Phosphate Fixation and P Mineralogy on Natural and Ca-Modified Zeolites During Simultaneous Nutrient Removal

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Abstract The recovery and recycling of nutrients (N&P) from wastewater are one of the major topics to save primary energy and resources, to raise the efficiency of wastewater treatment plants, and to foster a future circular economy. In the present study, the removal of ammonium ($NH_4^+$) and phosphate ($PO_4^{3-}$) using natural and Ca-treated zeolite is investigated in detail. Special emphasis is put on the simultaneous removal of both species from model solutions followed by elaborate mineralogical analyses (XRD, EPMA, FEG-SEM) for zeolite characterization and in order to determine the type, structure, and crystal sizes of CaP-phases precipitating on the zeolites surface. The effectivity of the phosphate segregation and chemical composition and the crystalline structure of the CaP-phase precipitating on the surface of the zeolite depend on the physico chemical conditions in particular on pH, molar ratio of Ca and P (due to zeolite modification), and the presence of $NH_4^+$. Results of simultaneous removal experiments of N&P revealed that Ca pretreatment enhances P segregation and increases the obtainable P-loadings of Ca–zeolites. Maximum P-loadings of 25 mg g$^{-1}$ Ca–zeolite in binary solutions containing both ammonium and phosphate were obtained. Simultaneous phosphate removal by surface precipitation of CaP-phases does not significantly influence ammonium ion exchange and the type of CaP-precipitates formed on the zeolite surface is assumed to be mainly brushite and apatite.

Keywords Surface precipitation · Ammonium removal · Clinoptilolite · Wastewater treatment

1 Introduction

In the actual discussions of promoting renewable energy, biogas and sewage treatment plants try to enhance their capacity for anaerobic digestion to produce green fuels (e.g., biomethane). This leads to increasing amounts of liquid digestates and sludge liquor, which contain ascending concentrations of dissolved ammonium and phosphorous due to intensified co-substrate fermentation. Legal regulations (maximum permissible values) and technological issues (ammonium inhibition) necessitate the reduction of both dissolved species by certain wastewater treatment methods. Nowadays, $NH_4^+$ and $PO_4^{3-}$ are usually removed separately by biological and chemical processes. All of these standard processes are expensive and do not offer the possibility to recover, concentrate, and utilize $NH_4^+$ and $PO_4^{3-}$ for example as nutrients in fertilizers. In contrast to these enormous amounts of N and P that are lost via global wastewater treatment systems, the food supply of an
increasing world population strongly depends on the availability of these nutrients for crop cultivation.

Phosphorus is one of the building blocks of life and essential for many cellular processes and DNA synthesis. Additionally, no chemical component can substitute P in the life cycle and therefore P is irreplaceable (Cordell et al., 2009). By now, nearly three times the amount of the natural P cycle is mined per year from phosphate rock for anthropogenic needs (e.g., for fertilizer in agriculture and chemical and pharmaceutical products) (Vučić et al., 2021). In this human-affected cycle, P is lost at all stages of the processing systems (Daneshgar et al., 2018). After Kok et al. (2018); 3.7 Mt a\(^{-1}\) of P deriving from food is discarded via wastewater, where it is currently lost.

A similar picture can be observed for nitrogen, but even at a larger scale: since the invention of the Haber Bosch process in the beginning of the twentieth century, which enables the synthesis of NH\(_3\) from natural gas and atmospheric N\(_2\), nitrogen is extensively used as fertilizer. The global nitrogen cycle accounts for \(\sim 400 \text{ Mt a}^{-1}\), with approximately 50% thereof deriving from anthropogenic activities (Fowler et al., 2013).

Given these values, the recovery and utilization of NH\(_4^+\) and PO\(_4^{3-}\) are of great interest considering sustainability and closing of material cycles. As phosphorous is also one of the critical raw materials identified by the EU (European Commission, 2014), its recovery would contribute to a sustainable raw material supply. Currently, many countries throughout Europe promote the use of secondary phosphates (e.g., Switzerland and Germany) by making P-recovery mandatory for wastewater treatment plants that exceed defined sizes, e.g., >50,000 population equivalent in Germany (Bundesministerium für Umwelt, Naturschutz und nukleare Sicherheit (BMU), 2017). Sweden and Austria are on the same path and committed themselves to put P-recovery into national legislation in the near future. Potential technologies for P-recovery comprise the precipitation of dissolved PO\(_4^{3-}\) from the liquid phase as magnesium ammonium phosphates (MAP) or MAP/calcium phosphate minerals (Law & Pagilla, 2018) as well as recovery from sewage sludge ash after incineration. The latter approach has some serious constraints like the costly technological facilities needed for monocombustion, storage and ash processing, and the production of certain amounts of (toxic) by-products. Hence, technologies for liquid phase recovery of P (and N) are largely investigated, under which ion-exchange and adsorption processes on natural and modified zeolites have become an interesting topic (Yin & Kong, 2014; Zhang et al., 2011). In contrast to synthetic zeolites, natural zeolites present an attractive option for applications in water purification due to their high abundance, availability, and low costs (Alshameri, Ibrahim, et al., 2014; Guaya et al., 2015b; Malekian et al., 2011; Zhang et al., 2011).

So far, natural zeolites have been rarely used for anion sorption due to the overall negative charge on their surface, caused by their crystal structure (Jiang et al., 2013). For anion and especially phosphate removal, modification of the surface of the zeolite is necessary. This is usually done by incorporation or impregnation of metal oxides and hydroxides in the zeolite pore structure (e.g., Fe, Al, La, and other rare earth elements; Alshameri, Yan, et al., 2014; Jiménez-Cedillo et al., 2011; Sang et al., 2020; Simsek et al., 2013), the replacement of natural cations by the aforementioned metal ions via selective ion exchange (Guaya et al., 2015b), or the use of special, metal-incorporated zeolites synthesized from fly ashes (e.g., Hermassi et al., 2020; Xu et al., 2022). The process of phosphate removal is usually related to surface precipitation of various insoluble chemical species, like monodentate and bidentate complexes on incorporated iron hydroxides inside the zeolite micropores (Guaya et al., 2015b).

Several studies have been conducted for the simultaneous removal of NH\(_4^+\) and PO\(_4^{3-}\) from aqueous solutions by modified zeolites; many of them are based on synthetic zeolites, fly-ash zeolites, or mixtures of reactive sorbents (e.g., Gao et al., 2018; He et al., 2016; Hermassi et al., 2018; Salam et al., 2021; Zhang et al., 2007). Guaya et al. (2015b) modified natural clinoptilolite with Fe(III)-ions to promote the formation of hydroxyl groups (\(\equiv\text{Fe-OH}\)). Although simultaneous removal of NH\(_4^+\) and PO\(_4^{3-}\) was found to be possible, the NH\(_4^+\) exchange capacity was slightly decreased by the competing ions and surface precipitation of various MAP and CaP- minerals (blocking of pores). Furthermore, Guaya et al. (2015a) studied adsorption on a natural zeolite modified with hydrated aluminum oxide and increased PO\(_4^{3-}\) sorption from 0.6 to 7 mg-P g\(^{-1}\) while NH\(_4^+\) sorption capacity was slightly decreased from 33 to 30 mg-N g\(^{-1}\). In both studies, recovery of phosphate
was found to be possible, but a reduction of phosphate uptake after several regeneration cycles was observed (Guaya et al., 2015a). Other studies show that simultaneous removal of $\text{NH}_4^+$ and $\text{PO}_4^{3-}$ with zeolites modified by Fe, Al, Mn, and La is possible for simulated wastewaters (e.g., Li et al., 2005; Ning et al., 2008; Li et al., 2009 and Huo et al., 2012). Performance depends on the competing ions in the applied wastewater and the pre-treatment. Simultaneous recovery of nitrogen and phosphorus was achieved by natural zeolites from sludge fermentation liquid in a study of Wan et al. (2017). Regeneration of expended zeolites can be carried out by $\text{H}_2\text{SO}_4$ (e.g., Choi et al., 2014) or NaOH-solutions (e.g., Guaya et al., 2015a). Generally, regeneration performance and products thereof depend on the type of zeolite surface modification and the phosphorous removal mechanism.

Some studies regarding P-removal with Ca-modified zeolites were published (e.g., Hermassi et al., 2016; Schick et al., 2012), but so far, Ca-modified zeolites were only rarely used for simultaneous N&P-removal from different media. Sun et al. (2011) made sorption experiments with CaCl$_2$-modified zeolite in aqueous solution. Phosphate removal efficiencies were pH-dependent, with maximum values between 7 and 9. They stated that the mechanism of ammonium adsorption on the modified zeolite was ion exchange, whereas the removal of phosphate was a matter of chemical precipitation. You et al. (2017) investigated Ca- and Mg-modified zeolites for the removal of $\text{NH}_4^+$ and $\text{PO}_4^{3-}$ from simulated treated wastewater effluents. They measured maximum sorption capacities for a Ca-treated zeolite of 123.1 ± 9 mg $\text{NH}_4^+$ g$^{-1}$ and 119.5 ± 7.5 mg $\text{PO}_4^{3-}$ g$^{-1}$ in an ammonium/phosphate binary system. Sorption mechanisms involved ammonium ion exchange and precipitation of Ca and Mg phosphates or mixed ammonium-magnesium phosphate minerals like apatite, brusite (CaHPO$_4$.H$_2$O), and struvite (NH$_4$MgPO$_4$.6 H$_2$O). Mitrogiannis et al. (2017) reported that Ca(OH)$_2$-treated zeolite can be effectively used for $\text{PO}_4^{3-}$ removal from aqueous solutions under low P concentrations (0.5–10 mg L$^{-1}$) with the predominant mechanisms for P-removal being ligand exchange and Ca-P surface precipitation. In subsequent studies, they applied Ca(OH)$_2$-treated zeolites for P-removal from urine (Mitrogiannis et al., 2018) and olive mill wastewater (Mitrogiannis et al., 2020), where maximum P removals of 73.9 and 85.9% for two different wastewater samples were achieved.

The present study further investigates the phosphate removal via natural and Ca-modified zeolites (clinoptilolite) by applying an elaborate set of chemical and mineralogical analyses for the liquid solutions as well as the solid materials. The type of P-phases formed on the applied zeolites is further investigated and sorption capacities in binary aqueous solutions containing $\text{NH}_4^+$ and $\text{PO}_4^{3-}$ are presented in order to improve simultaneous ammonium and phosphate removal and recovery from real effluents in the ILS process (Ellersdorfer, 2018) and other potential future applications.

### 2 Materials and Methods

Sorption experiments for single and simultaneous phosphate segregation were conducted using natural and Ca-modified natural zeolite. For Ca-modification, natural zeolite from Eastern Slovakia was treated with CaCl$_2$ solution (0.5 M; pH 10) for 24 h in columns using a volume of 20 L of treatment solution and 2.3 kg of natural zeolite (grain size 1.5–2 mm). Diameter of the column was 120 mm and flow rate 2.5 L min$^{-1}$. To ensure a complete submersion of the zeolite, the modification process was run in up flow mode. After modification, the zeolite was flushed with deionized water until pH 7 was reached. The modified zeolite was dried at 105 °C and stored in hermetically sealed bottles until sorption experiments.

Screening and single phosphate sorption experiments were conducted in batch operation with dibasic sodium phosphate dodecahydrate solutions (Na$_3$PO$_4$.12 H$_2$O) in a concentration range from 500 to 2500 mg PO$_4^{3-}$ L$^{-1}$. For preliminary screening experiments, a concentration of 2500 mg PO$_4^{3-}$ L$^{-1}$ was used. For simultaneous sorption from binary solutions (N&P), an additional amount of 1000 mg L$^{-1}$ of $\text{NH}_4^+$ was added to each phosphate solution using ammonium sulfate (($\text{NH}_4$)$_2$SO$_4$). If necessary, pH-values were adjusted with sulfuric acid (H$_2$SO$_4$; 96%). The batch system used for sorption experiments was the same for both experimental approaches: 500 mL of initial single/binary solution was treated with 20 g of natural and Ca-modified zeolite for 24 h in an overhead shaker at 20 °C respectively. In order
to avoid abrasion of the material due to agitation, the zeolite was packed into cotton bags.

Liquid samples were taken before and after the sorption experiments and filtrated by syringe filters (0.22 µm). Ammonium was detected via Kjeldahl method using boric acid and HCl (0.1 M)/Tashiro’s indicator for titration. For phosphate analysis, the photometer method was used (Merck Spectroquant® 114848). The difference in $\text{PO}_4^{3-}$ and $\text{NH}_4^+$ concentration of the initial and final solution represents the amount of removed phosphate and ammonium and therefore the effective segregation capacity for different equilibrium concentrations which yields to the sorption isotherm under the given conditions.

For data interpretation, the equilibrium loadings ($q_{\text{PO}_4^{3-}/\text{NH}_4^+}$) were evaluated by the amount of adsorbed phosphate/ammonium per unit mass of zeolite (in mg $\text{PO}_4^{3-} \text{g}^{-1}$ zeolite or mg $\text{NH}_4^+ \text{g}^{-1}$ zeolite), calculated according to the following equation:

$$q_{\text{PO}_4^{3-}/\text{NH}_4^+} = \frac{(C_0 - C_e) \cdot V}{m}$$

where $C_0$ and $C_e$ are the phosphate/ammonium concentrations (mg L$^{-1}$) of the stock solutions at initial and final stage of the experiment, $V$ is the total solution volume (L), and $m$ is the mass of the zeolite (g).

Mineralogical investigations were done before and after the modification step and at the end of the segregation experiments using dried samples (105 °C), which were sealed in plastic containers and stored for subsequent solid analysis. The material was characterized by X-ray diffraction (XRD) as well as electron microprobe (EPMA) and scanning electron microscopy (FEG-SEM). XRD was conducted with a PANalytical X’Pert Pro, Cu K$\alpha$, $\lambda = 1.541$ nm at the Chair of Petroleum Geology, Montanuniversität Leoben. XRD patterns were evaluated with the software X’Pads with its attached ICDD databases. Electron microprobe technique was used for chemical mapping using a Superprobe Jeol JXA8200 instrument at the Eugen F. Stumpf Microprobe Laboratory provided by UZAG at Montanuniversität Leoben. Element mappings were made from carbon-coated polished sections. Morphology and surface studies of the zeolite before and after the sorption experiments were undertaken using FEG-SEM technique (LEO 1525, Carl Zeiss SMT at Erich Schmid Institute of Material Science, Leoben). Zeolite grains were sticked on a carbon tab placed on an aluminum stub and coated with a carbon layer. BSE high resolution microphotographs of the zeolite were taken at an acceleration voltage of 5 kV.

### 3 Results and Discussion

#### 3.1 Characterization of the Untreated and Treated Raw Material

The XRD spectra represent the whole rock mineral composition of both the untreated and Ca-treated zeolitic rock (Fig. 1). The X-ray diffraction pattern of the untreated zeolite identifies a HEU-type zeolite (clinoptilolite) as the main mineralogical phase in the raw material. Minor phases include biotite (annite?), plagioclase (dominantly andesine), alkali feldspar, quartz, and cristobalite. Minor amounts of plagioclase and biotite differ slightly between the untreated and treated zeolite but do not indicate significant changes in the mineralogical composition of the different samples nor mineral transformation from pre-treatments (Vollprecht et al., 2019). The different cation loading after the treatment has no significant impact on the peak position of the clinoptilolite. A slight enhancement of the background signal between 20 and 30° may indicate minor amounts of amorphous tuffic glass (Ostrooumov et al., 2012).

The mean chemical composition ($n=22$) of the clinoptilolite obtained by EPMA and the cations per formula units were calculated based on 72 oxygens, deriving from the ideal formulas of HEU type zeolites (Heulandite (Na, K)Ca$_4$(Al$_8$Si$_{28}$)O$_{72}$:24 H$_2$O and clinoptilolite (Na, K)$_6$(Al$_6$Si$_{30}$)O$_{72}$:20 H$_2$O) and are listed in Table 1.

| Element | Formula |
|---------|---------|
| Si      | $\text{Si}_{4+}$ |
| Al      | $\text{Al}^{3+}$ |
| Na      | $\text{Na}^{+}$ |
| K       | $\text{K}^{+}$ |

Si and Al are assigned to the tetrahedral site and Na, K, Ca, and Mg to the exchangeable cation sites. Spots for the WDS analysis were selected using back scattered electron (BSE) images and EDS spectra, in order to avoid contamination by iron hydroxides and to distinguish minor minerals (quartz and feldspar) from the zeolite matrix.

#### 3.2 Sorption Experiments

Results from preliminary screening experiments for phosphate segregation show that the $\text{PO}_4^{3-}$-loading...
on natural zeolite accounts for 12.3 mg PO$_4^{3-}$ g$^{-1}$ zeolite (Table 2). The pH value of the initial phosphate solution (~2500 mg PO$_4^{3-}$ L$^{-1}$) slightly decreases from 12.23 to 11.93 after 24 h of contact with the natural zeolite sample. Adjustment of the pH-value of the initial solution to 8.5 leads to a small increase in the phosphate removal capacity (14.5 mg PO$_4^{3-}$ g$^{-1}$) and a larger decline in the pH-value during 24 h of PO$_4^{3-}$-segregation (8.5 to 7.67) as a consequence of the logarithmic pH-scale. Phosphate loading of Ca-pretreated zeolite (6.5 mg PO$_4^{3-}$ g$^{-1}$) is only half of the loading of natural zeolite at high pH-values (~12) but is more than doubled when the pH-value is adjusted to values <8. With a value of 13.1 mg PO$_4^{3-}$ g$^{-1}$, the final loading of Ca-pretreated zeolite after 24 h is higher than the value of untreated zeolite without pH-adjustment. These results emphasize the importance of the pH-value for phosphate removal by the formation of CaP-phases. In any of the conducted experiments, the pH-value of the loading solution is

![Fig. 1 X-ray diffraction (XRD) patterns of zeolites with different cation loading (bt: biotite, cpt: clinoptilolite, plg: plagioclase, qz: quartz, crs: cristobalite, or: orthoclase)](image)

Table 1 Mean chemical composition of zeolite minerals obtained by EPMA (calculations based on 72 oxygen atoms, apfu atoms per formula unit)

|               | Si  | Al  | Na  | K   | Ca  | Mg  |
|---------------|-----|-----|-----|-----|-----|-----|
|               | $n$ | [apfu] | [apfu] | [apfu] | [apfu] | [apfu] | [apfu] |
| Natural zeolite | 22  | 30.53 | 5.93 | 0.08 | 0.44 | 1.61 | 0.14 |
| Ca-treated zeolite | 13  | 30.43 | 5.63 | 0.05 | 0.85 | 1.67 | 0.09 |

Table 2 Phosphate segregation on natural and Ca-pretreated zeolites after 24 h of contact with model solutions (2500 mg PO$_4^{3-}$ L$^{-1}$; Na$_3$PO$_4$.12 H$_2$O) at different pH-values

| Sample         | pH$_{t=0}$ | pH$_{t=24}$ | PO$_4^{3-}$-loading | pH$_{t=0}$ | pH$_{t=24}$ | PO$_4^{3-}$-loading |
|----------------|------------|-------------|---------------------|------------|-------------|---------------------|
| Natural zeolite | 12.23      | 11.93       | 12.3                | 8.5        | 7.67        | 14.5                |
| Ca-treated zeolite | 12.23      | 12.00       | 6.5                 | 7.37       | 7.02        | 13.1                |
lower after phosphate segregation, which is a result of the acidic reactions responsible for the CaP-phase formation.

The single phosphate sorption experiments (Fig. 2; left) reveal a continuous increase of the phosphate loading with a maximum of around 20 mg PO₄³⁻ g⁻¹ at initial solution concentrations of 2500 mg PO₄³⁻ L⁻¹ for both natural and Ca-treated zeolites. Whereas natural zeolite P-loading is higher at solution concentrations <250 mg L⁻¹; Ca-treatment leads to higher phosphate removal capacities between 250 and 1000 mg PO₄³⁻ L⁻¹. Nevertheless, the observed maximum P-loadings are similar, which may be an effect of the establishment of an equilibrium between the dissolved cations in the solution (mostly Na⁺) and the exchangeable ions on the zeolite (mainly Ca²⁺; Table 1), which is limited by the amount of available cations in the solution.

The addition of ammonium changes the phosphate sorption capacities of natural zeolite and leads to significantly higher P-loadings up to a concentration of 1000 mg PO₄³⁻ L⁻¹ (Fig. 2; right). Nevertheless, the maximum P-loading of natural zeolite at 2500 mg PO₄³⁻ L⁻¹ in binary solution is similar to that of the single phosphate experiment (~20 mg PO₄³⁻ g⁻¹). The P-loadings of Ca-modified zeolite reveal significantly higher values over the whole concentration range with a maximum of 25 mg PO₄³⁻ g⁻¹ at 2500 mg PO₄³⁻ L⁻¹. These results indicate that the P-loading capacity of the natural zeolite is affected by Ca-limitation at higher P-concentration levels, where the natural zeolitic Ca²⁺-ions are already expended

and the additional ammonium-ions in the loading solutions cannot deliberate extra Ca-ions. Ca pretreatment increases the available amounts of Ca²⁺ in the zeolite structure and cations like NH₄⁺ can mobilize these additional Ca-ions in order to increase Ca-phosphate precipitation.

Ammonium sorption for natural and Ca-treated zeolite is between 12 and 16 mg NH₄⁺ g⁻¹ and tends to increase slightly with rising P-concentrations. As indicated by the similar values of NH₄⁺-loading on both zeolites, the ammonium ion exchange is hardly influenced by the formation of CaP-phases, although a small decrease in the NH₄⁺-loading can be observed for the maximum phosphate binding zeolite (Ca-treated zeolite at 2500 mg L⁻¹ PO₄³⁻). In comparison to single ammonium sorption experiments (1000 mg NH₄⁺ L⁻¹ without PO₄³⁻-addition; dotted lines in Fig. 2), NH₄⁺-uptake during simultaneous operation is in the same range as in single ion exchange experiments, which is another proof that ammonium exchange is not negatively influenced by CaP-precipitation (e.g. blockage of pores).

3.3 Identification of the Solid Phases After Precipitation

Several different techniques were applied in order to identify the precipitated mineral phase on the zeolites surface. Observation with FEG-SEM technique gives information about the morphological aspects like crystal form, shape, size, and grade of crystallinity. Analyses with EPMA technique provided information

![Fig. 2 Results of P (left) and N&P-sorption experiments (right) as a function of initial PO₄³⁻-concentrations in the solution](https://example.com/figure2.png)
about the chemical composition of the precipitate. The formation of the precipitates was observed only in pH-adjusted solutions combined with the Ca-treated zeolite.

A BSE image of the cross section after sorption experiments presented in Fig. 3 shows the typical texture of a zeolitic tuff as it was used in this study. Coarser grained, partially idiomorphic mineral phases are embedded in a finer grained matrix. The main different components distinguishable by different gray scales were identified by EDS and consist of clinoptilolite, plagioclase, K-Feldspar, and minor amounts of quartz and biotite. Clinoptilolite occurs thereby as distinct grains filling pore spaces, extremely fine-grained matrix material, and clinoptilolite pseudomorphs of former glass shards representing its volcanic origin. This is consistent with the mineral composition obtained by XRD (Fig. 1).

Mapping of the elemental distribution by EPMA in the same area is shown in Fig. 4a–f. Aluminum and silicon are revealed in all regions of the zeolite grain as they are considered as the ions in the tetrahedral sites of zeolite crystal lattice. K, Na, and Ca as exchangeable cations of the zeolite are also distributed over the whole cross section. Distinct higher amounts of Ca, Na, and Al in some regions of the cross section can be explained by feldspar crystals which occur in the fine-grained matrix of the zeolite rock. High concentrations of calcium and phosphorous are less uniformly distributed. Notably, phosphorous is largely absent in the center of the grain but is enriched on the rim of the zeolite forming the precipitate together with calcium (Fig. 4e and f).

BSE pictures from polished cross sections obtained during EPMA in Fig. 5 reveal details of zeolite grains after precipitation at higher magnification; an accumulation of fine-grained CaP-phases especially in small cavities and niches along the rim of the zeolite grain is particularly visible and shows two different types of crystal shapes occurring as described in detail below.

Precipitates of calcium phosphate (CaP) produced during pH-adjusted screening experiments were observed using FEG-SEM technique following the protocol described in the “Materials and Methods” section. The CaP-crystals shown in Fig. 5 are representative of precipitates produced at pH 7 during 24 h sorption from a 2500-mg L⁻¹ solution (Table 2). The experiments resulted in the formation of acicular crystals up to a grain size of 5 µm (Fig. 6a–c) and platy, radiating rose-like aggregates of tabular crystals up to a size of 100 µm (Fig. 6d–f). Due to crystal habit and crystal form, it can be assumed that these are tabular crystals of monoclinic

![Fig. 3 Photomicrograph of a representative cross section of the zeolite obtained by EPMA showing rock fabric and mineral assemblage (bt: biotite, cpt: clinoptilolite, plg: plagioclase, qz: quartz, or: orthoclase)](image-url)
brushite with perfect cleavage in [001] and [010]. Besides, brushite acicular crystals, especially contact twinned, can also be assumed as hydroxylapatite.

In summary, the simultaneous removal of ammonium and phosphate is possible both with natural and Ca-modified zeolites. The applied method of Ca pretreatment with CaCl₂-solutions, which increases the Ca-loading and subsequently also the P-removal efficiency, can be executed by simple immersion of the zeolite grains in liquid CaCl₂-solutions without several ion-exchange stages (e.g., Guaya et al., 2015a, b) subsequent drying, calcination, or other (thermal) treatments. This is of specific interest for the further development of the ILS-process, where columns filled with zeolite grains (1–3 mm) together with sodium hydroxide solutions are used for ammonium removal via acid scrubbing. CaCl₂-solutions can be applied for a simple in-line treatment of zeolite grains to remove phosphate from wastewater in a subsequent extra-column, which is loaded with CaCl₂ and recovered by acid flushing from time to time. Hence, the ILS-process itself is not changed but only expanded by an additional column used for P-removal, which enables simultaneous removal but selective recovery of N&P in order to recycle these valuable nutrients.

Fig. 4 a–f Elemental distribution obtained by EPMA of the cross section of a representative zeolite grain after sorption experiments with 2500 mg L⁻¹ PO₄³⁻ for 24 h and adjusted pH to 7: a silicon map, b aluminum map, c sodium map, d potassium map, e calcium map, and f phosphorous map; brighter regions correspond to higher amounts of the respective element.
Conclusions

Results from the sorption experiments show that Ca-treated natural zeolite can be used to simultaneously remove ammonium and phosphate from aqueous solutions. Natural zeolitic calcium is exchanged with sodium ions in \( \text{Na}_3\text{PO}_4 \)-solutions leading to the precipitation of free \( \text{PO}_4^{3-} \) as CaP-phases on the zeolite. The same effect can be observed in presence of additional \( \text{NH}_4^+ \), which enhances CaP-precipitation due to higher cation concentrations and ammonium selectivity, which deliberates more Ca-ions, if they are available (Ca-treatment). For natural zeolites, Ca-limitation impedes CaP-precipitation at higher \( \text{PO}_4^{3-} \)-concentrations; wherefore, the following conclusions can be drawn:

- P segregation using zeolites is feasible and similar for natural and Ca-pretreated zeolites in phosphate solutions.
- Ca pretreatment enhances P segregation and increases the obtainable P-loadings of Ca–zeolites.
- P segregation is higher in solutions containing additional \( \text{NH}_4^+ \) (binary solutions) for both natural zeolites and Ca-treated zeolites as long as natural zeolitic Ca-ions are sufficiently available.
- Ca-treatment significantly increases P-loadings in binary solutions with higher \( \text{PO}_4^{3-} \)-concentrations (>100 mg L\(^{-1}\)) and maximum P-loadings of 25 mg g\(^{-1}\) Ca–zeolite were measured.
CaP-phase formation does not significantly influence ammonium ion exchange in the experimental setup used for this study.

The type of CaP-precipitates formed on the zeolites surface is assumed to be brushite and apatite.

This leads to the assumption that the natural Ca-loading of the applied zeolite is not sufficient for a complete CaP-precipitation and P-loadings are limited by the available Ca-cations in the zeolite structure. The effectivity of the phosphate segregation as well as the chemical composition and crystalline structure of the CaP-phases precipitating on the surface of the zeolite depends on the physicochemical conditions in particular on pH, molar ratio of Ca and P (influenced by zeolite modification), and the presence of NH$_4$$^+$.

For the effective recovery of phosphate from real effluents of wastewater treatment plants, pH-values and cation concentrations have to be considered due to their influence on the amount and mineralogical type of precipitates and therefore on the method used for selective P-recovery in the future. In view of a potential application of simultaneous N&P sorption in wastewater treatment, the fixation of the precipitated CaP-phases to the zeolites surface is a major benefit in order to selectively recover phosphorous and avoid the need for subsequent filtration of fine-grained precipitates.

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