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Environmental significance

Phosphate removal by *ex situ* generated Fe (hydr) oxides from scrap iron electrocoagulation: the critical role of coprecipitation[†]

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The immobilization of phosphate by Fe (hydr)oxides is a promising method with low cost and high efficiency. However, the complicated synthesis process and particle aggregation limited their application. In this study, we utilized electrocoagulation with a scrap iron anode to generate Fe (hydr)oxides in the prepared NaCl electrolyte, which are ready to use for phosphate removal on-site. The Fe (hydr)oxides generated *via* scrap iron electrocoagulation without aeration removed 86.7% of phosphate and showed an energy cost of 35.9 W h g⁻¹ P. The structure and morphology of Fe precipitates containing phosphate were characterized by X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), and transmission electron microscopy (TEM). The results showed that vivianite, strengite, and amorphous Fe–P precipitates were formed immediately after the mixing of Fe (hydr)oxides with phosphate. Compared to coprecipitation, adsorption by Fe (hydr)oxides (*e.g.*, lepidocrocite) plays a minor role in phosphate removal. The findings gained from this study shed light on the feasibility and mechanism of scrap iron materials in the immobilization of phosphate through on-site electrocoagulation.

The looming P crisis makes the removal of P from wastewater the main challenges of the twenty-first century. We report a new strategy of *ex situ* production of Fe (hydr)oxides by scrap iron electrocoagulation in a prepared NaCl electrolyte that is facile and low cost for the sequestration of phosphate, expanding the scope of application of Fe-based materials for phosphate sequestration in wastewater.

1. Introduction

Inefficient use and loss of phosphorus pose increasing risk to food and water security, freshwater biodiversity, and human health.^{1,2} For example, P fertilizer manufacturing emits around 200–280 million tons of phosphogypsum waste per year, which leads to serious water and soil contamination during the storage.^{3,4} Increasing P load possibly leads to the eutrophication of freshwater bodies as well as harmful algal blooms. These phenomena are especially serious in closed waterbodies, such as lakes and ponds, as revealed by exhaustion of dissolved oxygen, death of fish, and bad smell.⁵ The looming P crisis makes the removal of P from wastewater the main challenges of the twenty-first century.⁶⁻⁸

In both China and the United states, the discharge of phosphate is limited to 0.5-1.0 mg L^{-1.5} To meet the strict discharge requirements on P content, technologies including chemical precipitation,9,10 adsorption,11-13 and enhanced biological P removal¹⁴ have been developed. Among the methods, adsorption by a suite of Fe (hydr)oxides (e.g., ferrihydrite, magnetite, and goethite) has shown promise for phosphate removal owing to the advantages of environmental friendliness, low cost, and easy regeneration.^{11,15} The adsorption of PO₄³⁻ on Fe (hydr)oxides was found to be endothermic and spontaneous, possibly via electrostatic attraction and surface precipitation.11,15-17 The adsorption capacities of Fe (hydr)oxides largely depend on the synthesis methods, (hydr)oxide structure, solution pH, and competitive anions.¹² For example, Fe hydroxides (e.g., ferrihydrite and green rust) showed higher adsorption capacity in comparison to oxides (e.g., magnetite and goethite).^{11,15} Moreover, freshly prepared Fe (hydr)oxides were found to have a higher adsorption capacity than aged ones.¹⁸ Fe (hydr)oxides are usually synthesized by co-precipitation or hydrothermal treatment.15,19 However, these methods are either time consuming or difficult to control, limiting the application of Fe (hydr)oxides as a P sequestrator.

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Paper

Electrocoagulation is an efficient method for the production of a range of high-quality Fe (hydr)oxides that are potential sequestrators of phosphate. In the electrocoagulation process, Fe ions are delivered in situ by the electrochemical dissolution of Fe(0) anodes, which react with the hydroxyl ions generated on the cathode to form Fe (hydr)oxides.^{20,21} This process could produce various Fe (hydr)oxide nanoparticles including green rust, magnetite, lepidocrocite, and goethite, depending on the current density, dis-For oxygen, and electrolyte.22,23 example. solved an electrocoagulation process using steel electrodes produced pure magnetite particles of 100 nm diameter, which showed significant magnetic susceptibility.24 Similarly, uniform magnetite nanorods were prepared by a pulsed current electrocoagulation method.25 Ying et al.26 found that magnetite particles could form in NaCl solution regardless of the presence of Mg²⁺ and Ca²⁺ ions by electrocoagulation with carbon steel electrodes. Although some studies reported that the electrocoagulation process was efficient for in situ treatment of excess P in wastewater,27-29 complexed wastewater components would lead to electrode passivation and energy increase.30 Besides, wastewater of low electricity conductivity would lead to high energy consumption in the electrocoagulation process due to the high ohmic drop. Thus, dosing Fe (hydr)oxides generated by electrocoagulation in well-controlled electrolytes such as prepared NaCl solution would be more attractive for the removal of phosphate. However, the knowledge on this aspect is still limited.

In this study, we utilized scrap iron as a sacrificial anode to generate Fe (hydr)oxides to reduce cost and promote waste recycle. The performance of P removal by *ex situ* generated Fe (hydr)oxides was evaluated by using the removal rate, P/Fe ratios, and energy consumption. Specifically, we first compared the performance of different types of scrap iron materials. Then, we investigated the impact of aerations and current intensity on the performance. Ultimately, we characterized the Fe precipitations containing Fe–P minerals by X-ray diffraction (XRD), X-ray photoelectron spectroscopy (XPS), and transmission electron microscopy (TEM). The results obtained from this study shed light on the feasibility of scrap iron materials in the treatment of P wastewater and recycling P to achieve the goal of turning waste into treasure.

2. Materials and methods

2.1. Experimental operation

The description and physical properties of waste materials used as electrodes are presented in Table 1. Electrocoagulation

experiments were carried out in a glass cell of 600 mL at room temperature (Fig. S1 in the ESI⁺). Electrodes made of scrap iron were installed in parallel with a distance of 2.0 cm. Prior to each experiment, the iron electrodes were polished mechanically, and cleaned with deionized (DI) water. 500 mL electrolyte was prepared with 0.2 M NaCl. NaCl was chosen as it is cheap and widely available, and retards the passivation of electrodes.31 After 25 min electrolysis, suspensions containing Fe (hydr)oxides were generated. Then, a stock solution of KH₂PO₄ was added into the suspensions to initiate the experiments of phosphate removal. The solution was thoroughly mixed with a magnetic stirrer during the electrocoagulation and phosphate removal tests. A schematic diagram of the experimental procedure is shown in Scheme 1. The aeration was supplied by an air pump (SB748, Zhongshan Songbao Electric Group) at a flow rate recorded by a gas flow meter (LZB-3WB, Shuanghuan). Constant currents were supplied by a DC power supply (GPS-3303C, Gw-INSTEK) during the electrocoagulation. The energy consumption during the processes was monitored by using a wattmeter (DPT1502, JUWEI). The initial pH values of the suspensions were adjusted with diluted hydrochloric acid (HCl) and sodium hydroxide (NaOH) solution during the phosphate removal tests.

2.2. Calculations

The removal rate (%) was calculated using eqn (1):

Removal rate =
$$\frac{C_0 - C_t}{C_0} \times 100\%$$
 (1)

where C_0 and C_t are the initial and final concentrations of phosphorus (mg L⁻¹).

The phosphate removal capacity of Fe (hydr)oxides (P/Fe, mg g^{-1}) was calculated using eqn (2):

$$P/Fe = \frac{C_0 - C_t}{C_{Fe}} \times 1000$$
⁽²⁾

where C_{Fe} is the final concentration of total Fe generated by electrocoagulation (mg L⁻¹).

The mean energy consumption for phosphate removal (Energy/P, W h g^{-1}) was calculated using eqn (3):

Energy/P =
$$\frac{E}{V(C_0 - C_t)} \times 100$$
 (3)

where E is the power consumption recorded by the wattmeter (W h), and V is the volume of the phosphate solution (L).

Table 1	Description and	physical	properties of	the waste	materials us	sed as	electrodes	in th	is study
---------	-----------------	----------	---------------	-----------	--------------	--------	------------	-------	----------

erap cast iron	Scrap mild steel		
ast iron	Mild steel		
e, C, Si	Fe, Mn, C		
35	7.85		
$00 imes 10^6$	$6.21 imes10^{6}$		
linder	Cylinder		
863	7.006		
2.22	60.70		
	rap cast iron st iron , C, Si 35 00×10^{6} linder 363 .22		



Scheme 1 Schematic diagram of the experimental procedure in this study. Electrodes made of scrap iron were polished and cleaned before usage. Electrolyte was prepared with 0.2 M NaCl solution. After 25 min electrolysis, suspensions containing Fe (hydr)oxides were generated. Then, a stock solution of KH_2PO_4 was added into the suspensions to initiate the experiments of phosphate removal.

2.3. Measurement and analysis

Two samples of the solution were taken at predetermined times during the phosphate removal experiments with one of these samples filtered immediately through a 0.22 μ m nylon membrane. The concentrations of phosphate in the filtered samples were determined by the molybdenum blue spectrophotometric method at 700 nm using an AOE UV-1800 spectrophotometer (Ao Yi Instrument, Shanghai). The solution pH values and the dissolved oxygen (DO) concentrations were measured by using a multifunctional water quality analyzer (Multi-350i, WTW Group, Germany). Dissolved Fe(II) was measured at a wavelength of 562 nm by a ferrozine method.³² Another set of samples acquired from the suspensions were digested in 3 M HCl for the determination of total Fe through the reduction of hydroxylamine hydrochloride.

The solid samples were collected during the phosphate removal tests for X-ray powder diffraction (XRD), X-ray photoelectron spectroscopy (XPS), transmission electron microscopy (TEM), and scanning TEM (STEM) combined with energydispersive X-ray detection (EDX). The suspensions generated after 25 min electrocoagulation were centrifugated and freezedried under anoxic conditions filled with N_2 gas (99.999%). XPS was collected on an AXIS SUPRA+ Spectrometer (SHI-MADZU, Japan) equipped with a monochromatic Al Ka source at 58.7 eV. The binding energy was calibrated using the C 1s XPS peak at 284.8 eV. XRD was carried out on an X-ray diffractometer (SmartLab SE, Rigaku, Japan) with Cu K radiation at a scanning speed of 1° per min and a step size of 0.02°. To characterize the morphology of precipitates, a field-emission gun TEM (Tecnai G² F20, FEI, USA) was operated at 200 kV. The elemental distribution was obtained on the same machine equipped with an EDX detector.

3. Results and discussion

3.1. Impact of aeration

The Fe (hydr)oxides generated during iron electrocoagulation depend on the DO content that is controlled by aeration. We

compared the performance of Fe (hydr)oxides generated under different flow rates of aeration for phosphate removal. As shown in Fig. 1a, phosphate was largely removed in the first 5 min, with a small fraction being removed in the following 25 min. The phosphate removal rates approached 64.16%, 53.26%, 38.94%, and 48.76% at 30 min when 0, 0.25, 0.5, and 1.0 L min⁻¹ air was supplied to the electrocoagulation processes, respectively. In line with the results of removal rates, the phosphate removal capacity of Fe (hydr)oxides decreased with the air flow rate from 0 to 0.5 L min⁻¹, but increased slightly from 0.5 to 1.0 L min⁻¹ (Fig. 1b). The energy consumption for phosphate removal increased from 48.4 W h g^{-1} to 99.5 W h g^{-1} as the air flow rate increased from 0 to 0.5 L min⁻¹, but decreased slightly to 75.1 W h g⁻¹ at 1.0 L min⁻¹ (Fig. 1c).

After 25 min of electrocoagulation, the majority of Fe ions dissolved from the anode were precipitated (Fig. 2a). Especially for 0 L min⁻¹, there was only 0.1 mg L⁻¹ of dissolved Fe detected (Fig. 2a). The low content of dissolved Fe at 0 L min⁻¹ might be due to the rise in solution pH to 10.4 (Fig. 2b), under which conditions both Fe²⁺ and Fe³⁺ were hydrolyzed and precipitated to Fe (hydr)oxides. However, as the solution pH decreased to about 7.3 during the phosphate removal experiments (Fig. 2b), Fe²⁺ would be dissolved and contributed to the increase of the phosphate removal rate after 5 min. Thermodynamic calculation by using Visual MINTEQ v3.1 showed that phosphate competed with hydroxyl ions for Fe²⁺ to form vivianite in the pH range of 4.5-8.0 (Fig. S2a⁺). However, the formation of vivianite would be unfavorable at pH > 9.³³ Therefore, coprecipitation might be a contributor to the phosphate removal under anoxic conditions.

In contrast, when $0.25-1.0 \text{ Lmin}^{-1}$ air was supplied to the electrolytes, the solution pH value was kept at about 6.5 and dissolved Fe could be observed (Fig. 2a and b). The concentrations of DO increase with increasing the air flow rate (Fig. 2c). Under oxic conditions, Fe(II) hydroxides were oxidized quickly, leading to Fe(III) oxides or hydroxides being the main components of precipitates. It has been found that lepidocrocite is the main product of precipitation in the





Fig. 1 (a) Variations of the phosphate removal rate, (b) removal capacity of phosphate, and (c) electricity energy input for the removal of 1 g phosphate by the Fe (hydr)oxides generated from scrap iron electrocoagulation under different air flow rates. Experiment conditions: [electrolyte] = 0.2 M NaCl, 25 min electrocoagulation using the cast iron anode at 0.2 A, and [phosphate]₀ = 10 mg L⁻¹.

Fig. 2 (a) Dissolved Fe content after 25 min electrocoagulation, change of (b) pH, and (c) DO during the phosphate removal processes by the Fe (hydr)oxides generated from scrap iron electrocoagulation under different air flow rates. Experiment conditions: [electrolyte] = 0.2 M NaCl, electrolysis at 0.2 A for 25 min, and [phosphate]₀ = 10 mg L⁻¹.

aerobic electrocoagulation system.^{23,32} The initial quick removal of phosphate might be induced by the dissolved Fe (mainly as Fe²⁺) that resulted in the formation of vivianite and strengite (Fig. S2[†]). Consistent with the concentrations of dissolved Fe, the phosphate removal rate at 5 min was in an order of 0.25 L min⁻¹ >1.0 L min⁻¹ >0.5 L min⁻¹. In the subsequent processes, the phosphate removal rate was slow

under all conditions. The adsorption of phosphate by the Fe(m) hydroxides was facilitated by the presence of hydroxyl groups (–OH and –OH₂).³⁴ However, the fraction of these hydroxyl groups decreased with the aggregation process of nanoprecipitates, and the formation of large flocs reduced the surface area available for phosphate adsorption.³⁵ Therefore,

the removal of phosphate slowed down with increasing reaction time, which was in line with previous observations.³⁶

3.2. Impact of scrap iron materials

To study the effects of different iron materials on the Fe (hydr) oxide production and phosphate removal, scrap cast iron and scrap mild steel were selected for the experiments. As shown in Fig. 3a, the phosphate removal rates were 64.16% and 56.99% in 30 min adsorption by the Fe (hydr)oxides generated from the electrocoagulation process using the cast iron electrodes and the mild steel electrodes, respectively. The phosphate removal capacity of the Fe (hydr)oxides was comparable between the two materials, both about 33.0 mg g⁻¹ (Fig. 3b). However, the energy



Fig. 3 (a) The phosphate removal rate during the experiments by the Fe (hydr)oxides generated from the electrocoagulation processes with different scrap iron materials. (b) P/Fe ratios and (c) energy consumption of the experiments. Experimental conditions: 0.2 M NaCl, 0.2 A for 25 min electrocoagulation, no aeration, and [phosphate]₀ = 10 mg L⁻¹.

consumption using cast iron (48.4 W h g^{-1}) was lower than that using mild steel (65.8 W h g^{-1}) (Fig. 3c).

The production of dissolved Fe ions from the anode is crucial for the phosphate removal by the electrocoagulation process since the Fe ions are the source of Fe (hydr)oxides. The Fe concentration generated by the cast iron electrodes was 181 \pm 20 mg L^{-1} at 30 min, higher than that by the mild steel electrodes (163 \pm 4 mg L⁻¹). The different Fe production might result from the different C contents in the materials. The C content in the cast iron is 5.0-7.9%, which is higher than that in the mild steel (0.6-1.9%). The carbon is mainly present as graphite in the cast iron. The release of Fe on the surface leaves behind a skeleton of graphite flakes, which would contribute to the electrode corrosion.37 Previous work also presented a higher production of Fe ions in the first 25 min using the cast iron anode than that in the case of the mild steel anode.²⁸ Altogether, the high removal rate and low energy consumption made the cast iron outperform mild steel when serving as the anode of the electrocoagulation process.

3.3. Impact of current intensity

The phosphate removal and energy cost performance under different current intensities differed remarkedly. Since high concentrations of Fe (hydr)oxides were generated under high current intensity, the reaction time with phosphate was prolonged to 300 min. The phosphate removal rate substantially increased from 28.0% at 100 mA to 86.7–99.3% when the applied current was \geq 200 mA (Fig. 4a). Among the experiments, the removal capacity of the Fe (hydr)oxides generated at 200 mA was highest being 44.5 mg g⁻¹ (Fig. 4b). Although a high removal rate was reached at 400 and 600 mA, surplus Fe (hydr) oxides generated did not contribute to the phosphate removal. Moreover, the energy consumption increased with the current intensity. For example, the energy consumption at 200 mA (35.9 W h g⁻¹) was less than a third of that at 400 mA (112.2 W h g⁻¹) (Fig. 4c).

Compared to the quick equilibrium of phosphate removal under aeration, the phosphate removal approached the steady stage until 180 min of reactions under no aeration. The DO concentration gradually increased from 0.3 mg L⁻¹ to 6.1 mg L^{-1} , and the solution pH decreased from 10.4 to 5.5 at 200 mA (Fig. S3[†]). During the phosphate removal process, the Fe(II) (hydr)oxides were gradually oxidized by the DO that diffused naturally, which resulted in the release of protons. Then, the weakly acidic conditions were favorable for PO₄-P precipitation as vivianite and strengite as we have calculated in Fig. S2.† However, the solution pH increased to 11.0 after 25 min of electrocoagulation at 600 mA, which only decreased to be pH 9.1 after 300 min reaction. The abundant hydroxyl ions generated during electrocoagulation at high current would not be neutralized during the oxidation. Thus, considering the phosphate removal and energy cost, 200 mA is the optimal current for Fe (hydr)oxide generation by electrocoagulation.

The phosphate removal capacity achieved by the Fe (hydr) oxides at 200 mA was comparable to those reported in the literature. For example, Ajmal *et al.* obtained the maximum



Fig. 4 (a) The phosphate removal rate during the adsorption experiments by the Fe (hydr)oxides generated from the electrocoagulation processes at different current intensities. (b) P/Fe ratios and (c) energy consumption of the experiments. Experimental conditions: 0.2 M NaCl, 25 min electrocoagulation using the cast iron anode, no aeration, and [phosphate]₀ = 10 mg L⁻¹.

adsorption capacity of 57.8 mg g⁻¹ for phosphate using wellprepared magnetite at pH 7.0.¹¹ The removal of phosphate by the *in situ* electrocoagulation is also a promising process, although it depends on wastewater compositions. For the treatment of simulated wastewater containing P of 1–3 mg L⁻¹, the removal capacity of electrocoagulation approached 50 mg P g⁻¹ Fe.³⁸ However, the removal capacities of electrocoagulation were found to be 21.7–93.5 mg P g⁻¹ Fe for the treatment of wastewater containing P of 1.3 mg L^{-1} .³⁶

3.4. Characterization of Fe precipitates

To reveal the phosphate removal mechanism by the Fe (hydr) oxides, XRD, XPS, and TEM were performed to characterize the Fe precipitates. As shown in Fig. 5, vivianite and strengite, formed by the reactions of PO_4^{3-} with ferrous and ferric ions (eqn (4) and (5)), were produced immediately after the reaction. However, the intensity of vivianite gradually disappeared at 120 min, which might result from the oxidation of vivianite.

$$3Fe^{2+} + 2PO_4^{3-} \rightarrow Fe_3(PO_4)_2 \tag{4}$$

$$Fe^{3+} + PO_4^{3-} \to FePO_4 \tag{5}$$

In contrast, the intensity of magnetite and lepidocrocite becomes strong with time, which provides further evidence of the oxidation of Fe hydroxides. The transformation of vivianite to strengite solids might improve the phosphate removal efficiencies, since strengite has a low solubility ($\log K_{sp} = -26$) and is easy to settle.^{39,40} These observations confirmed that coprecipitation contributed to the initial removal of phosphate, and the oxidation of Fe(II) minerals promoted the removal efficiency in the later stage.

The coprecipitation of Fe–P and the transformation of $Fe_3(PO_4)_2$ to $FePO_4$ were also evidenced by XPS analyses (Fig. 6). The P $2p_{3/2}$ at 133.4 eV and $2p_{1/2}$ at 134.2 eV were ascribed to the peaks of PO₄.^{41,42} As shown in the P 2p spectra, PO₄ appeared on the surface once the reaction started, which increased to a high level at 30 min and kept stable during the subsequent processes (Fig. 6a–c). The Fe $2p_{3/2}$ data were successfully modeled using a combination of ferrous and ferric multiplet patterns (Fig. 6d–f and Table S1†). The multiplet peaks at 710.0 and 711.0 eV were ascribed to Fe(II)-PO₄, and multiplet peaks at 711.6, 712.7, 713.8, and 715.1 eV belong to Fe(II)-OH.⁴² Vivianite (Fe(II)-PO₄)



Fig. 5 XRD pattern of solid samples collected at 0, 30, and 120 min after phosphate dosing into the Fe (hydr)oxide suspensions generated under 200 mA without aeration. In the figure, S is the abbreviation of strengite, V is vivianite, L is lepidocrocite, and M is magnetite.



Fig. 6 XPS spectra of solid samples collected at (a and d) 1 min, (b and e) 30 min, and (c and f) 120 min after phosphate addition to the Fe (hydr) oxide suspensions generated under 200 mA without aeration.

accounted for 46% of the surface Fe precipitates in the beginning. However, at 30 min, the content of Fe(II)-PO₄ decreased to 40%, suggesting that oxidation occurred during the process, in line with the findings of XRD results. The oxidation might be partial, since the content of Fe(II)-PO₄ still accounted for 40% at 120 min, due to the stability of the electronic configuration of a Fe(II)_B-Fe(III)_B pair relative to a Fe(II)_B-Fe(III)_B pair.⁴² Thus, the vivianite on the surface was partly oxidized during the reactions.

The TEM images of Fe precipitates collected at 1 min, 30 min, and 120 min of the experiments are shown in Fig. 7a-c and S4.† At 1 min, the Fe precipitates showed spherical-shaped substances at locations 1 and 3, planar-shaped substances with smooth surface textures at location 2, and individual lath-like particles with lengths of 20-50 nm. These lath-like particles are lepidocrocite that forms under conditions of rapid oxidation of Fe²⁺.43 The element composition of the Fe precipitates was further examined by STEM-EDX, with 3 locations reported on the right of Fig. 7. The atomic contents of phosphate are less than 0.8% at locations 1 and 2, while it is up to 6.6% at location 3. This finding suggests that the planar-shaped substances and spherical-shaped substances with crystal edges are Fe precipitates with low phosphate adsorption capacity. The sphericalshaped substances devoid of crystal edges and lattice fringes at location 3 might be amorphous Fe-phosphate precipitates, since the Fe polymerization is largely limited to the stage of monomers or oligomers by the presence of phosphate.43,44 At 30 min, the Fe precipitates exhibited more lath-like particles with lengths of 100-200 nm, which could be ascribed to the oxidation of Fe(II) precipitates to lepidocrocite.³⁴ Furthermore, the STEM-EDX results showed a phosphate content of about 3%

at all locations examined, which suggested adsorption of phosphate on these lepidocrocite nanoparticles and formation of Fe-phosphate polymers. At 120 min, beyond individual lathlike particles, there are spherical-shaped substances containing high contents of P (6.3% at location 1). Since the oxidation of $Fe(\pi)$ precipitates decreased the solution pH to about 5.5, the dissolution and reformation of Fe precipitates might result in the reoccurrence of spherical-shaped substances. The average contents of P increased from 2.66% to 3.57% as the reaction time increased from 1 min to 120 min, which is consistent with the decrease of phosphate concentrations in the solution. The phosphate content in the precipitate is much higher than the reported sorption capacities of lepidocrocite at near neutral pH (1.5%).44 Thus, phosphate removal could not be explained by merely adsorption. In contrast, the coprecipitation of phosphate and Fe would not be limited by the surface area. As a result, the high content of P might mainly come from coprecipitation of Fe-P.

3.5. Environmental implications

A conceptual model of phosphate removal by electrocoagulation using scrap iron electrodes is described in Scheme 2. As a lowcost and easy-access material, scrap iron (*e.g.*, cast iron stick) could be used for the electrode after a simple treatment, such as cutting and mechanical clean. Electrode passivation is a major challenge of electrocoagulation. To solve this issue, we proposed to use the prepared NaCl solution as the electrolyte, instead of electrocoagulation performed directly in wastewater. As evidenced in this work, we found that the current efficiencies



Fig. 7 Bright field TEM images of the Fe-P solid samples obtained at (a) 1 min, (b) 30 min, and (c) 120 min. The element contents of the locations 1–3 identified by EDX are presented on the right.



Scheme 2 A conceptual model of on-site dosing of Fe (hydr)oxides for phosphate removal by electrocoagulation with scrap iron electrodes.

were always close to 100% independent of aeration, anode material, and current intensity of 100–600 mA (Fig. S5†). The electrocoagulation in NaCl solution could be installed on-site for Fe (hydr)oxide dosing in P-burden wastewater treatment facilities. The Fe (hydr)oxides generated by this method exhibited high efficiency (86.7–99.3%) for phosphate removal. The normalized specific energy consumption is 35.9–112.2 kW h per kg P, or 18.0–56.1 CNY per kg P assuming that the

price of electricity is 0.5 CNY per kW h. The phosphate coprecipitated or adsorbed with Fe (hydr)oxides could be separated from the wastewater by sedimentation or filtration. The strategy proposed in this work utilizing available and low-cost materials in daily life provides a promising solution for phosphate removal from wastewater.

4. Conclusions

This study evaluated the feasibility of an ex situ electrocoagulation utilizing scrap iron electrode for phosphate removal over a range of conditions representative of wastewater. The Fe precipitates generated at 25 min of electrocoagulation with cast iron electrodes could remove 86.7% of phosphate (10 mg L^{-1}), which showed a phosphate removal capacity of 44.5 mg g⁻¹ Fe and energy cost of 35.9 W h g⁻¹ P. In a 0.2 M NaCl electrolyte, the current efficiency of electrocoagulation could approach 100%, which avoids electrode passivation. Compared to the electrocoagulation under air aeration, the Fe precipitates generated by electrocoagulation without aeration showed better performance, which also saved the energy consumption. The high removal rate and low energy consumption for phosphate removal made the cast iron outperform mild steel when serving as the anode of the electrocoagulation process. The phosphate retained in the Fe precipitates was mainly incorporated into the structural as

vivianite, strengite, and amorphous Fe–P precipitates. We acknowledge that the efficiency of this method for the treatment of real wastewater with varied concentration of phosphate and other contaminants is still to be explored. Ongoing research is required to apply this new strategy for phosphate removal from domestic wastewater on different scales.

Author contributions

Shiwei Xie: conceptualization, writing-original draft, writingreviewing and editing, funding acquisition, and supervision. Zhengkang Bai and Wei Shao: writing-original draft, investigation, and visualization. Chen Wang, Jianglong Qin and Ze Liu: methodology and investigation. Peng Liao: writing-reviewing and editing, funding acquisition, and supervision.

Conflicts of interest

The authors declare no conflict of interest.

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