Silicate minerals control the potential uses of phosphorus-laden mineral-engineered biochar as phosphorus fertilizers

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Abstract
Silicate minerals constitute the main components in silicon (Si)-rich biomass, affecting the phosphorus (P) adsorption and release competencies of mineral-engineered biochar; however, the mechanisms underlying their differences remain largely unresolved. To examine these interactions, we investigated the mineralogical compositions and quantified the P-adsorption capacities of Al-, Fe-, Mn-, Zn-, and Mg-engineered biochars from Si-rich rice husk material. The potential uses of P-laden mineral-engineered biochar for P fertilizers were assessed using citric acid extraction. The results from X-ray diffraction, scanning electron microscopy, and Fourier transform infrared spectrometry revealed that mixed metal (oxyhydr)oxides and metal-silicate compounds precipitated in the biochar structure and acted as the main P adsorbents. Micro-crystalline silicates derived from the biomass-induced metal-silicate precipitates in all engineered biochars, which effectively retained the aqueous P with varying excellent capacities (25.6–46.5 mg/g) but relatively slow kinetics (48 h). The suitability of the Zn-, Mg-, Mn-, and Fe-biochars as P-recycled fertilizers was confirmed by the high amounts of citric acid extractable P (19–69% of the total P). Varying amounts of Zn, Mg, and Mn (34–47% of the total host metals) were also released from the engineered biochar through ligand-promoted dissolution. Our data shed light on the novel potential utilization of Mn-, Mg- and Zn-biochars from Si-rich biomass for P retrieval and their use for P, Mg, and micronutrient (Mn and Zn) fertilizers. Regarding the P removal capacity, the mineral-engineered biochar needed a longer adsorption period than conventional metal-engineered biochar.

Graphical Abstract

Extended author information available on the last page of the article

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1 Introduction

Biochar utilization constitutes a clean and green technology for the effective recycling of biomass and organic wastes because it can purify pollutants from wastewater and aquatic environments (Ahmad et al. 2014). Biochar loaded with metal (oxyhydr)oxides—mineral-engineered biochar—has been recognized as an adsorbent that can improve the competency of biochar for contaminant removal. The competency for anionic pollutant retention is primarily controlled by the biomass type and added metals (El-Naggar et al. 2019; Mohan et al. 2018; Prakongkep et al. 2015). Many studies have loaded biochar with Al, Fe, Mn, and Mg to form metal (oxyhydr)oxides to act as the main effective sorbents for enhancing their removal capacity (Luo et al. 2021; Peng et al. 2019; Takaya et al. 2016; Vikrant et al. 2018; Wang et al. 2015; Yang et al. 2019a; Zhang and Gao 2013). Mg-biochar has been studied the most for its potential use in P removal because of its high retention capacity (95–835 mg/g) and its subsequent regeneration as P-recycled fertilizer (Takaya et al. 2016; Yao et al. 2013). Recent studies showed that Fe/Al biochars retained P up to 215 mg/g (Peng et al. 2019; Sornhiran et al. 2021). To our knowledge, there has been no investigation on Mn-biochar for use as a P adsorbent, although in an arsenate sorption study it was reported to have low arsenate retention capacity of 0.59 mg/g (Wang et al. 2015). A study using Zn-biochar for P removal reported a maximum retention of only 5.1 mg/g (Namasivayam and Sangeetha 2004).

Very limited studies have examined Al-, Fe-, Mn-, and Zn-engineered biochars and reused them as P-recycled fertilizers, attributed to Fe and Al oxides showing a strong binding capacity towards P and thus lowering P release (Peng et al. 2021). A very recent study revealed the possible use of Fe- and Al-biochars for P-recycled fertilizers as they could release adequate available P concentrations to sand material (Sornhiran et al. 2021). Most importantly, Fe, Mn, and Zn are plant micronutrients; however, the production of the micronutrient-biochars as a P adsorbents and their regeneration as both P- and micronutrient-recycled fertilizers that potentially offer a greener and cleaner technology for environmental sustainability are unresolved.

Rice husk is a promising agricultural residue for biochar production; globally approximately 140 million t, is generated annually (Moraes et al. 2014; Shen 2017). Silicate compounds in such biomass can substantially affect the adsorption and release properties of the mineral-engineered biochar. Amorphous silicates with high solubility in the biomass could polymerize or coprecipitate with metal (oxyhydr)oxides (Christl et al. 2012) and might decrease the efficiency of mineral-engineered biochar for P reclamation. Such evidence might unintentionally support a belief that Si-rich material is unsuitable for preparing engineered biochars for P removal, resulting in it simply not to being used as biochar feedstock. Contrariwise, we hypothesized that the polymerization and coprecipitation mechanisms between metal (oxyhydr)oxides and silicates could offer additional benefits by enhancing P release from adsorbents with strong P-adsorption bonding energy with Al, Fe, Mn, and Zn-engineered biochars, and thereby further promote their potential reuse as P-recycled fertilizers rather than being used solely as P adsorbents. To date, key information remains seriously lacking on the P-sorption capacity and the possible use of engineered biochar after P addition (P-laden biochar) from Si-rich material for P-recycled fertilizers.

Therefore, this research aimed: (i) to investigate the capacity and suitable pH conditions of Al-, Fe-, Mn-, Zn-, and Mg-biochars prepared from rice husk for P removal and (ii) to assess the P and metal extractability of the
mineral-engineered biochars for potential uses as combined P and micronutrient fertilizers using citric acid. This organic acid is the most common extractant for assessing P fertilizer potential use and is the acid released from plant roots to promote P and micronutrient availability through ligand-promoted dissolution (Mizuki and Katoh 2021; Yang et al. 2019b).

2 Materials and methods

2.1 Preparation of pristine and mineral-engineered biochars

Rice husk, as one of the primary agricultural residues in several rice-producing countries around the world, was selected for biochar production. The pristine biochar (PB) was produced as described by Klüpfel et al. (2014); this procedure has proved to successfully obtain biochar with excellent chemical activity. Briefly, the intact raw material was air-dried, tightly packed into a heat-resistant crucible (35 mL) to minimize oxygen diffusion and then placed in a muffle furnace (Carbolite 91E; Carbolite; UK) preheated at 600 °C. The closed vessel was carbonized under an ambient atmosphere for 1 h before removal and cooling in a desiccator before further analyses.

The pretreatment process was selected for mineral-engineered biochar production since many studies reported that this method provided higher P-retention capacity than the post-treatment process (Takaya et al. 2016). The mineral-engineered biochars were produced as described by Zhang et al. (2012) with some modifications. About 10 g samples of the air-dried biomass were individually saturated with 60 mL each of AlCl3, FeCl3, MnCl2, ZnCl2, and MgCl2 (67% w/v). The equilibrium periods varied between 48 and 72 h, depending on the saturation of each metal-treated biomass. The metal-impregnated biomass was oven-dried at 80 °C for at least 24 h and carbonized using the same procedure as for PB production. The PB and mineral-engineered biochar samples were washed several times with deionized water to remove impurities and were dried at 80 °C for 24 h. All samples were finely ground using an agate mortar and passed through a 0.5–2.0 mm sieve. The resultant material was collected and preserved in an airtight container for further investigation. The engineered biochar samples treated with Al, Fe, Mn, Zn, and Mg were designated as EB_{Al}, EB_{Fe}, EB_{Mn}, EB_{Zn}, and EB_{Mg}, respectively.

2.2 Characterization of pristine, metal-engineered, and phosphorus-laden biochars

The mineral and chemical compositions and physical properties of all engineered biochar and P-laden biochar samples were analyzed using X-ray diffraction (XRD), Fourier transform infrared spectrometry (FTIR), scanning electron microscopy (SEM) imaging, acid digestion, and physisorption analysis. The powder XRD patterns of all biochar samples were collected using a PANalytical X-ray diffractometer equipped with Cu Kα radiation (λ = 1.54059 Å) over the range 5–70° 2θ, using a step size of 0.015° 2θ, and a dwell time of 0.02° 2θ/s. The FTIR spectra were recorded in the 4000–400 cm⁻¹ region with a resolution of 2 cm⁻¹ using an FTIR spectrometer (Bruker Tensor 27; Bruker; USA). The size, shape, and chemical composition of minerals (oxides and phosphate compounds) in the mineral-engineered biochar and P-laden biochar were investigated in detail using SEM (JEM-2010 (HR); JEOL; Japan) combined with energy-dispersive X-ray spectrometry (EDS; Oxford Instruments; UK). The total P in the samples was quantified based on digestion of 0.20 g finely ground samples with 12 mL of a mixture of 1:3 hydrochloric acid (HCl): nitric acid (HNO₃; aqua regia) at 130 °C for 3 h (Chen and Ma 2001). The inorganic P concentration in the digests was colorimetrically measured using a spectrophotometer. The specific surface area was analyzed using the Brunauer–Emmett–Teller (BET) method on an adsorption analyzer (3Flex Physisorption; Micromeritics; USA) with the volume and size (mesopore = 2–150 µm; and micropore = < 2 µm) of pores calculated from the respective Density Functional Theory and Barrett-Joyner-Halenda analysis.

2.3 Determination of zeta potential of pristine and mineral-engineered biochars

The zeta potential was measured following the procedure of Szewczuk-Karpisz et al. (2020) to investigate the surface charge behavior of the PB and EB samples. Briefly, the samples were ground to pass through a 0.053 mm sieve. About 40 mg of the solid was mixed with 400 mL of 1 mM NaCl solution. The suspension pH value of each sample was adjusted to 2.0–10.0 using HCl and NaOH, and the samples were equilibrated for 3 days. The zeta potential was determined using a Zetasizer instrument (Zetasizer nano series; Malvern Panalytical; UK). The plot between the suspension pH and zeta potential of the biochar was used to identify the isoelectric point (IEP).
2.4 Phosphorus sorption of mineral-engineered biochar

2.4.1 Sorption isotherm

The P-sorption isotherms of all biochar samples (PB, EBAl, EBFe, EBMn, EBZn, and EBMg) were performed in triplicate. Solid materials were weighed in separate centrifuge tubes and mixed with varying P concentrations of KH2PO4 (20–500 mg/L) using a solid-to-solution ratio of 1:500. The vials were agitated on a mechanical shaker at room temperature (25 ± 2 °C) for 24 h. The aliquots were kept after centrifugation at 2205×g for 15 min for P analysis and metal analyses, as described above. Details of the Langmuir and Freundlich’s models are given in supporting information.

2.4.2 Sorption kinetics

The P-sorption kinetics were investigated to obtain information on how quickly the mineral-engineered biochars retained the dissolved P. Each mineral-engineered biochar was weighed into a centrifuge tube with the P solution at the defined the P sorption maximum (Qm) concentration. The experiment was run in triplicate. The vials were agitated on a mechanical shaker at room temperature (25 ± 2 °C) for periods of up to 72 h. After defined periods of 1, 2, 4, 6, 12, 24, 48, and 72 h, the aliquots were retrieved after centrifugation and preserved for P analysis and metal analyses. Details of the kinetics models are given in supporting information.

2.4.3 Sorption envelope

The P-sorption envelope was determined on the mineral-engineered biochar samples to obtain the suitable pH conditions for maximizing P removal. Each mineral-engineered biochar sample was weighed into a centrifuge tube containing the P solution at the Qm concentration and was adjusted to pH 4.0 ± 0.02, 6.0 ± 0.02, and 8.0 ± 0.02 using HCl or NaOH. The vessels were equilibrated using a mechanical shaker at room temperature (25 ± 2 °C) for defined periods of 1, 2, 4, 6, 12, 24, 48, and 72 h. The supernatants were collected for P analysis after centrifugation at 2205×g for 15 min.

2.5 Phosphorus release of phosphorus-laden biochar

All mineral-engineered biochar samples were treated with different P concentrations and solution pH values to obtain approximately 10% of their Qm concentration. The defined concentrations were expected to obtain the P contents in P-laden biochar after P retrieval from real wastewaters containing 0.9–6.4 mg P/L with a 1:500 solid-to-solution ratio (Zheng and Gao 2019). The sample aliquots were equilibrated using a mechanical shaker for 48 h before centrifugation at 2205×g for 15 min and then heated at 80 °C. All engineered biochar samples (EBAl-P, EBFe-P, EBMn-P, EBZn-P, and EBMg-P) were designated as P-laden biochar (EBAl-P, EBFe-P, EBMn-P, EBZn-P, and EBMg-P). The pH values of the P-laden biochar samples were measured using a pH meter at a solid-to-solution ratio of 1:20.

Citic acid extraction was selected to obtain the capacity of P released from the P-laden biochar samples because this solution has been widely accepted for considering the potential of biochar as P fertilizer (Weber et al. 2014). In addition, citric acid is an organic acid commonly released from plant roots to enhance the acquisition of P and micronutrients (Bais et al. 2006). The P-laden biochar was mixed with 2% citric acid (w/v) with a solid-to-solution of 1:100. The tubes were agitated on a shaker at 25 ± 2 °C for 30 min. The supernatants were kept, and the P concentrations in the aliquots were colorimetrically analyzed using spectrophotometry. The experiment was run in triplicate.

3 Results and discussion

3.1 Characterization of pristine, metal-engineered, and phosphorus-laden biochars

3.1.1 BET analysis

The BET properties of the biochars are given in Table 1. Impregnating biomass with Al, Fe, Mn, Zn, or Mg increased the mesopore area, cumulative pore volume, and median pore size of the engineered biochar samples relative to the pristine biochar. The lowest specific surface area value was for EBMg (246 m²/g), and the highest value was for EBZn (848 m²/g). The extraordinarily high specific surface area of the EBZn was attributed to the preferred formation of Zn-silicate with large crystal sizes (> 10 µm) (Figs. 1, 2) and thereby contributing solely to the mesopore area. Notably, the biochar engineering with metals promoted the mesopore area formation, which was 51–898% higher than that in the PB sample. Biochar engineering with metals increased both the cumulative pore volume (0.142–1.421 cm³/g or 213–3058% of the PB value) and median pore width (0.47–0.77 nm, or 9–79% of the PB value) compared to the respective PB samples (0.045 cm³/g and 0.43 nm).

3.1.2 XRD and chemical analysis

While quartz and calcite were the major minerals in the PB sample (Fig. S1), many metal-(oxyhydr)oxides and...
metal-silicate compounds dominated in all the mineral-engineered biochar samples (EBAl, EBFe, EBMn, EBZn, and EBMg) after the rice husk had been treated with the metals (Fig. 1). Boehmite (AlOOH) was the sole Al (oxyhydr)oxide in the EBAl sample, while various amounts of Fe (oxyhydr)oxides, including hematite (α-Fe\textsubscript{III}O\textsubscript{3}), magnetite (α-Fe\textsubscript{III}Fe\textsubscript{II}O\textsubscript{4}), amakinite (Fe\textsubscript{III}(OH)\textsubscript{2}), and crenstedtite (Fe\textsubscript{II}Fe\textsubscript{III}([SiFe\textsubscript{III}]O\textsubscript{4}[OH]\textsubscript{5}) were present in the EBFe sample. Manganite (Mn\textsubscript{III}O\textsubscript{2}OH), kanoite (Mn\textsubscript{II}SiO\textsubscript{3}), and quartz occurred in the EBMn sample. Willemite (Zn\textsubscript{2}SiO\textsubscript{4}) and K\textsubscript{2}Zn(SO\textsubscript{4})\textsubscript{2} were found in the EBZn sample, while periclase (MgO), brucite (Mg(OH)\textsubscript{2}), and sepiolite [Mg\textsubscript{4}Si\textsubscript{6}O\textsubscript{15}(OH)\textsubscript{2}·6H\textsubscript{2}O] were observed in the EBMg sample.

There were no metal phosphate compounds detected in any P-laden biochar samples using the XRD technique (Fig. 1). This was attributable to the P loading being only about 10% of the P-sorption maximum capacity for each engineered biochar, which suggested that the metal-phosphate compounds were formed in low quantities below the detection limit (0.2–5%) of the conventional XRD technique for minerals with a good crystalline structure (Newman et al. 2015).

The total P concentrations in the P-laden biochar were in the range 2.38–3.89 mg/g, with the lowest and highest values for EBZn-P and EBAl-P, respectively (Table S1). Most engineered biochar samples after P loading were acidic with the range in pH values covering very strongly acidic for EBFe-P (pH = 4.37), strongly acidic for EBZn-P (pH = 5.51), and moderately acidic for EBMn-P (pH = 5.65) and EBAl-P (pH = 5.86). Only EBMg-P had a neutral pH value (pH = 7.34).

### 3.1.3 SEM analysis

The engineered and P-laden biochar samples were examined in detail using the SEM–EDS technique. The SEM–EDS data showed diverse types, sizes, and shapes of metal compounds in the engineered biochar (Fig. 2), and the corresponding P-laden biochar samples (Fig. 3) were primarily governed by the added metals and Si compounds in the raw material. In addition to the XRD data, the SEM–EDS analysis provided new information that most of the metal (oxyhydr)oxides in the engineered biochar occurred as different crystals of metal-silicate mixtures, including Al–silicate (anhydrous), Fe–silicate (compact granular), Mn–silicate (cubic), Zn–silicate (straw-bundle-like), and Mg–silicate (anhydrous). The information suggested that the engineered metals were likely to react with dissolved microcrystalline Si minerals in the raw rice husk biomass during the metal saturation process, which could be newly precipitated as metal-silicate compounds upon pyrolysis. The particle sizes of the metal silicate compounds from the SEM analysis were typically smaller than 1 µm for the EBAl, EBFe, EBMn, and EBMg samples, but those for the EBMg samples were larger than 10 µm.

Although metal silicate mixtures were undetectable using XRD, the SEM–EDS analysis demonstrated that they were the primary adsorbents for P (Fig. 3). The particle shapes of the P-metal-silicate compounds noticeably differed from those of the engineered biochars. The particle sizes of the P-metal-silicate mixtures were also smaller than those of the mineral-engineered biochars. The data suggested that the dissolution of metal-oxide and metal-silicate compounds and their subsequent coprecipitation with the added P could be the main mechanism for P retention within the engineered biochar structure, in addition to the electrostatic attraction mechanism during the P adsorption process.

### 3.1.4 FTIR analysis

The FTIR spectrum of the PB sample indicated the presence of Si–O–Si, aliphatic CH\textsubscript{2}, and aromatic C=C and C=O functional groups (Fig. 4). For all metal engineered biochar samples, the intensity of the bands in the carboxylate region (1700–1400 cm\textsuperscript{-1}), particularly the peaks at 1441, 1561, and 1651 cm\textsuperscript{-1}, became more intense relative to the PB sample, suggesting that carboxylic functional groups

| Sample | Specific surface area (m\textsuperscript{2}/g) | Micropore area (m\textsuperscript{2}/g) | Mesopore area (m\textsuperscript{2}/g) | Relative mesopore area (% of total) | Cumulative pore volume (cm\textsuperscript{3}/g) | Average pore diameter (nm) | Median pore width (nm) |
|--------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| PB     | 255             | 170             | 85              | 33              | 0.045           | 5.11            | 0.43            |
| EB\textsubscript{Al} | 341 (34) | 40 (–76) | 301 (254) | 88              | 0.258 (473)    | 4.69 (–8)       | 0.59 (37)        |
| EB\textsubscript{Fe} | 290 (14) | 163 (–4) | 128 (51) | 44              | 0.158 (251)    | 8.60 (68)       | 0.47 (9)         |
| EBM\textsubscript{n} | 299 (17) | 124 (–27) | 175 (106) | 59              | 0.137 (204)    | 4.34 (–15)      | 0.50 (16)        |
| EB\textsubscript{Zn} | 848 (233) | 0 (–100) | 848 (898) | 100             | 1.421 (3,058)  | 6.86 (34)       | 0.77 (79)        |
| EBM\textsubscript{g} | 246 (–4) | 106 (–38) | 140 (140) | 57              | 0.141 (213)    | 5.62 (10)       | 0.50 (16)        |
could be a primary adsorption site for the added Al, Fe, Mn, Zn, and Mg. The broad peak at 482 cm⁻¹ for all the engineered biochar samples was less intense than that for the PB sample, which could be interpreted as the dissolution of polymeric silicates from the engineered biochar. The band of Si–O–Si at 742 cm⁻¹ was more pronounced in some engineered biochar samples (EBAl, EBAl-P, EBMn, and EBMn-P), which may indicated the coprecipitation of the metal with silicates. The broad peak at around 600 cm⁻¹ was assigned to Al–O bonding being visible in the EBAl sample (Thabet and Ismaiel 2016). The small peak at 850 cm⁻¹ in the EBMg and EBMg-P samples corresponded to Mg–O bonding (Akgul et al. 2019). The small peaks that occurred at 1250 cm⁻¹ in all P-laden biochars could be interpreted as the symmetric vibrations of metaphosphate formation (El Batal et al. 2016). Additionally, the region around 800–1200 cm⁻¹ indicated the presence of PO₄³⁻ (Arai and Sparks 2001) and the presence of the peak at about 3450 cm⁻¹ (data not shown) indicated the OH stretching of free water molecules, which coincided with the high water level of carbon-coated montmorillonite nanocomposite (Wei et al. 2019).

3.1.5 Zeta potential analysis

Potentiometric titration data for the pristine and mineral-engineered biochar samples are given in Fig. 5. The zeta potential of all biochar samples became more negative with increasing suspension pH, which indicated the more negatively charged sites were on solid surfaces. The IEP value of the PB sample was 2.80, which was close to the point of zero charge (PZC) of quartz (Essington 2015). This value was lower than that for other EB samples. The IEP values of the EB samples increased in the order EB Mn (2.84) < EB Mg (3.63) < EB Zn (4.54) < EB Fe (4.90) < EB Al (8.45), which were in strong accordance with their mineralogical compositions, showing the presence of metal silicates including kanoite, sepiolite, willemite, and cronstedite in the respective EB Mn, EBMg, EB Zn, and EB Fe samples and boehmite in the EBAl sample (Fig. 1). These data demonstrated that the silicate compounds in the biomass did play an important role in changing the chemical composition and particle charge behavior.

3.2 Phosphorus sorption of mineral-engineered biochar

3.2.1 Adsorption isotherms

The maximum P-adsorption capacities of all the engineered biochar samples were higher than that for the pristine biochar (Table 2). The P sorption data were well described by both the Langmuir and Freundlich models with coefficient of determination (R²) values in the ranges 0.72–0.99 (mean = 0.91) and 0.65–0.97 (mean = 0.89), respectively (Fig. 6). The EBMg sample was the most effective engineered biochar for aqueous-P removal with a maximum sorption capacity of 46.5 mg/g, followed by the EB Mn sample that adsorbed P up to 38.6 mg/g. The EB Al and EB Fe samples retained comparable P quantities of 32.3 and 31.7 mg/g, respectively. The EB Zn sample had the lowest P maximum sorption capacity of 25.6 mg/g. The Langmuir K L coefficient (relating to P-binding energy) varied considerably from
Fig. 2 Backscattered electron micrographs using SEM–EDS of engineered biochars from rice husk pyrolyzed at 600 °C showing metal-silicate mixtures of EB_{Al} (a), EB_{Fe} (b), EB_{Mn} (c), EB_{Zn} (d), and EB_{Mg} (e). Small orange squares represent the area of SEM–EDS analysis. Atomic ratios: Al:Si ≈ 1.0:1.7, Fe:Si ≈ 1.0:1.7, Mn:Si ≈ 1.0:0.2, Zn:Si ≈ 1.0:0.4, and Mg:Si ≈ 1.0:0.2
Fig. 3  Backscattered electron micrographs using SEM–EDS of P-laden biochars showing metal-silicate mixtures containing P in $\text{EB}_{\text{Al-P}}$ (a), $\text{EB}_{\text{Fe-P}}$ (b), $\text{EB}_{\text{Mn-P}}$ (c), $\text{EB}_{\text{Zn-P}}$ (d), and $\text{EB}_{\text{Mg-P}}$ (e). Small orange squares represent the area of SEM–EDS analysis. Atomic ratios: Al:Si:P ≈ 1.00:0.02:0.01, Fe:Si:P ≈ 1.00:3.76:0.12, Mn:Si:P ≈ 1.00:0.15:0.76, Zn:Si:P ≈ 1.00:0.15:0.76, and Mg:Si:P ≈ 1.00:0.37:0.39.
0.01 to 0.53 L/mg. The lowest $K_L$ value in the EB$_{Mg}$ sample and the highest value in the EB$_{Fe}$ sample suggested low and high P-binding energies for the EB$_{Mg}$ and EB$_{Fe}$ samples, respectively. The Freundlich $K_f$ coefficient (associated with the sorption site quantity) ranged from 0.12 for the EB$_{Fe}$ to 12.21 for the EB$_{Al}$ samples. The Freundlich $n$ constant, interpreted as differences in the adsorption sites of adsorbents, substantially differed from 0.15 for the EB$_{Al}$ to 0.80 for the EB$_{Fe}$ samples, with a low value indicating heterogeneity of the reactive adsorption sites and a high value denoting homogeneity of the adsorption sites (Sposito 1980). The $n$ constant value of below unity also demonstrated that the chemical mechanisms were more favorable under the tested conditions for P sorption than the physical processes (Ajmal et al. 2018; Ozcan et al. 2005). This evidence conformed to the data showing that enhancing the specific surface area and the cumulative pore volume of the engineered biochar caused by metal oxide impregnation demoted their maximum P-adsorption capacities (Fig. S2).

### 3.2.2 Adsorption kinetics

The kinetics data demonstrated that P adsorption onto the EB samples was slow and reached equilibrium after 48 h (Fig. S3). With the addition of the P contents in the samples at $Q_m$ concentrations of 32.3, 31.7, 38.6, 25.6, and 46.5 mg/g for the respective EB$_{Al}$, EB$_{Fe}$, EB$_{Mn}$, EB$_{Zn}$, and EB$_{Mg}$ samples (Table 2), the EB$_{Mn}$ and EB$_{Mg}$ samples removed 76 and 74%, respectively, of the total added P after 72 h. The EB$_{Al}$, EB$_{Zn}$, and EB$_{Fe}$ samples removed 62, 55, and 45%, respectively, of the total added P.

Five kinetic models (zero-order, first-order, parabolic diffusion, power function, and Elovich model) were used to describe the P-adsorption kinetics on the engineered biochar. The results demonstrated that the Elovich model provided the best description for the P sorption kinetics...
in most samples with $R^2$ values in the range 0.79–0.99 (mean = 0.92, Table 3), suggesting that the adsorption of P onto the mineral-engineered biochar could be a diffusion-controlled mechanism. These data agreed well with Wang et al. (2015) and Zhang and Gao (2013), showing that the As(V) adsorption kinetics on Mn-engineered biochar and Al-engineered biochar were well fitted by the Elovich model ($R^2 = 0.98$ and 0.96, respectively). This may have been due to the colloidal particles of metal (oxyhydr)oxides in the biochar acting as the primary adsorbents for added P. However, other studies suggested that first- and second-order equations better explained the sorption kinetics data for As (V) onto Fe biochar and P sorption onto Mg-biochar (Yao et al. 2013; Zhang et al. 2013), indicating that multiple mechanisms might control the adsorption of oxyanions onto mineral-engineered biochar. The $R^2$ values for the other models decreased in the following order: power function ($R^2 = 0.84–0.99$, mean = 0.90) and parabolic diffusion ($R^2 = 0.77–0.99$, mean = 0.90) > zero-order ($R^2 = 0.66–0.91$, mean = 0.79) > first-order ($R^2 = 0.67–0.75$, mean = 0.65).

### 3.2.3 Adsorption envelope

From the P sorption of each EB sample at the defined $Q_m$ levels and different pH values of 4.0, 6.0, and 8.0, it was evident that the initial pH values of the $KH_2PO_4$ solution made a substantial contribution to the P-sorption capacity of the engineered biochar. The highest P adsorption for most samples (EBAl, EBFe, EBMn, and EBMg) occurred at solution pH 4.0, except for the EBMn samples with the highest P at pH 6.0 (Fig. 7). The P-retention capacities for the EB samples at pH 8.0 decreased substantially by 17–76% and 24–67% compared to those at pH 4.0 and 6.0, respectively. This could be due to the initial solution pH of 4.0 largely deviating from the PZC values of the pure (oxyhydr)oxides of Al (boehmite: 8.5), Fe (magnetite and synthetic hematite: 4.8–7.0), Mn (manganese oxide: 4.0–4.5), Zn (zinc oxide: 8.7–9.7), and Mg (magnesium oxide: 12–13) (Adair et al. 2001; Wood et al. 1990). The strong acidic conditions at pH 4.0 could have promoted the positively charged sites of the metal (oxyhydr)oxides and metal-silicate compounds within the engineered biochar and thus promoted the electrostatic attraction forces between the metal (oxyhydr)oxides and the added aqueous P (Yao et al. 2011). These results corroborated those acidic conditions more efficient than alkaline conditions for the EB samples in P removal from aqueous solution. Nonetheless, the low IEP values of the EBMn, EBFe, EBZn, and EBMg biochar samples suggested that the electrostatic attraction between metal oxides in biochar particles and P was unlikely to be the primary retention mechanism for P in the current study. However, chemisorption as revealed by the Freundlich $n$ constant (Table 2) and coprecipitation as shown by SEM analysis (Figs. 2, 3) did control aqueous-P reclamation under the studied conditions.

### 3.3 Regeneration and extractability study of phosphorus-laden biochar

The absolute and relative amounts of P extracted using a citric acid solution are given in Fig. 8. The results showed that the EBMg-P sample had the highest value of citric acid extractable P (2.23 mg/g), followed by the EBZn-P (1.65 mg/g), EBFe-P (1.24 mg/g), EBAl-P (0.45 mg/g), and EBZn-P samples (0.0017 mg/g), respectively. Considering the relative amounts of extracted P, the EBZn-P samples provided the highest P release of 69% of the total P and the other extracted P values decreased in the following order: EBMg-P (65%) > EBMn-P (35%) > EBFe-P (19%) > EBAl-P (0.044%). This demonstrated that EBMg-P remained the best material for both P-adsorbent capacity and P-recycled fertilizer. Furthermore, the results suggested a new potential application of EBZn-P, EBMn-P, and EBFe-P as P-recycled fertilizers. However, the EBAl-P samples had a very low value of citric acid extractable P, reflecting its poor potential use as a P fertilizer.

Furthermore, the concurrent release of the host metals from each engineered biochar through the citric extraction
corresponded to 46.80, 47.47, 34.14, and 0.53%, respectively, of the total host metal concentrations. This suggested a potential use of the Mg-, Zn-, and Mn-biochars for releasing the host metals as plant nutrients, in addition to the P release. However, strong P binding to Fe oxides in the Fe-biochar remained to limit the Fe release and availability.

3.4 Metal-silicate biochar as a possible new recycled fertilizer of phosphate and micronutrients

Many studies on engineered biochar from different biomass materials with Al, Fe, Mn, or Mg showed that metal (oxyhydr)oxides were the main components in the designed biochar, including boehmite in Al-biochar.

Fig. 6 Phosphorus adsorption isotherms of engineered biochars produced from rice husk treated with Al (EB_{Al}), Fe (EB_{Fe}), Mn (EB_{Mn}), Zn (EB_{Zn}), and Mg (EB_{Mg}) (a–e, respectively) compared to pristine biochar (PB, f). Symbols are for experimental data, and error bars show the standard error of triplicates.
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maghemite in cottonwood Fe-biochar (Zhang et al. 2013), manganosite in loblolly pine Mn-biochar (Wang et al. 2015), brucite in tomato leaf Mg-biochar (Yao et al. 2013) and periclase in Mg-biochar from several agricultural wastes namely tomato leaves, sugar beet tailings, sugarcane bagasse, cottonwood, pinewood, and peanut shells (Yao et al. 2013; Zhang et al. 2012). Our data obtained using the XRD and SEM–EDS techniques for engineered biochar rice husk provided new insight into the role of silicate minerals in the biomass as contributing substantially to the formation of the metal-silicate compounds in the engineered biochar (Figs. 2, 3), which could act as the primary host for P in the studied mineral-engineered biochar samples. The large variations in the compounds, which were newly formed in the mineral-engineered biochar, strongly controlled their P-sorption capacity and their potential use as P-recycled fertilizers.

All the engineered biochar samples could be used as effective adsorbents for P removal from aqueous solutions with different levels of efficiency and sorption kinetics, suggesting that varying the solid-to-solution ratios should be considered for each engineered biochar to achieve its maximum P-removal effectiveness. Considering the P-sorption maximum capacities, the EBAl and EBMg samples (38.6 and 46.5 mg/g, respectively) retained lower P contents than Al-biochar and Mg-biochar in other studies prepared at the same temperature (600 °C), but with different initial biomass, including cottonwood, sugar beet tailings, tree leaves, and tomato leaves that retained P up to 835 mg/g (Zhang et al. 2012), 135 mg/g (Zhang and Gao 2013), 122 mg/g (Luo et al. 2021), and 100 mg/g (Yao et al. 2013), respectively. The P-retention capacity of the present Fe-biochar (31.7 mg/g) was comparable to the Fe(III) biochar from

| Sample  | Kinetic model       | Parameter 1* | Parameter 2* | R²  |
|---------|---------------------|--------------|--------------|-----|
| EBAl    | Zero order          | k=0.12       | q₀=15.3      | 0.66|
|         | First order         | k=0.0063     | q₀=15.3      | 0.67|
|         | Parabolic diffusion | a=1.2        | b=13.2       | 0.77|
|         | Power function      | a=0.12       | b=13.4       | 0.84|
|         | Elovich model       | a=5.03       | b=13.0       | 0.79|
| EBFe    | Zero order          | k=0.10       | q₀=8.9       | 0.81|
|         | First order         | k=0.0089     | q₀=8.3       | 0.74|
|         | Parabolic diffusion | a=1.0        | b=7.1        | 0.92|
|         | Power function      | a=0.17       | b=7.4        | 0.90|
|         | Elovich model       | a=4.2        | b=7.0        | 0.92|
| EBMn    | Zero order          | k=0.30       | q₀=6.9       | 0.91|
|         | First order         | k=0.022      | q₀=5.9       | 0.75|
|         | Parabolic diffusion | a=3.0        | b=1.8        | 0.99|
|         | Power function      | a=0.44       | b=4.1        | 0.99|
|         | Elovich model       | a=12.0       | b=1.7        | 0.96|
| EBZn    | Zero order          | k=0.16       | q₀=4.9       | 0.78|
|         | First order         | k=0.023      | q₀=2.6       | 0.49|
|         | Parabolic diffusion | a=1.7        | b=1.9        | 0.91|
|         | Power function      | a=0.51       | b=2.1        | 0.86|
|         | Elovich model       | a=7.3        | b=1.4        | 0.99|
| EBMg    | Zero order          | k=0.55       | q₀=4.4       | 0.79|
|         | First order         | k=0.049      | q₀=0.77      | 0.59|
|         | Parabolic diffusion | a=5.6        | b=5.6        | 0.91|
|         | Power function      | -            | -            | -   |
|         | Elovich model       | a=23.5       | b=-6.6       | 0.94|

*aCoefficients in kinetic equations: zero-order equation ($q_t = -kt + q_0$), first-order equation ($\ln q_t = -kt + \ln q_0$), parabolic diffusion ($q_t = at^{0.5} + b$), power function ($q_t = br^a$), and Elovich model ($q_t = a \log r + b$)

Fig. 7 Absolute (a) and relative percentage (b) of P adsorption concentration of engineered biochars derived from rice husk treated with Al, Fe, Mn, Zn, and Mg (EBAl, EBFe, EBMn, EBZn, and EBMg) at pH solution values of 4, 6, and 8. Bar graphs indicate mean values of experimental data with error bars showing standard error.
rice straw of 32.6 mg/g (Wu et al. 2020). No evidence has been found regarding the P-sorption abilities of Mn- and Zn-biochars. The current study showed that EB_{Fe} and EB_{Mn} retained P at levels as high as EB_{Al}, but EB_{Zn} removed the lowest P amount of 25 mg/g. The main difference between the P retention capacities in the current study and those of other studies could be mainly attributable to the different chemical compositions of the raw biomass and the designed metals, which could have resulted in the substantial variations in the sizes and shapes of the metal-silicate precipitates in the biochar structures (Figs. 2, 3). Compared to other adsorbents such as montmorillonite-biochar and Zn/Al-LDH-biochar composites, the mineral-engineered biochars in the current study (25.6–46.5) showed 2–5 times less P adsorption capacity than other adsorbents such as montmorillonite-biochar and Zn/Al-LDH-biochar composites with P retention of about 150 mg/g (Chen et al. 2017; Yang et al. 2019a).

Furthermore, the metal-silicate mixtures observed in the SEM investigation were important host phases for P in the biochar produced from Si-rich rice husk. The dissolved silicates from the Si-rich material could be coated onto the reactive surfaces of the metal (oxyhydr)oxides or coprecipitated with the host metals and thereby could shift the IEP of the metal (oxyhydr)oxides downward and close to the IEP for quartz (Fig. 5) (Essington 2015). This process resulted in low magnitudes of positive charge sites in the designed metal (oxyhydr)oxides for P and other oxyanion retention (Hiemstra et al. 2007; Schwertmann and Fechter 1982). In addition, a study on the competitive effect of silicates on arsenite (As(III)) and arsenate (As(V)) retention on the hematite surface showed that a higher degree of Si polymerization onto the hematite induced by the longer residence time lowered As(III) and As(V) adsorption on hematite (Christl et al. 2012). The more pronounced effect on the PZC of the metal (oxyhydr)oxides could prevail if the dissolved silicates were coprecipitated with metal (oxyhydr)oxides (Schwertmann and Fechter 1982). This coprecipitation mechanism was likely to be the primary mechanism in the transformation in the surface charge behavior and anionic binding capacity of the metal (oxyhydr)oxides in the current study because the particles of the metal-silicate mixtures under SEM examination typically occurred as discrete particles rather than as Si coated onto the metal (oxyhydr)oxides. The low IEP values of the Fe-, Mn-, Zn-, and Mg-engineered biochars were consistent with the metal silicate formation (Figs. 2, 3), presumably through coprecipitation mechanisms. The EDS analysis of the mineral grains under the SEM investigation also revealed the higher Si content in the mineral grains and the lower host metal content (Fig. 9a). Moreover, the more elevated the Si content, the lower the P content in the mineral grains (Fig. 9b). These data confirmed that silicate in the biomass could react with the added host metals and form metal-silicate precipitates that lowered the available sites for P sorption and retarded its sorption kinetics. Therefore, the roles of silicates importantly affected both the retention and release of anions by the metal (oxyhydr)oxides.
Most P-laden biochar samples (except for the EB_{Al-P} samples) released considerable amounts of P and host metals (Fe, Mn, Zn, and Mg) when extracted using 2% citric acid (Fig. 8, Table S2). The EB_{Zn-P} samples released the highest P concentration (69%), followed by the EB_{Mg-P} (65%), EB_{Mn-P} (35%), and EB_{Fe-P} (19%) samples. Notably, other studies reported that only 4.4% of the total P (10.5 mg/g from 239 mg/g) could be extracted using citric acid from corn Mg-biochar (Fang et al. 2014), 14% of the total P (8.9 mg/g from 65 mg/g) extracted by KCl from oak wood Mg-biochar (Takaya et al. 2016), and 7.6% of the total P (7.6 mg/g from 100 mg/g) extracted by Mehlich-3 from tomato leaf Mg-biochar (Yao et al. 2013). The higher extractable P content from the P-laden biochar in the current study than in other studies could be attributed to the metal-silicate precipitates that could have weakened the binding energy for P on the designed metal oxides. The metal-silicate precipitates enhanced the P extractability from EB_{Zn}, EB_{Mn}, and EB_{Fe} that are strong adsorbents for P. More importantly, other studies reported that 20–30% (Hedley et al. 1988) or 10–43% (Santos et al. 2019) of the total P dissolved from rock phosphate was extracted using 2% citric acid and provided an excellent indication of the high reactivity of rock phosphates (Hedley et al. 1988; Santos et al. 2019).

Although the EB_{Al-P} sample released the lowest P concentration (0.044% of the total P), a recent study on a sand-EB_{Al-P} mixture revealed a sufficient amount of Olsen extractable P (11–12 mg/kg) constantly released from the mixture throughout 45 days of incubation (Sornhiran et al. 2021). This result suggested the possible use of Al-biochar as both P-recycled fertilizer in sandy soils with a low P availability and a high risk of P loss.

The current results highlighted the potential of mineral-engineered biochar with abundant Si as both adsorbents for P removal and subsequent reuse as P-recycled fertilizers. Furthermore, the release of the host metals (Mg, Zn, and Mn) from each engineered biochar through extraction suggested the additional advantage of the mineral-engineered biochar in the current study for supplying secondary (Mg) and micronutrient (Zn and Mn) fertilizers in addition to the recycled P. However, further studies should examine the kinetics and suitable chemical conditions for releasing P and micronutrients from the mineral-engineered biochar.

### 4 Conclusions

This work investigated the retrieval and reuse of P from aqueous solution using mineral-engineered biochars. Silicon-rich rice husk was used as the raw material for the novel design of Al-, Fe-, Mn-, Zn-, and Mg-(oxyhydr)oxide–biochar composites. The silicate compounds in the raw biomass induced the formation of metal-silicate compounds in the biochar framework and acted as the main absorbents for P. All mineral-engineered biochars caused slow sorption kinetics but offered reasonable sorption ability to P in the aqueous solution because the metal-silicate compounds had lower chemically reactive surfaces for P than the pure metal (oxyhydr)oxides typically found in diverse engineered biochars from other materials with less silicon content. The silicate compounds could weaken the P-binding capacity through adsorption onto or coprecipitation with metal (oxyhydr)oxides in the engineered biochar. Therefore, the silicate compounds showed the greater significance and novelty of the metal-silicate biochars as P-recycled fertilizer than the mineral-engineered biochar in other studies where strong binding capacity makes it difficult to release P for plant utilization.

![Fig. 9 Bivariate plots of Si content with host metal (a) and P (b) contents of mineral grains within mineral-engineered biochars based on SEM–EDS investigation](image-url)
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Availability of data and materials All data generated or analyzed during this study are included in this published article and its supplementary information files.

Declarations

Competing interests The authors have no conflicts of interest to declare that are relevant to the content of this article.

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