Fe$^{3+}$ concentration (Biosorbent dose 2 g/L; Sorption solution pH 4; Initial phosphate concentration 15 mg P/L; Temperature 20±3°C; Error bars represent the deviation of two repeats).