Phosphorus sorption capacity of different types of opoka

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Abstract: Phosphorus sorption capacity of different types of opoka. The bedrock opoka has been lately reported as an appropriate reactive media for onsite wastewater treatment systems due to its high phosphorus (P) sorption capacity. However, variations on its chemical composition may affect its reactivity with P, therefore leading to a variable P removal efficiency. In this paper, the P-sorption capacity of three different types of opoka from the region of Miechów, Poland, is reported. According to the silica and carbonate content, opoka samples were classified as light-weight and heavy-weight opoka. When heated over 900°C, opoka showed a very high P-sorption capacity that was well correlated to its Ca content. P-sorption isotherms from batch experiments with an artificial P solution were plotted and fitted to the Langmuir and Freundlich adsorption models. The Freundlich isotherm appeared to model better the P-sorption of light opoka and the Langmuir isotherm of heavy opoka, suggesting different dominating mechanisms of P-sorption by light and heavy opoka.

Key words: calcium; freundlich isotherm; langmuir isotherm; opoka; phosphorus removal; silica; sorption.

INTRODUCTION

Reactive media with high phosphorus (P) sorption capacity are required in wetland and infiltration systems with the purpose of P removal from wastewater (Mann and Bavor, 1993; Zhu et al., 1997; Drizo et al., 1999). There has been extensive research on materials with noticeable P sorption capacity (Sakadevan and Bavor, 1998; Angel, 1999; Johansson, 1999; Johansson Westholm, 2006). Natural calcareous soils and limestone are also known for its P retention ability (Zhou and Li, 2001). The P sorption capacity of a soil is closely related to the content of Al, Fe and Ca as well as soil pH (Sakadevan and Bavor, 1998; Zhu et al., 1997). The bedrock opoka is a Ca rich sedimentary deposit with moderate to high P-sorption capacity. It mainly consists of SiO2 and CaCO3 but also contains significant amounts of Al2O3 and Fe2O3 (Brogowski and Renman, 2004). Depending on the silica and carbonate content there is a large variability of opoka deposits ranging from 37.5 to 52.1% of silica and 34.5–50.4% of carbonate. Thus, opoka can be classified as light-weight (more SiO2) and heavy-weight opoka (more CaCO3) (Brogowski and Renman, 2004). Polish literature gives a wider range of silica content from 17.06 to 51.88% (Bolewski and Turnau-Morawska, 1963). This type of rock can also be classified as geza when the silica dominates.

It is known the ability of opoka to remove P. The process occurs mainly through Ca-P interactions (Johansson and Gustafsson, 2000). Therefore, variations on its Ca content may account
for differences in P-sorption. The P-sorption efficiency increases significantly when heated over 900°C due to the transformation of CaCO₃ into CaO, with an estimated sorption capacity of 119 g P kg⁻¹ material (Brogowski and Renman, 2004).

The objective of this study was (i) to characterize opoka from three different quarries in the region of Miechów, Poland, and (ii) to evaluate its efficiency for P removal from wastewater. The short-term P-sorption was studied in batch experiments and the maximum sorption capacity was estimated through the Langmuir isotherm approximation.

MATERIALS AND METHODS

Opoka was acquired from three different quarries in the region of Miechów, Poland, located 60 km north of Cracow (Tab. 1). The samples were taken at a depth between 2–4 meters. The material was dried in the air and then in the oven at 105°C.

The material was crushed and sieved to different fractions. Triplicate 2 g samples of the powdered fraction of natural opoka (Opk) and opoka heated to 900°C for 1 hour (900Opk), were used for analyse. Extraction with nitric and perchloric acids by heating for 3–4 days and later filtration, prepared the samples for atomic absorption and emission spectrometry. AAS Solar M6 and ICP-AES JY 238 Ultrace were used. The silica content was calculated from the weight of the filter remain after burning it at a temperature of 900 °C. The CaCO₃ content was analysed using the Scheibler’s method. The pH was measured in a 1M KCl solution in a proportion 1:2.5.

Triplicate 0.1 g samples of the powdered opoka heated to 900°C (900Opk) were placed in 50 ml flasks with artificial P-solutions (KH₂PO₄) of different P concentrations (0, 10, 20, 50, 100, 200, 300, 500 mg P·dm⁻³) and then mixed in a rotator at 60 rpm for 1 hour at room temperature (21°C). After that time, the pH was measured and next, the samples were centrifuged at 6000 rpm and then filtered through a 0.45 μm membrane filter for P determination. The P was analysed using the standard ammonium molybdate method (Murphy and Riley, 1962) in a Beckman DU 600 spectrophotometer. The difference between the initial and final P concentration at equilibrium was supposed to be sorbed to the material.

Sorption parameters were calculated using the Langmuir and Freundlich isotherms by plotting graphically the inverse and logarithm, respectively, of the sorbed portion (qs) against the inverse and logarithm of the solute concentration (Cs) at equilibrium, giving a straight line. The Langmuir and Freundlich constants are obtained from the intercept “y” axis and the slope.

| Sample | Quarry | Bearings | Altitude |
|--------|--------|----------|----------|
| Opk1   | Strzeżów | N 50°23′04,8″ EO 20°03′11,6″ | 355 m |
| Opk2   | Cisie (Antolka) | N 50°24′23,4″ EO 20°05′36,7″ | 348 m |
| Opk3   | Widnica | N 50°24′05,0″ EO 20°01′55,0″ | 354 m |
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Langmuir: \[ q_s = \frac{k_L \cdot C_s}{1 + a_L \cdot C_s} \rightarrow \frac{1}{q_s} = \frac{1}{C_s \cdot k_L} + \frac{a_L}{k_L} \]

where \( k_L (\text{dm}^3\cdot\text{g}^{-1}) \) reflects the adsorptivity and \( a_L (\text{dm}^3\cdot\text{mg}^{-1}) \) is related to the energy of adsorption. \( \frac{k_L}{a_L} (\text{mg} \cdot \text{g}^{-1}) \) is obtained from the intercept “y” axis and reflects the maximum adsorption capacity.

Freundlich: \[ q_s = a_F \cdot C_s^{b_F} \rightarrow \log q_s = b_F \cdot \log C_s + \log a_F \]

where \( a_F (\text{dm}^3\cdot\text{g}^{-1}) \) expresses the adsorbent capacity (the larger value the higher the capacity) and \( b_F \), which ranges from 0 to 1, is the heterogeneity factor.

RESULTS AND DISCUSSION

Chemical composition

Table 2 shows the chemical composition of all three types of opoka in its natural and heated form. According to the silica and carbonate content, opoka can be classified as light-weight opoka (more SiO₂) and heavy-weight opoka (more CaCO₃). Figure 1 shows the deposits of opoka in the quarry of Strzęzów. Opoka from Strzęzów (Opk1) could be then classified as light opoka and the other two types, Opk2 from Cisie (Antolka) and Opk3 from Widnica, with higher carbonate content, could be catalogued as heavy opoka. There is a noticeable higher content of Al and Fe oxides in light opoka (Opk1). The rest of the components appear in similar proportions for all three types of opoka.

Light opoka has a high porosity whereas heavy opoka has a more compact structure due to its higher content of CaCO₃ (Brogowski and Renman, 2004). This could favour P movement through cavities and make more sorption places available. However, heavy opoka has a considerably higher content of Ca and a larger sorption capacity is expected.

Most of the CaCO₃ and other possible Ca forms in opoka convert to CaO after heating to 900°C. This has been already reported and it is the base of a strong Ca-P interaction due to the higher reactivity of CaO (Johansson and Gustafsson, 2000; Brogowski and Renman, 2004). The pH values increase drastically after heating, which also favours the Ca-P reaction.

It has been observed a significant reduction in the Al and Fe content after heating of light opoka (Opk1) while it did not change much for Opk2 and Opk3. There is no noticeable change in the content of the other elements after heating to 900°C with the exception of a slight reduction of Cu for all three types of opoka.

P-sorption capacity

The batch experiments showed a high P-sorption capacity for powder fractions of all three materials heated to 900°C. The natural form of opoka was not tested in this study since it has already been reported a low P-sorption capacity by Johansson, 1999. Light opoka (900Opk1) removed up to 70 mg P·g⁻¹ from a solution of 500 mg P·dm⁻³ and its maximum sorption capacity was estimated in 79 mg P·g⁻¹ by the Langmuir adsorption isotherm (Tab. 3). However, the experimental data fitted better to the Freundlich isotherm as shown in Figure 2.

The heavy opoka, both 900Opk2 and 900Opk3, showed a higher sorption capacity than light opoka, removing 100 mg P·g⁻¹ and 120 mg P·g⁻¹ respectively.
TABLE 2. Chemical composition of opoka samples (Opk: natural; 900Opk: heated)

| Element | Opk1   | Opk2   | Opk3   | 900Opk1 | 900Opk2 | 900Opk3 |
|---------|--------|--------|--------|---------|---------|---------|
| g kg⁻¹  |        |        |        |         |         |         |
| Si      | 231.32 | 169.34 | 154.57 | 310.67  | 248.37  | 210.05  |
| Ca      | 171.34 | 296.55 | 311.58 | 220.79  | 364.39  | 419.75  |
| Mg      | 2.56   | 2.77   | 2.83   | 2.02    | 2.78    | 3.11    |
| Al      | 8.96   | 6.45   | 7.32   | 6.41    | 5.87    | 7.11    |
| Fe      | 10.07  | 7.30   | 6.87   | 6.72    | 6.20    | 5.76    |
| K       | 0.42   | 0.29   | 0.34   | 0.39    | 0.32    | 0.32    |
| Na      | 1.32   | 0.94   | 0.99   | 0.49    | 0.44    | 0.46    |
| P       | 0.19   | 0.22   | 0.22   | 0.13    | 0.18    | 0.21    |
| mg kg⁻¹ |        |        |        |         |         |         |
| Mn      | 41.43  | 50.15  | 42.33  | 36.87   | 75.18   | 43.41   |
| Zn      | 19.87  | 20.99  | 19.58  | 15.19   | 18.31   | 15.74   |
| Cu      | 5.31   | 5.37   | 6.16   | 2.32    | 2.56    | 3.41    |
| Co      | 1.29   | 1.05   | 1.11   | 1.43    | 1.77    | 0.91    |
| Cd      | < 0.5  | < 0.5  | < 0.5  | < 0.5   | < 0.5   | < 0.5   |
| pH (KCl)| 7.48   | 7.65   | 7.55   | 12.40   | 12.41   | 12.41   |

FIGURE 1. Quarry at Strzeżów (Miechów)
FIGURE 2. Sorption isotherms of heated opoka with artificial P solutions.
from a solution containing 500 mg P·dm$^{-3}$ (Fig. 2). The experimental data for heavy opoka fitted better to the Langmuir isotherm and its estimated maximum adsorption capacity is given in Table 3.

There has been observed a very good correlation between the total

TABLE 3. P-sorption parameters from the Langmuir and Freundlich isotherms

| Sample | $R^2$ | $k_L$ (dm$^3$·g$^{-1}$) | $k_L/a_L$ (mg·g$^{-1}$) | $R^2$ | $a_F$ (dm$^3$·g$^{-1}$) | $b_F$ |
|--------|-------|------------------------|-------------------------|-------|------------------------|-------|
| 900Opk1 | 0.9777 | 0.3909 | 79.37 | 0.9836 | 0.9768 | 0.6950 |
| 900Opk2 | 0.9973 | 0.5126 | 136.99 | 0.9833 | 0.8714 | 0.8192 |
| 900Opk3 | 0.9991 | 0.4405 | 181.82 | 0.9994 | 0.8794 | 0.8227 |

The results are comparable to those obtained by Brogowski and Renman who reported a sorption capacity of 119 mg P·g$^{-1}$ using a processed opoka from different origin (Brogowski and Renman, 2004).

The Freundlich equation, although empirical, implies that the affinity for adsorption decreases exponentially with increasing saturation, which may be closer to reality than the assumption of constant binding energy inherent in the Langmuir equation.

The Freundlich equation appear to model the equilibrium P-adsorption data better than the Langmuir equation for many other reactive media studied before such as slags, zeolites, calcareous soils and limestone (Sakadevan and Bavor, 1998; Zhou and Li, 2001). The Freundlich $a_F$ values give a measure of the relative P-adsorption capacity. However, these values are not directly related to the adsorption maxima from the Langmuir equation in the studied system. This has been previously observed by Sakadevan and Bavor, 1998, when using other reactive substrata and it was attributed to the precipitation of P in addition to P adsorption. Thus, adsorption mechanisms dominate in light opoka while precipitation may contribute considerably to the P-sorption process in heavy opoka.

Ca content in heated opoka and its estimated (Langmuir) and observed (at 500 mg P·dm$^{-3}$) maximum P adsorption capacity, with correlation coefficients of $R = 0.99$ and $R = 0.98$ respectively. This gives consistency to explain that Ca-P interaction drives the P-sorption mechanism in powdered opoka heated to 900°C and the more CaO the higher the sorption capacity.

Light opoka (Opk1) removed about 95% of P from a 10 mg P·dm$^{-3}$ solution while heavy opoka (Opk2 and Opk3) removed 100%. The removal efficiency of Opk1 dropped to nearly 50% with a 50 mg P·dm$^{-3}$ solution while it was necessary 200 mg P/l or more to drop the efficiency of heavy opoka (Opk2 and Opk3) to 50%. The pH values decreased with increasing P concentration from 11 to about 7. The P-sorption process in opoka is highly dependent on pH but this has not been the focus of this study.

These results are not applicable when using other fractions of opoka and they should be taken carefully. Courser fractions have higher hydraulic conductivity and therefore are more suitable for infiltration systems, however, they have lower porosity and its expected...
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sorption capacity will be lower than that reported in this paper. The results of a column experiment using heated opoka with courser particle size (2–5.5 mm) showed a much lower P-sorbed load (Hylander et al., 2006). Furthermore, differences between light and heavy opoka regarding porosity (Brogowski and Renman, 2004) may cause the system to work properly or fail. Heavy opoka has a lower porosity and more compacted structure what leads often to clogging the system.

It is important to remember that this research has been performed with artificial P solutions and the results may differ when using wastewater due to the presence of other substances that could interfere the P interaction.

It is not known exactly what Ca-P compounds are formed in opoka. Depending on the Ca : P ratio, pure CaCO₃ can form four main different species (House, 1998). At low P concentrations, the main form is hydroxiapatite. P sorbed may be more or less available depending on the strength of the interaction and type of compound formed. It is known that hydroxiapatite is a strong component, in which form P is not available to plants (Johansson and Gustafsson, 2000). Further research shall test the plant availability of P sorbed to different types of opoka.

CONCLUSIONS

Opoka deposits in the region of Miechów, Poland, appear to vary considerably. Depending on its chemical composition, the rock was grouped as light-weight opoka (Strzeżów), with predominant silica, and heavy-weight opoka (Cisie and Widnica), with higher carbonate content. There is a noticeable higher content of Al and Fe oxides in light opoka. The P-sorption capacity increases considerably when heating the material up to 900°C and in that form, its P-sorption capacity is directly related to the CaO content. P-sorption maximum capacities of about 70 and 100–120 mg P g⁻¹ have been observed for powdered light and heavy opoka respectively. The Langmuir isotherm estimates in 80 mg P·g⁻¹ and 137–181 mg P·g⁻¹ the P-sorption capacities of the tested powdered light and heavy opoka respectively. The P-sorption isotherms fitted well to the Freundlich adsorption model for light opoka and to the Langmuir isotherm for heavy opoka, showing that different mechanisms govern the P-sorption process in light and heavy opoka. Further research shall focus on the study of the plant available P of different types of opoka.

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REFERENCES

ANGEL R. 1999: Removal of phosphate from sewage as amorphous calcium phosphate. Environ. Technol. 20 (7): 709–720.
BOLEWSKI A., TURNAU-MORAWSKA M. 1963: Petrografia. Wydawnictwa Geologiczne. Warszawa.

BROGOWSKI Z., RENMAN G. 2004: Characterisation of Opoka as a Basis for its use in wastewater treatment. Polish J. of Environ. Studies 13 (1): 15–20.

DRIZO A., FROST C., GRACE J., SMITH K. 1999: Physico-chemical screening of phosphate-removing substrates for use in constructed wetland systems. Water Res. 33 (17): 3595–3602.

GRIFFIN R.A., JURINAK J. J. 1974: Kinetics of the phosphate interaction with calcite. Soil Sci. Soc. Amer. Proc., vol 38.

HYLANDER L., KIETLINSKA A., RENMAN G., SIMÁN G. 2006: Phosphorus retention in filter materials for wastewater treatment and its subsequent suitability for plant production. Biore. Technol., 97: 914–921.

HOUSE W.A. 1999: The physico-chemical conditions for the precipitation of phosphate with calcium. Int. conf. on phosphorus recovery, Warwick University, UK.

JOHANSSON L. 1998: Industrial by-products and natural substrata as phosphorus sorbents. Environ. Technol. 20: 309–316.

JOHANSSON L., GUSTAFSSON J.P. 2000: Phosphate removal from wastewaters using blast furnace slags and opoka – Mechanisms. Water Res. 34 (1): 259–265.

JOHANSSON WESTHOLM L. 2006: Substrates for phosphorus removal-Potential benefits for on-site wastewater treatment. Water Res. 40: 23–36.

MANN R.A., BAVOR H.J. 1993: Phosphorus removal in constructed wetlands using gravel and industrial waste substrata. Wat. Sci. Tech. 27 (1): 107–113.

MURPHY J., RILEY J.P. 1962: A modified single solution method for the determination of phosphate in natural waters. Anal. Chim. Acta 27: 31–36.

SAKADENAV K., BAVOR H. 1998: Phosphate adsorption characteristics of soils, slags and zeolites to be used as substrates in constructed wetland systems. Water Res., 22 (2): 393–399.

ZHOU M., LI Y. 2001: Phosphorus-sorption characteristics of calcareous soils and limestone from the southern Everglades and adjacent farmlands. Soil. Sci. Soc. Am. J. 65: 1404–1412.

ZHU T., JENSSEN P.D., MAEHLUM T., KROGSTAD T. 1997: Phosphorus sorption and chemical characteristics of lightweight aggregates (LWA) – potential filter media in treatment wetlands. Wat. Sci. Tech. 35 (5): 103–108.