Effects of fresh and aged biochars from pyrolysis and hydrothermal carbonization on nutrient sorption in agricultural soils

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Abstract

Leaching of nutrients from agricultural soils causes major environmental problems that may be reduced with biochar amendments to the soils. Biochars are characterised by a high adsorption capacity, i.e., they may retain nutrients such as nitrate and ammonium. However, biochar properties strongly depend on feedstock and the production process. We investigated the nutrient retention capacity of biochars derived from pyrolysis (pyrochar) as well as from hydrothermal carbonization (hydrochar; produced at 200 and 250 °C) from three different feedstocks (digestates, Miscanthus, woodchips) mixed into different soil substrates (sandy loam and silty loam). Moreover, we investigated the influence of biochar degradation on its nutrient retention capacity using a seven-month in-situ field incubation of pyrochar and hydrochar. Pyrochars showed the highest ability to retain nitrate, ammonium and phosphate, with pyrochar from woodchips being particularly efficient in nitrate adsorption. Ammonium adsorption of pyrochars was controlled by the soil type of the soil-biochar mixture. We found some ammonium retention on sandy soils, but no pyrochar effect or even ammonium leaching from the loamy soil. The phosphate retention capacity of pyrochars strongly depended on the pyrochar feedstock with large phosphate leaching from digestate-derived pyrochar and some adsorption capacity from woodchip-derived pyrochar. Application of hydrochars to agricultural soils caused small, and often not significant, effects on nutrient retention. In contrast, some hydrochars did increase the leaching of nutrients compared to the non-amended control soil. We found a surprisingly rapid loss of the biochars’ adsorption capacity after field application of the biochars. For all sites and for hydrochar and pyrochar, the adsorption capacity was reduced by 60–80 % to less or no nitrate and ammonium adsorption. Thus, our results cast doubt on the efficiency of biochar applications to temperate zone soils to minimize nutrient losses via leaching.
1 Introduction

Excessive application of mineral fertilizers to agricultural soils is one of the major drivers for various threats to the environment (Laird et al., 2010; Liang et al., 2006). An excess of nutrients may induce soil acidification, increase direct and indirect greenhouse gas emissions (Karaca et al., 2004) and cause eutrophication of the receiving water bodies. However, mineral fertilization has also been the major driver for increased global agricultural production during the last decades. Therefore, technologies are required to both decrease nutrient leaching from soils and enhance nutrient use efficiency with the result that less fertilizer is needed. Biochar amendment to soils is proposed as one promising option to retain nutrients and prevent leaching (Lehmann, 2009).

Biochar is the solid charcoal product derived from the thermal transformation of a variety of organic feedstocks such as digestates, sewage sludge, woods and other forestry or agricultural residues (Hale et al., 2013; Yao et al., 2012). At present, two main production processes for biochar are used: the first production process, slow pyrolysis, is the combustion and conversion of biomass at processing temperatures above 450 °C under oxygen-free conditions. In the following, we will refer to the solid product derived from pyrolysis as pyrochar. Pyrochars are characterized by a high degree of aromaticity (Keiluweit et al., 2010; Lehmann et al., 2006) and recalcitrance against degradation or mineralization (Glaser et al., 2002). Second, hydrothermal carbonization (HTC) is a low-temperature production process (temperatures between 180 and 300 °C) under high pressure (2–2.5 MPa) with water for several hours (Funke and Ziegler, 2010; Libra et al., 2011; Wiedner et al., 2013). In the following, we will refer to the solid product from the HTC as hydrochar. Hydrochars have recently received increasing attention since wet feedstock can also be carbonized without drying pre-treatment (Funke and Ziegler, 2010). Hydrochars are characterized by a lower degree of carbonization and thus more aliphatic carbon (C) but smaller amounts of aromatic C and lower specific surface area (SSA) compared to pyrochars (Eibisch et al., 2013; Titirici et al., 2008). Besides general differences between pyrochar and hydrochar,
their properties differ strongly depending on the feedstock, carbonization processes parameters, and subsequent thermochemical reactions (Cantrell et al., 2012; Cao et al., 2011; Eibisch et al., 2013, 2015; Yao et al., 2012).

For the past ten years, the application of biochar derived from pyrolysis, and later from HTC to agricultural soils, has become a centre of attention as an option to store atmospheric C in soil to mitigate global warming. Additionally, a variety of positive co-benefits are attributed to pyrochar amended soils: an increase in water retention capacity (Glaser et al., 2002; Abel et al., 2013), reduction of greenhouse gas emissions such as nitrous oxide (\(\text{N}_2\text{O}\)) and methane (\(\text{CH}_4\)), and an enhanced crop productivity due to the retention of plant available nutrients in the rhizosphere (Lehmann, 2009), increased soil pH and soil cation exchange capacity (CEC) (Liang et al., 2006), and preservation of toxic compounds (Chen and Yuan, 2011).

Both pyrochars and hydrochars contain nutrients which can be released slowly into the rhizosphere (Eibisch et al., 2013; Spokas et al., 2011; Taghizadeh-Toosi et al., 2011) but more important is the pyrochars’ ability to adsorb nutrients due to its high surface charge density and CEC. The leaching and adsorption of nitrate (\(\text{NO}_3^-\)), ammonium (\(\text{NH}_4^+\)), and phosphate (\(\text{PO}_4^{3-}\)) to various activated C and charcoals has been studied (Bandosz and Petit, 2009; Ding et al., 2010). However, studies concerning the sorption behavior of biochar, and especially hydrochars, are rare. Previous studies focusing on soil-biochar mixtures have shown that leaching of \(\text{NO}_3^-\), \(\text{NH}_4^+\), and \(\text{PO}_4^{3-}\) from soils amended with biochar from pyrolysis and HTC was frequently reduced due to adsorption on the respective biochar (Bargmann et al., 2014b; Ding et al., 2010; Laird et al., 2010; Sarkhot et al., 2012). Laird et al. (2010) applied 20 g kg\(^{-1}\) pyrochar from hardwood to an agricultural soil, which decreased the leaching of \(\text{NO}_3^-\) from swine manure by 10%. Yao et al. (2012) reported increased \(\text{NO}_3^-\) adsorption of up to 4%, but also leaching rates of up to 8% from aqueous solution. Other studies showed that \(\text{NO}_3^-\) (Castaldi et al., 2011; Hale et al., 2013; Jones et al., 2012), as well as \(\text{NH}_4^+\) leaching was decreased by 94% due to pyrochar application to a ferralsol in a 37 day soil column leaching experiment (Lehmann et al., 2003). Furthermore, both \(\text{NH}_4^+\) adsorption...
by up to 15% from aqueous solution, but also leaching up to 4% in to solution was observed (Yao et al., 2012). Also other nutrients which are not particularly prone to leaching, such as \( \text{PO}_4^{3-} \), have been reported to be retained by application of pyrochar (Laird et al., 2010; Morales et al., 2013; Xu et al., 2014). For example, Laird et al. (2010) reported up to 70% reduced \( \text{PO}_4^{3-} \)-P leaching in a soil column experiment mixed with 20 g kg\(^{-1}\) pyrochar. In contrast, Yao et al. (2012) observed up to 5% \( \text{PO}_4^{3-} \)-P leaching from aqueous solution for pyrochars from bamboo and hydrochars from peanut-hull. In summary, these studies implicate a strong variation of leaching or retention behavior of biochars, which seems to depend on feedstock and production process.

Biochar application has been promised to be multi-beneficial. However, benefits have been tested mostly for biochar from slow pyrolysis-amended tropical soils with few comparative studies for temperate soils or hydrochars. This is one of the main reasons why neither pyrochar nor hydrochar application is considered in agricultural practice in the temperate zone at the moment. Even though biochars, especially pyrochars, are relatively stable in soils, an increasing number of studies suggest that biotic and abiotic processes can lead to degradation of biochar and thus change its surface properties and sorption behavior (Cheng et al., 2008; Hale et al., 2011; Liu et al., 2013; Steinbeiss et al., 2009). The physical structure and chemical properties of hydrochars result in a lower recalcitrance towards microbial degradation compared to pyrochars (Bargmann et al., 2014a; Hale et al., 2011; Steinbeiss et al., 2009). Furthermore, hydrochars release a higher amount of DOC which might be easily mineralized. Hence, soil amended with hydrochars increases microbial-biomass production and immobilization of mineral nitrogen (Bargmann et al., 2014a; Lehmann et al., 2011), and an increased nitrification from \( \text{NH}_4^+ \) to \( \text{NO}_3^- \) may occur. Over time, slow biochar aging due to oxidation may lead to carboxylic and phenolic functional groups on the chars’ surface and thus negative charges. On the other hand, the atomic C content and positive surface charge on the edge sites of aromatic compounds will be reduced (Cheng et al., 2008; Cheng et al., 2006; Glaser et al., 2000). Furthermore, surface oxidation increases CEC per unit C and the charge density (Liang et al., 2006), but a higher anion exchange capacity
In summary, according to the majority of studies (Hale et al., 2013; Knowles et al., 2011; Lehmann et al., 2003; Morales et al., 2013; Xu et al., 2014), biochar may be a potential melioration for soils by decreasing nutrient leaching via improved adsorption properties. However, there is only little knowledge of the nutrient sorption potential of pyrochars compared to hydrochars, and the influence of ageing/degradation on nutrient sorption.

The influence of biochar properties resulting from different carbonization methods and different feedstock materials on nutrient sorption potential is also insufficiently understood. Furthermore, no systematic comparison of different feedstock materials on nutrient sorption has yet been conducted, and the effect of aging of biochars on their sorption potential has not yet been investigated. Therefore, the objectives of this study are to first determine the nutrient sorption potential of nine different biochar-soil mixtures in laboratory batch experiments and to investigate the influence of (i) biochar type (pyrochar vs. hydrochar), (ii) soil type (sandy loam vs. silty loam), and (iii) biochar feedstock (woodchips, digestate and Miscanthus). Secondly, we want to assess the effect of aged vs. fresh biochars (pyrochar and hydrochar from Miscanthus) on nutrient sorption potential in a field experiment.
2 Materials and methods

2.1 Production and general properties of pyrochars and hydrochars and their corresponding feedstocks

The nine biochars that were used for laboratory batch experiments originated from the same setup as the chars described in Eibisch et al. (2013, 2015). They derived from HTC and pyrolysis and were produced from three feedstock materials with different physico-chemical properties (digestates (99% maize), woodchips (95% poplar, 5% willow), and Miscanthus). The hydrochars were carbonized with water (1 : 10, w/w) in a batch reactor for 6 h, 2 MPa at 200 (hereafter referred to as Hydro200) and 250°C (hereafter referred to as Hydro250; SmartCarbon AG, Jettingen, Germany). Pyrochars were produced in a Pyreg reactor (PYREG GmbH, Dörth) for 0.75 h at 750°C (designated hereafter as Pyro750). Detailed information on biochar preparation and methods of analysis (e.g., specific surface area (SSA), pore volume, average pore size) can be found in Eibisch et al. (2013, 2015).

Hydrochar and pyrochar produced from Miscanthus was used for the field incubation. The hydrochar was carbonized with water (1 : 10, w/w) in a tabular reactor (3 m$^{-3}$) for 11 h, 2 MPa at 200°C by AddLogicLabs/SmartCarbon (Jettingen, Germany). Citric acid was added as catalyst for the dehydration process and to increase the C content in the solid product (Wang et al., 2010). Pyrochars were produced in a Pyreg reactor for 0.75 h at 750°C. Analyzes of general properties of the chars and raw material were carried out by Andrea Kruse (KIT Karlsruhe). All chars were dried at 40°C and sieved ≤ 2 mm. Basic characteristics of feedstocks, pyrochars, and hydrochars for the laboratory batch and field incubation experiment are listed in Table 1.

In order to simulate field ageing, we compared unwashed biochars with washed biochars in the laboratory experiment. Washing was assumed to be capable of simulating ageing of the char as initially bound nutrients or salts would be removed. Washing was carried out by shaking 4.5 g biochar with 1 L deionized water in an overhead-shaker at 9 rpm for 4 h and thereafter solution was filtered with pleated paper filter (Grade:
3 hw; Diameter: 150 mm; 65 g m\(^{-2}\)) and filtrate (pyrochar or hydrochar) was dried for 24 h at 105°C. Washing effects were only studied in the pyrochar and hydrochar applied to silty loam mixtures, because highest nutrient leaching or adsorption effects were expected for this soil.

### 2.2 Field ageing

For the investigation of the effect of ageing of the chars in the field, biochars were incubated in-situ at three cropland sites in the North German lowland (mean annual temperature 8.8°C, around 600 mm precipitation). The three sites differ mainly in their soil texture (Table 2) and are located in Bortfeld (sandy loam (SL); 52°28′N, 10°41′E, 80 m a.s.l.), Volkmarsdorf (sandy loam (SL); 52°36′N, 10°89′E, 105 m a.s.l.) and Querenhorst (loamy sand (LS); 52°33′N, 10°96′E, 112 m a.s.l.). All sites were managed according to common regional practice with conventional tillage and fertilizing. Crop rotations were barley (2012), winter wheat (cover crop), sugar beet (2013) (Querenhorst); barley (2012), mustard (cover crop), sugar beet (2013) (Volkmarsdorf); potatoes (2012), sugar beet (2013) (Bortfeld). At all three sites, mini-plots (plot size: 70 × 70 cm; plot depth: 25 cm) were dug out in triplicate in March 2013, and the hydrochar and pyrochar were mixed into the soil in a cement mixer in an amount that aimed to double the soils’ C-content (corresponds to around 100 t ha\(^{-1}\) biochar). The experimental setup was a randomized plot design carried out in three rows for each site so that every row consisted of three treatments: (i) control (soil only), (ii) soil + hydrochar, and (iii) soil + pyrochar. In order to distinguish the soils’ C-contents from treated or non-treated soil, and to quantify any blending or attenuation with the surrounding soil, e.g., due to tillage, 105 g Zinc as an inert tracer was added to each treatment in the cement mixer (control, pyrochar + soil, hydrochar + soil). The mini-plots were not fenced off, so the farmers were able to manage the fields exactly like to the rest of the field.
Sampling was carried out twice: the first set of soil samples was taken in March 2013 right after mixing the soil with biochars \((T_0)\). After seven months (October 2013) a second sampling was carried out \((T_1)\). Soil samples were obtained by taking five randomly distributed soil cores to a depth of 25 cm with a Split-Tube sampler (5 cm diameter) from each mini-plot. Afterwards, samples were dried at 40°C and sieved \(\leq 2\) mm. Zinc concentrations at \(T_0\) and \(T_1\) were used to calculate a correction factor \(F_Z\), which determines the recovery-rate of incubated biochars in the field study

### 2.3 Batch sorption experiments

Soil-biochar mixtures used solely in the laboratory were produced by mixing 0.5 g of biochar with 10 g soil in order to roughly double the soil’s C content. Two soils were used for the char-soil mixtures: a silt loam (Blagodatskaya et al., 2014) from a cropland site at the Thünen-Institute in Braunschweig, Germany \((52°17'\ N, 10°26'\ E,\ 80\ m\ a.s.l.)\) and a sandy loam from a cropland site of the University of Göttingen (Reinshof), Germany \((51°28'\ N, 9°58'\ E,\ 205\ m\ a.s.l.)\). The soil was dried at 105°C to inhibit any microbial activity and sieved \(\leq 2\) mm. The pH-value of soils and biochars was measured in 0.01 M \(\text{CaCl}_2\) with a ratio of 1 : 5 (volume soil / volume solution). Carbon and N contents were determined using dry combustion with an elemental analyzer (LECO TrueMac CN LECO Corp., St. Joseph (MI), USA). Soil texture was determined by the combined sieve and pipette method.

Preliminary sorption kinetic experiments were conducted to determine the sorption equilibrium by shaking the batches for 4, 8, 12, 24, and 48 h at 9 rpm in an overhead shaker. Based on the results of the kinetic experiments, shaking time for the determination of the sorption isotherms was set to 24 h. An amount of 10.5 g of soil only (control) and soil-biochar mixtures were added to 40 mL of a nutrient solution in a 50 mL plastic centrifuge tube. Six concentration levels of a nutrient solution containing several nutrients that were chosen in order to mimic a “typical” agricultural soil solution were used (Table 3). In addition, the pH-value of the solution was adjusted to 6 by adding HCl. Triplicates were measured for each concentration level. Cation exchange...
capacity (CEC) and pH were measured immediately after shaking in the biochar / soil-solution mixtures. Thereafter, suspensions were centrifuged at 4500 rpm for 30 min. The supernatant was aspirated with a syringe and filtered through 0.45 µm membrane filters (CHROMAFIL PET-45/25 disposable syringe filters, Macherey-Nagel). The ion-concentrations of the filtrates were analyzed using ion chromatography (METROHM 761) for anions (NO$_3^-$, PO$_4^{3-}$) and inductively coupled plasma chromatography (ICS-90 Dionex/Thermo Fisher Scientific) for cations (NH$_4^+$). Moreover, contents of Ca$^{2+}$, Mg$^{2+}$, K$^+$, and SO$_4^{2-}$ were also determined and fitted isotherms can be found in Table S1 in the Supplement.

Soil-biochar mixtures from the field experiment were used directly in the batch sorption experiments (NO$_3^-$, NH$_4^+$, PO$_4^{3-}$), which were carried out as described above. To calculate the biochar adsorption effect relative to the control we used the following equations:

Relative adsorption of the control:

$$Q_{Ctrl} = \left( 1 - \left( \frac{IC (Ctrl)}{IC (Blind)} \right) \right) \times 100. \quad (1)$$

Relative adsorption of the biochar treatment to control:

$$Q_{BChar} = \left( 1 - \left( \frac{IC (BChar)}{IC (Ctrl)} \right) \right) \times F_Z \times 100. \quad (2)$$

Whereby $F_Z$ was only used to calculate relative adsorption for field incubated biochars. IC is the equilibrium ion content of the nutrient solution after shaking for blinds (IC$_{Blind}$), control (IC$_{Ctrl}$) or soil biochar mixtures (IC$_{BChar}$).
2.4 Statistical analyses

Adsorption data were fit to Freundlich and linear adsorption isotherms:

Freundlich isotherm: \( Q_e = K_F \times IC^{1/n} \)  \( (3) \)
Linear isotherm: \( Q_e = a \times IC + Y_0. \)  \( (4) \)

\( Q_e \) is the amount of ion adsorbed, while IC is the ion concentration in the solution after 24 h equilibration. A positive \( Q_e \) indicates adsorption of ions in the nutrient solution on an adsorbent and a negative \( Q_e \) desorption from adsorbent to the nutrient solution.

Logarithmized equilibrium-concentration and log adsorbed amount was used to calculate the Freundlich sorption partitioning coefficients (\( K_F \)) and the Freundlich exponents (\( 1/n \)) following nonlinear fitting. For linear isotherm, \( Y_0 \) is the intercept.

The Akaike information criterion (AIC) was used to select the best fitting isothermal model. Significance of treatment effects on shape of isotherms was tested using two procedures:

1. If, for two treatments, the same model type resulted in the best fit, their difference was tested with a likelihood-ratio test. It was tested whether fitting the model to the data separately resulted in a better fit than fitting the model to the combined data. If the separately fitted model resulted in a better fit than the combined model, treatments were different with their corresponding \( p \) value. This test could only be conducted if it was numerically possible to fit the model to the combined data.

2. Generalized additive models (GAM, R package \( \text{gam} \), (Hastie, 2013)), including and excluding treatment as a predictor, were fitted and compared using analysis of deviance with a \( \chi^2 \) statistics.

All \( p \) values were adjusted for multiple testing using the procedure of Benjamini and Hochberg (1995). All statistical analyses were conducted using R 3.1.1 (RCoreTeam, 2014). The results of the statistical analyses can be found in the Supplement (Tables S1, S3, S5, S7 and S8).
3 Results

3.1 Biochars physico-chemical properties

The pH values for the used hydrochars were acidic ranging from 3.8 to 6.2, and 4.2 to 5.7, for Hydro200 and Hydro250, respectively (Table 1). The pH-values of Pyro750 were alkaline, ranging from pH 8.7 to 9.8. The ash content increased with increasing carbonization temperature and was highest for pyrochars from wood chips (24.6 %). Generally, the wood chips had the highest C concentration (48.6 % C) as a raw material, but after carbonization, Pyro750 from Miscanthus had the highest C concentrations (Lab: 76.9 % C; Field: 81.8 % C). The highest amounts of total N and P were found in Hydro200 and Hydro250 from digestates. After carbonization, highest SSA was observed for pyrochars and decreased in the order Pyro750 > Hydro200 > Hydro250 (Table 1). Pyro750 showed the highest pore volume, followed by Hydro200 and Hydro250. In general, Pyro750 showed smaller average pore size than Hydro200 and 250 by factor 10.

3.2 Influence of soil, feedstock and carbonization type on nutrient sorption (laboratory experiments)

Figures 1, 2, and 3 show the relative change of ion concentration of the biochar treatments from the three feedstocks (triangles = Miscanthus, circles = digestates, squares = wood chips) to the control (0 % line) at all applied nutrient concentration levels. Positive values correspond to adsorption and negative values to leaching.

3.2.1 Sorption of nitrate

The pure sandy loam (control in Table 3) showed neither NO$_3^-$ sorption nor release (all data points are around 0 %). In contrast, the pure silty loam tended to a high NO$_3^-$ release of around 60 % (N concentration in batch solution ($IC_{Blind}$): 5.19 mg N L$^{-1}$; N
concentration in batch solution with control soil (IC\textsubscript{Ctrl}): 8.22 mg N L\(^{-1}\)) at the lowest concentration level of the nutrient solution. This release decreased to 5 % with increasing concentrations of the nutrient solution.

Mixing soil with Pyro750 significantly reduced NO\textsubscript{3}\textsuperscript{-} leaching, independent of the soil and feedstock used (Fig. 1a, b). At the lowest concentration level of the nutrient solution (5 mg N L\(^{-1}\)), application of Pyro750 raised NO\textsubscript{3}\textsuperscript{-} adsorption between 2–15 % (silty loam) and 7–30 % (sandy loam) compared to the respective control soil (Fig. 1a, b). The relative adsorption on Pyro750 decreased with increasing nutrient solution concentration to 5–12 %. For both soil types, the fitted isotherms for Pyro750 were significantly different from the control (\(p \leq 0.01\)) and to both Hydro200 and Hydro250 (\(p \leq 0.01\)). Further, isotherms of NO\textsubscript{3}\textsuperscript{-} adsorption by Pyro750 mixed with sandy loam were significantly different to those of silt loam (\(p \leq 0.01\)). The effects of nutrient retention in Pyro750 mixtures depended on the carbonized feedstock (\(p \leq 0.01\); Fig. 1a, b). For Pyro750, adsorption increased in the order digestates < Miscanthus < woodchips in both soil types. Values for digestates ranged from 8 % (N concentration in batch solution with control soil (IC\textsubscript{Ctrl}): 5.23 mg N L\(^{-1}\); N concentration in batch solution with soil biochar mixtures (IC\textsubscript{BChar}): 5.08 mg N L\(^{-1}\)) and decreased to 3 % with increasing NO\textsubscript{3}\textsuperscript{-}-N concentration level (sandy loam) or remains at the same 3–5 % level (silty loam). For Pyro750 from Miscanthus, relative NO\textsubscript{3}\textsuperscript{-} adsorption was higher with 14 % (IC\textsubscript{Ctrl}: 5.23 mg N L\(^{-1}\); IC\textsubscript{BChar}: 4.78 mg N L\(^{-1}\)) for low NO\textsubscript{3}\textsuperscript{-}-N concentrations and 10 % at high NO\textsubscript{3}\textsuperscript{-}-N concentrations (IC\textsubscript{Ctrl}: 60 mg N L\(^{-1}\); IC\textsubscript{BChar}: 55.13 mg N L\(^{-1}\)). For Pyro750 from woodchips, the relative adsorption was highest and ranged from 15 % (IC\textsubscript{Ctrl}: 5.23 mg N L\(^{-1}\); IC\textsubscript{BChar}: 4.10 mg N L\(^{-1}\)) and decreased to 10 % with increasing NO\textsubscript{3}\textsuperscript{-}-N concentration level.

After addition of hydrochars (both, Hydro200 and Hydro250), significant effects on NO\textsubscript{3}\textsuperscript{-} retention were observed neither in the sandy loam nor in the silty loam (Fig. 1c, d). Fitted isotherms showed no differences between Hydro200 and Hydro250 and the control soil but significant differences between both control soils (\(p \leq 0.01\)). Hydrochars
from the three carbonized feedstocks showed no significant differences in their relative NO$_3^-$ adsorption (Fig. 1c, d) or fitted isotherms.

In summary, the relative amount of adsorbed NO$_3^-$ in pyrochar amended soils was higher in sandy loam than in silty loam and adsorption increased in the order digestates $<\text{Miscanthus}<\text{woodchips}$ in both soil types. Addition of hydrochar had no effect on NO$_3^-$ adsorption irrespective of the used carbonization temperature, feedstock or soil type.

### 3.2.2 Sorption of ammonium

The NH$_4^+$ sorption in the control soil was around 3–4 times higher in the silty loam than in the sandy loam (Table 3). Values were around 55% (IC$_{\text{Blind}}$: 4.68 mg N L$^{-1}$; IC$_{\text{Ctrl}}$: 2.16 mg N L$^{-1}$) at the first concentration level, and decreased to 32% with increasing nutrient concentrations, while the sandy loam adsorbed around 15% at all concentration levels.

Addition of pyrochars to the sandy loam increased the adsorption relative to control between 0% (Pyro750 from Miscanthus) and 17% (Pyro750 from digestates) (IC$_{\text{Ctrl}}$: 3.85 mg N L$^{-1}$; IC$_{\text{BChar}}$: 3.15 mg N L$^{-1}$) at the first concentration level ($p \leq 0.01$; Fig. 2a). This effect decreased with increasing ion concentration level. The fitted isotherms for Pyro750 mostly showed significant differences to control soil ($p \leq 0.01$). For the silty loam, addition of pyrochars did not raise relative NH$_4^+$ adsorption but led to leaching compared to the control. Comparison of fitted isotherms of both soils mixed with Pyro750 showed significant differences between sandy loam and silty loam ($p \leq 0.01$). The effect of feedstock on relative NH$_4^+$ adsorption was soil-dependent and significant for both soils (Fig. 2a, b; $p \leq 0.05$). Pyro750 from digestates caused the strongest increase of relative NH$_4^+$ adsorption when mixed with sandy loam (17–7% from lowest to highest nutrient solution concentration level). Pyro750 from woodchips raised NH$_4^+$ adsorption by only 8–2%. Pyro750 from Miscanthus showed no effect. When Pyro750 from digestates was added to silty loam, the highest reduction of relative NH$_4^+$ adsorp-
tion through leaching was observed for the first two NH\textsubscript{4} concentration levels (level P1: −45 % (IC\textsubscript{Ctrl}: 2.16 mg N L\textsuperscript{-1}; IC\textsubscript{BChar}: 3.15 mg N L\textsuperscript{-1})) (Fig. 2b). For silty loam amended with Pyro750 from Miscanthus, relative NH\textsubscript{4}\textsuperscript{+} adsorption was reduced by 20 % (IC\textsubscript{Ctrl}: 2.16 mg N L\textsuperscript{-1}; IC\textsubscript{BChar}: 2.62 mg N L\textsuperscript{-1}) at the first concentration level and decreased to 0 % at highest level.

Application of hydrochars to either soil type had only marginal effects. These ranged from leaching to adsorption with relative values between +10 and −20 %, respectively (Fig. 2c, d). In general, NH\textsubscript{4}\textsuperscript{+} adsorption by the control soil was significantly different to that in the soil amended with hydrochars (\(p \leq 0.01\)) for both sandy loam and silty loam. For Hydro200, NH\textsubscript{4}\textsuperscript{+} adsorption was close to zero when compared to the control at all concentration levels. A significant relative adsorption effect was observed for only some concentration points (Fig. 2c). Hydro250 showed both NH\textsubscript{4}\textsuperscript{+} release at the lowest concentration level and little adsorption of NH\textsubscript{4}\textsuperscript{+} at the higher concentration levels reaching up to about 10 % (Fig. 2d). The fitted isotherms for Pyro750 are significantly different from those for hydrochars and pure soil (depending on soil type), but there are no differences between Hydro200 and Hydro250. For hydrochars, no effect of feedstock on NH\textsubscript{4}\textsuperscript{+} adsorption was observed except for lower adsorption of Hydro200 from digestates compared to Miscanthus and woodchips (\(p \leq 0.01\)).

In summary, only pyrochars enhanced NH\textsubscript{4}\textsuperscript{+} adsorption (and only as a mixture with sandy loam), but hydrochars had either no effect or led to NH\textsubscript{4}\textsuperscript{+} release. The effect of pyrochar feedstock on NH\textsubscript{4}\textsuperscript{+} adsorption was soil-dependent. Relative NH\textsubscript{4}\textsuperscript{+} adsorption increased in the following order for pyrochars in the sandy loam: Miscanthus < woodchips < digestates. For the silty loam, the order for reduction of relative NH\textsubscript{4}\textsuperscript{+} adsorption was: woodchips < Miscanthus < digestates only at the first three nutrient concentration levels.
3.2.3 Sorption of phosphorous

The sandy loam leached $\text{PO}_4^{3-}$ at the lowest concentration level ($\text{IC}_{\text{Blind}}: 1.25 \text{ mg P L}^{-1}$; $\text{IC}_{\text{Ctrl}}: 2.29 \text{ mg P L}^{-1}$), but this changed to 65% adsorption at higher levels ($\text{IC}_{\text{Blind}}: 14.07 \text{ mg P L}^{-1}$; $\text{IC}_{\text{Ctrl}}: 4.88 \text{ mg P L}^{-1}$), while the silt loam adsorbed $\text{PO}_4^{3-}$ at all concentration levels (up to 80%; Fig. 3a, b).

Addition of pyrochars had significant effects on $\text{PO}_4^{3-}$ adsorption (the fitted isotherms for pyrochars are significantly different to the respective control soil used ($p \leq 0.01$)), but this strongly depended on feedstock material. Comparison of fitted isotherms showed significant differences between carbonized feedstocks for Pyro750 ($p \leq 0.01$). Pyro750 from Miscanthus mixed with the sandy loam resulted in a relative $\text{PO}_4^{3-}$ adsorption of 20–30% (Fig. 3a) ($\text{IC}_{\text{Ctrl}}: 2.29 \text{ mg P L}^{-1}$; $\text{IC}_{\text{Bchar}}: 1.82 \text{ mg P L}^{-1}$) but 20% less adsorption (leaching) when mixed with the silty loam (Fig. 3b). Woodchip pyrochar was most effective in adsorbing $\text{PO}_4^{3-}$ (15–40% for the silty loam and 60–70% for the sandy loam). However, pyrochar from digestates showed strong leaching in both sandy and silty loam (Fig. 3a, b). Addition of Pyro750 from digestates enriched the nutrient solution by up to 1000% (sandy loam; $\text{IC}_{\text{Ctrl}}: 1.25 \text{ mg P L}^{-1}$; $\text{IC}_{\text{Bchar}}: 24.87 \text{ mg P L}^{-1}$) and 1300% (silty loam; $\text{IC}_{\text{Ctrl}}: 1.25 \text{ mg P L}^{-1}$; $\text{IC}_{\text{Bchar}}: 14.47 \text{ mg P L}^{-1}$) at the lowest $\text{PO}_4^{3-}$-P concentration level, and decreased to 100% at the highest $\text{PO}_4^{3-}$-P concentration. Overall, although relative $\text{PO}_4^{3-}$ adsorption was higher in the sandy loam than in the silty loam after addition of Pyro750, these differences were not significant.

The addition of hydrochar (both Hydro200 and Hydro250) to soil led mainly to leaching of $\text{PO}_4^{3-}$ from biochars (Fig. 3c, d). Fitted isotherms showed significant differences between Hydro200 and Pyro750 ($p \leq 0.01$) but no differences to control or Hydro250. The adsorption of the soil was lowered by maximum values of around 40% for the sandy loam and 60% for the silty loam due to $\text{PO}_4^{3-}$ leaching. Values depended on the feedstock used and $\text{PO}_4^{3-}$ adsorption was significantly different in both soil types ($p \leq 0.01$). Again, the effect of feedstock (or any effect at all) was less pronounced for
hydrochars than pyrochars: Hydrochars from digestates tended to reduce the relative \( \text{PO}_4^{3-} \) adsorption by leaching. Mixing soil with Hydro200 and Hydro250 from Miscanthus and woodchips resulted in no effect on \( \text{PO}_4^{3-} \) adsorption (Fig. 3c). For both soil types, differences between Hydro200 from digestates to Miscanthus and to woodchips were significant \((p \leq 0.01)\). For Hydro250 only digestates to Miscanthus and to woodchips were significantly different \((p \leq 0.01)\) in the sandy loam.

In summary, only pyrochars enhanced \( \text{PO}_4^{3-} \) adsorption (especially as a mixture with the sandy loam), but hydrochars had either no effect or led to \( \text{PO}_4^{3-} \) release. The effect of pyrochar feedstock on \( \text{PO}_4^{3-} \) adsorption was soil-dependent. Relative \( \text{PO}_4^{3-} \) adsorption increased in the following series for pyrochars in the sandy loam: digestates (only leaching) < Miscanthus < woodchips. For the silty loam, series for relative \( \text{PO}_4^{3-} \) sorption was: digestates (only leaching) < Miscanthus (little leaching) < woodchips.

### 3.3 The effect of biochar ageing on nutrient sorption (field experiment)

At all three experimental sites \( \text{NO}_3^- \) was leached from pure soil with no biochar addition (control; data not shown). However, leaching was less pronounced at \( T_1 \) than \( T_0 \) \((p < 0.01)\). Amending the soils with biochar led to adsorption of \( \text{NO}_3^- \) for both pyrochar and hydrochar at all experimental sites (Fig. 4a–c). However, adsorption was higher for pyrochars than hydrochars \((p \leq 0.01,)\). Pyrochar reduced \( \text{NO}_3^- \) leaching up to 58 % relative to the control soil at the lowest nutrient solution concentration while hydrochar reduced leaching up to 25 % (Fig. 4a–c). After 7 months of ageing in the field \((T_1)\), adsorption by pyrochars decreased by 60 to 80 % often ending up with no nutrient retention relative to control \((p < 0.01; \text{Fig. 4a–c})\). Slight differences were observed between the three investigated sites but they were not significant. The effect of hydrochar addition diminished in a similar way after seven months: relative adsorption decreased by 10 to 100 %, ending up with no nutrient retention at Bortfeld (Fig. 4a) or even nutrient leaching (site Querenhorst and site Volkmarsdorf, Fig. 4b–c), as compared to the non-amended control soil. In four of our six cases, sorption effects of both pyrochar
and hydrochar were found to be significantly different for the aged biochar-soil mixture as compared to fresh biochars mixed into soils.

Highest adsorption of $\text{NH}_4^+$ was observed for fresh biochars ($T_0$) and adsorption was higher for pyrochar than for hydrochar at two sites (Bortfeld and Volkmarsdorf, $p \leq 0.01$), but was similar at the third site (Querenhorst) (Fig. 4d–f). For soils amended with fresh pyrochar, adsorption of $\text{NH}_4^+$ was up to 40% higher than observed for the control soil. After seven months, $\text{NH}_4^+$ adsorption of pyrochar-soil mixtures was significantly lower at all experimental sites than right after the biochar application ($p < 0.01$). Little relative $\text{NH}_4^+$ adsorption was found for fresh hydrochar and for aged hydrochar in the field. The relatively low adsorption capacity of hydrochars sometimes even changes to $\text{NH}_4^+$ leaching. The effect of pyrochar ageing on $\text{PO}_4^{3-}$ adoption was different from the other nutrients: Ageing increased the $\text{PO}_4^{3-}$ retention capacity of pyrochar soil mixtures at all three sites (Fig. 4g–i). The effect of hydrochar on $\text{PO}_4^{3-}$ was minor. Hydrochar was a source for $\text{PO}_4^{3-}$ in most soils with no consistent changes due to biochar ageing.

### 3.4 Effects of biochar preparation (washing)

Washing was carried out in order to reduce initial leaching effects from biochars, i.e., it was assumed that nutrients and salts were removed from the surface of the chars by washing. Figure 5 shows relative changes of ion concentration to control (0% line; $\text{IC}_{\text{Blind}}$: 20.23 mg N L$^{-1}$; $\text{IC}_{\text{Ctrl}}$: 23.37 mg N L$^{-1}$) at nutrient concentration level P3 (Table 3). Positive values indicate higher, and negative values indicate lower, removal of ions from nutrient solution compared to control due to adsorption or leaching, respectively. Washing of both Hydro200 and Hydro250, increased pH of the nutrient solution by 0.1 to 0.2 pH-units and for Pyro750, pH was decreased by 0.2 to 0.4 pH-units due to washing. The sorption behavior of both pyrochars and hydrochars significantly changed due to washing (Fig. 5). Washing increased the potential $\text{NO}_3^-$ adsorption of pyrochars by 3–4% ($p \leq 0.05$; Fig. 5a). For hydrochars, a similar effect was only observed for Hydro200 from digestates, turning the soil-hydrochar mixture from a $\text{NO}_3^-$
source (leaching) into a sink (absorption) \((p \leq 0.05)\). In the case of \(\text{NH}_4^+\), a decrease in net leaching was observed for all treatments (Fig. 5b). For most hydrochars, washing even turned soil-hydrochar mixtures from \(\text{NH}_4^+\) sources (leaching) into net sinks (adsorption (Fig. 5b). Strongest reductions in leaching were observed for Pyro750 (−37\%) and Hydro200 from digestates (−35\%). Washing effects on \(\text{PO}_4^{3-}\) sorption were inconsistent. Pyro750 showed increased \(\text{PO}_4^{3-}\) leaching (digestates), decreased adsorption (wood chips) and leaching instead of sorption (Miscanthus) (Fig. 5c). In the case of Hydro200 from digestates, \(\text{PO}_4^{3-}\) leaching was reduced by up to −950\%. For all other hydrochar mixtures, washing reduced both \(\text{PO}_4^{3-}\) leaching and sorption close to zero. Overall, washing seemed to be an effective measure to reduce the ion leaching of those ions that were adsorbed to the surface of fresh biochars.

4 Discussion

4.1 Char-induced effects on nutrient sorption: effects of carbonization process and feedstock material

Pyrochars and hydrochars showed general differences in their sorption behavior. In most cases, pyrochars could remove \(\text{NO}_3^-\), \(\text{NH}_4^+\), and \(\text{PO}_4^{3-}\) from soil solution. This is in line with previous studies (Hale et al., 2013; Sarkhot et al., 2013; Yao et al., 2012). Hydrochars showed marginal or no sorptive effect on \(\text{NO}_3^-\), \(\text{NH}_4^+\), and \(\text{PO}_4^{3-}\). Similar to our findings, Yao et al. (2012) found no sorptive effect of hydrochar from peanut hulls on \(\text{NO}_3^-\), \(\text{NH}_4^+\), and \(\text{PO}_4^{3-}\). Previous studies indicate that increasing carbonization temperature results in higher SSA of the produced char (Cantrell et al., 2012), which in turn leads to higher \(\text{NO}_3^-\) adsorption (Hale et al., 2013; Lehmann, 2009; Yao et al., 2012). However, Akaike information criterion (AIC) was used to select the best fitting isothermal model. For \(\text{NO}_3^-\) sorption on pyrochars, AIC prefers the fitted linear model rather than the Freundlich isotherm, which indicates a non-saturated surface of biochars at increasing ion concentration of the nutrient solution. This contradicts previous studies...
which prefer Freundlich or Langmuir (Hale et al., 2013; Mizuta et al., 2004). In most cases, hydrochars showed no sorptive effect but partly, in particular for hydrochars from digestates, PO$_4^{3-}$ release into aqueous solution was observed. This finding is corroborated by Yao et al. (2012) who also found 4% PO$_4^{3-}$ leaching into aqueous solution by a hydrochar (from peanut hull) – sandy soil mixture. The digestate feedstocks and the digestates carbonized to pyrochar and hydrochar contained 10 times more phosphorus (2.51%, Table 1) than the biochars from the other two feedstock materials, which explains the high PO$_4^{3-}$ leaching.

Besides carbonization process, the feedstock material had a marked influence on the sorption behavior, which is in accordance with findings from other studies: while NO$_3^-$ sorption was observed for pyrochar from Monterey Pine (Knowles et al., 2011), sugarcane bagasse and bamboo (Mizuta et al., 2004; Yao et al., 2012), pyrochar from pure washed cacao shell and corn cob without soil led to NO$_3^-$ release (Hale et al., 2013). This implies strong adsorption capacity variations with carbonized feedstock. The three carbonized feedstocks we tested (Miscanthus, digestates, and woodchips) for pyrochars showed high correlations between NO$_3^-$ adsorption and logarithmized SSA ($R^2 = 0.57; p \leq 0.05$ for amended loamy soil / $0.64; p \leq 0.01$ amended sandy soil), and average pore size ($R^2 = 0.64$ for amended loamy soil / $0.72$ for amended sandy soil; both $p \leq 0.01$). We also found strong correlations between H : C (indicates carbonization temperature) and NO$_3^-$ adsorption ($R^2 = 0.65/0.75$ for amended loamy and sandy soil respectively; both $p \leq 0.01$). The NH$_4^+$ sorption is strongly nonlinear with increasing solution concentration (Freundlich coefficient $n = 1.1–1.5$), which indicates a limited number of cation exchange sites of biochar (Hale et al., 2013). For all pyrochars, irrespective of feedstock, pore volume ($R^2 = 0.52, p \leq 0.01$), and ash content ($R^2 = 0.66, p \leq 0.01$) correlated with NH$_4^+$ adsorption. No saturation was found for PO$_4^{3-}$, with increasing solution concentration, especially evident for pyrochars from Miscanthus and also from woodchips for our used concentration range (2.5–15 mg P L$^{-1}$). This indicates that pyrochars could remove more PO$_4^{3-}$ at higher solution concentra-
tions, which is supported by Sarkhot et al. (2013), who tested 2 g pyrolysed hardwood biochars (without soil) in 40 mL nutrient solution at higher solution concentrations in comparison to ours (up to 50 mg P L$^{-1}$).

Generally, nutrient retention potential of biochar is a result of cation or anion exchange combined with the large surface area, internal porosity and polar and nonpolar surface sites of functional groups (Hale et al., 2013; Laird et al., 2010; Lehmann, 2009). Additionally, Keiluweit and Kleber (2009) reviewed cyclic aromatic $\pi$-systems which showed specific $\pi$-electron donor-acceptor (EDA) interactions (i.e., cation-$\pi$; hydrogen-$\pi$; $\pi$-$\pi$ EDA; and polar-$\pi$-interaction) with bonding energies between 4 and 167 kJ mol$^{-1}$ to nutrients. Thus, biochars’ surface charge is assumed to be negative, resulting in low anion exchange capacity and repulsion of NO$_3^-$ and PO$_4^{3-}$ (Hale et al., 2013; Mukherjee et al., 2011). However, our results and results from previous studies showed anion adsorption the processes of which are not yet fully understood. Chun et al. (2004) and Chen et al. (2008) disproved the ability of PO$_4^{3-}$ ions to bind with negatively charged biochar surface functional groups like hydroxyls, carbonyls, carboxyls and phenolics. However, Sarkhot et al. (2013) proposed the exchange of surface hydroxyl groups on biochar with PO$_4^{3-}$ inducing a pH controlled anion sorption capacity. Another mechanism is the ability of PO$_4^{3-}$ ions to form bridge bonds using the residual charge of electrostatically attracted or ligand-bonded multivalent cations (Ca$^{2+}$, Mg$^{2+}$, Al$^{3+}$, Fe$^{3+}$) (Mukherjee et al., 2011). We could not directly verify this assumption in our study because Ca$^{2+}$ and Mg$^{2+}$ were strongly leached (see Supplement; Table S1), but we suspect residual charge of electrostatically attracted cations to bind PO$_4^{3-}$ in the double layer sheet. Klasson et al. (2014) showed that pore blocking ash-content could be reduced by washing biochars with rainwater, thereby micropore volume, total pore volume, and SSA increased. Hale et al. (2013) suggests enhanced PO$_4^{3-}$ sorption due to increasing availability of binding sites on biochar’s surface after washing. However, in our lab-experiment we did not find increasing PO$_4^{3-}$ adsorption due to washing for any type of biochar. We assume that primary bonding agents for PO$_4^{3-}$ (Ca$^{2+}$, Mg$^{2+}$, Al$^{3+}$,
Fe$^{3+}$) are leached out, which results in no adsorption to the biochar surface. Secondly, PO$_4^{3-}$ compounds from the biochar matrix itself are rinsed.

### 4.2 Soil induced effect on nutrient sorption

Our results show that pyrochars could remove NO$_3^-$ and PO$_4^{3-}$ from soil solution when added to different soils (sandy and silty loam). NH$_4^+$ was retained only in the sandy loam which confirms the findings of Yao et al. (2012), who also mixed pyrochars to a sandy soil. For pyrochars mixed with loamy soil, we found reduced sorption capacity for NO$_3^-$, NH$_4^+$, and PO$_4^{3-}$, which is corroborated by Hale et al. (2011) who reported a reduction in the sorption capacity of biochars mixed with a fine-loamy soil. Hydrochars showed little (silty loam) or no (sandy loam) sorptive effect on NO$_3^-$, NH$_4^+$, and PO$_4^{3-}$.

The adsorption capacity of biochars for nutrients interacts with the amended soil type. Generally, soil's adsorption capacity for NO$_3^-$, NH$_4^+$, and PO$_4^{3-}$ is determined by pH, CEC, AEC, SSA, organic matter content, and soil texture. Hale et al. (2011) suggest a decreased reduction in the sorption capacity of biochars caused by blocking of sorption sites by dissolved organic carbon (DOC), which could leach out from soil and may adsorb to biochars. The solubility of DOC can be increased by increasing negative charge on the DOC due to a raised pH through biochar application to soils (Alling et al., 2014). In our study, application of pyrochars led to a stronger rise in pH in the silty loam than in the sandy loam (Table S2). According to Hale et al. (2011), this could have induced higher DOC solubility in the sandy loam and the leached DOC was adsorbed by pyrochars resulting in blocked binding sites. Further, the soils tested in this study differed strongly in their texture and CEC. The silty loam contained higher amounts of multi-layer clay minerals, which led to higher adsorption competition between biochar and clay mineral surfaces. Ersahin et al. (2006) report SSA between 46.5 and 90.38 as well as 20.60 and 61.95 m$^2$ g$^{-1}$ for silty loams and loamy sands, respectively. The pyrochars we tested had SSAs between 210 and 448 m$^2$ g$^{-1}$, which are considerably higher than the SSA of the used soils. The difference in SSA between pyrochar and...
soil was larger for the sandy loam than the silty loam. This resulted in stronger adsorption potential for ions from sandy loam or nutrient solution to the pyrochars. However, the larger SSA of the silty loam enhanced the adsorption competition for ions between loamy sand and pyrochars. In addition, ions from the nutrient solution are more attracted to the silty loam than to the sandy loam or to the pyrochars. Furthermore, soil-bound ions such as NO$_3^-$, Mg$^{2+}$, Ca$^{2+}$ were leached from the silty loam and were directly adsorbed by pyrochars, suggesting that this direct adsorption may result in occupied binding sites on the pyrochars, which led to no or less adsorption of NO$_3^-$, NH$_4^+$ and PO$_4^{3-}$ from the nutrient solution.

### 4.3 Effect of biochar ageing on nutrient sorption

The ability of both pyrochar and hydrochar to adsorb NO$_3^-$ and NH$_4^+$ from soil solution was stronger for fresh biochar as compared to aged biochar (i.e., after seven months field incubation). This was an unexpected behavior and often lead to a complete loss of the biochar’s nutrient retention capacity and has rarely been studied to date. Since the overall adsorption capacity of hydrochar observed in our study was small, the ageing effect was also less important compared to pyrochars. Explanations for the decreasing nitrogen adsorption capacity of pyrochar may include: (a) binding sites of both types of biochar may be blocked with organic matter or mineral particles such as clay, (b) binding sites of pyrochar may be reduced by microbial degradation changing the char’s surface properties, which in turn leads to a diminished number of negatively charged binding sites (Cheng et al., 2008; Cheng et al., 2006; Glaser et al., 2000). But for our study, we could not explain decreasing adsorption with these mechanisms.

Such a trend of decreasing adsorption capacity over time was also reported by Bargmann et al. (2014b) who incubated 2 and 4 % hydrochars from beet-root chips with a loamy soil for 8 weeks in the laboratory. A diminished number of negatively charged binding sites may result in higher leaching of positively charged ions (such as NH$_4^+$, Ca$^{2+}$, Mg$^{2+}$, K$^+$). In our experiment, the adsorption-rate of NH$_4^+$ was reduced...
over time and \(\text{Ca}^{2+}\) as well as \(\text{Mg}^{2+}\) showed higher leaching after seven months (Table S5). The biochars used in the field experiment had not been pretreated by washing. The increased adsorption capacity of biochar for \(\text{PO}_4^{3-}\) may thus be partly a result of initially bound \(\text{PO}_4^{3-}\) that was leached from fresh biochars \((T_0)\), and was leached less after seven months \((T_1)\). However, washing did not reduce \(\text{PO}_4^{3-}\)-leaching but increased the adsorption capacity in the laboratory study. Phosphate adsorption on biochar depends strongly on pH. For our used biochars, effect on pH in the nutrient solution was lower for washed than unwashed biochars.

5 Conclusions

The nutrient retention potential of biochars (i.e., nitrate, ammonium, and phosphate) differs strongly with nutrient, biochar, and type of carbonized feedstock, as well as amended soil type. Among nine different types of biochars tested in a laboratory batch experiment, only pyrochars showed the ability to effectively retain nitrate, ammonium, and phosphate. Moreover, the nutrient retention effect was of very limited duration. After seven months in the field, around 60 to 80% of the adsorption capacity of pyrochars was lost. Underlying mechanisms are poorly understood, but our results cast doubt on the efficiency of biochar to minimize the problems of nutrient leaching from agricultural soils to the groundwater and adjacent ecosystems.

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References


Influence of different biochars on nutrient sorption

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Table 1. General properties of feedstock materials and biochars used in the laboratory study (“Lab”) and field incubation (“Field”). Data for chars used in the laboratory only derived from Eibisch et al. (2013, 2015); n.d. = not determined.

| Experiment | Feedstock | Char type | \( C \)° | pH (CaCl\(_2\)) | Ash content | C [%]  | N [%]  | S [%]  | O:C | H:C | P [%]  | Ca [%] | Mg [%] | Na [%] | K [%]  | SSA [m\(^2\) g\(^{-1}\)] | Pore volume [cm\(^3\) g\(^{-1}\)] | Average pore size [Å] |
|------------|-----------|----------|---------|-----------------|-------------|-------|-------|-------|-----|-----|-------|-------|-------|-------|-------|--------|----------------|----------------|------------------|
| Lab        | Digestates raw – – | 11.9 | 41.9 | 1.57 | 0.28 | 0.87 | 0.14 | 1.28 | 0.87 | 0.66 | 0.05 | 2.88 | 8.6 | 0.03 | 61     |
|            | Hydrochar 200 | 6.2 | 10.3 | 53.8 | 2.59 | 0.30 | 0.46 | 0.10 | 1.23 | 1.39 | 0.48 | 0.03 | 0.98 | 13 | 0.09 | 192    |
|            | Hydrochar 250 | 5.7 | 13.6 | 61.8 | 2.98 | 0.22 | 0.29 | 0.08 | 1.56 | 1.60 | 0.85 | 0.02 | 1.41 | 2.8 | 0.02 | 167    |
|            | Pyrochar 750 | 9.8 | 46.0 | 69.7 | < 1.0 | 0.18 | 0.17 | 0.04 | 2.51 | 2.91 | 1.12 | 0.24 | 8.10 | 448 | 0.28 | 12     |
|            | Miscanthus raw – – | 2.9 | 45.6 | < 1.0 | 0.07 | 0.86 | 0.13 | 0.09 | 0.22 | 0.07 | 0.01 | 0.53 | 1.0 | 0.01 | 154    |
|            | Hydrochar 200 | 4.6 | 3.9 | 58.0 | < 1.0 | 0.07 | 0.46 | 0.10 | 0.13 | 0.30 | 0.05 | 0.27 | 5.2 | 0.05 | 180    |
|            | Hydrochar 250 | 4.2 | 4.5 | 69.0 | < 1.0 | 0.07 | 0.27 | 0.08 | 0.17 | 0.30 | 0.06 | 0.01 | 0.30 | 5.8 | 0.05 | 179    |
|            | Pyrochar 750 | 9.0 | 15.0 | 76.9 | < 1.0 | 0.12 | 0.10 | 0.02 | 0.41 | 1.14 | 0.30 | 0.18 | 2.12 | 279 | 0.19 | 14     |
|            | Woodchips raw – – | 4.2 | 48.6 | < 1.0 | 0.05 | 0.71 | 0.12 | 0.07 | 0.62 | 0.07 | 0.27 | 1.6 | 0.02 | 206    |
|            | Hydrochar 200 | 4.6 | 5.0 | 59.7 | 1.07 | 0.06 | 0.40 | 0.10 | 0.08 | 0.90 | 0.07 | 0.25 | 10 | 0.09 | 180    |
|            | Hydrochar 250 | 4.8 | 5.4 | 67.7 | 2.22 | 0.06 | 0.27 | 0.08 | 0.11 | 0.59 | 0.06 | 0.03 | 0.21 | 3.5 | 0.04 | 207    |
|            | Pyrochar 750 | 8.7 | 24.6 | 68.4 | < 1.0 | 0.13 | 0.10 | 0.02 | 0.35 | 3.43 | 0.29 | 0.15 | 0.87 | 210 | 0.17 | 17     |
| Field      | Miscanthus raw – – | 2.9 | 46.3 | < 1.0 | < 0.1 | 0.28 | 0.13 | 0.09 | 0.11 | 0.09 | 0.01 | 0.52 | n.d. | n.d. | n.d.  |
|            | Hydrochar 200 | 3.8 | 3.9 | 63.8 | < 1.0 | < 0.1 | 0.15 | 0.08 | 0.13 | 0.11 | 0.13 | 0.21 | 0.13 | n.d. | n.d.  |
|            | Pyrochar 750 | 9.0 | 15.0 | 81.8 | < 1.0 | 0.10 | 0.09 | 0.01 | 0.39 | 0.35 | 0.39 | 0.03 | 1.50 | n.d. | n.d.  |
**Table 2.** General properties of the soils used for the lab and field study.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Site</th>
<th>Soil type</th>
<th>Soil texture class</th>
<th>sand [%]</th>
<th>silt [%]</th>
<th>clay [%]</th>
<th>C_{org} [%]</th>
<th>N_{tot} [%]</th>
<th>C / N</th>
<th>pH (CaCl$_2$)</th>
<th>CEC [cmol$_c$ kg$^{-1}$]</th>
</tr>
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<tbody>
<tr>
<td>Lab</td>
<td>Goettingen</td>
<td>haplic Luvisol</td>
<td>Sandy loam</td>
<td>61.5</td>
<td>32.8</td>
<td>5.8</td>
<td>1.23</td>
<td>0.10</td>
<td>12.3</td>
<td>5.6</td>
<td>4.0</td>
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<tr>
<td></td>
<td>Braunschweig</td>
<td>haplic Cambisol</td>
<td>Silty loam</td>
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<td>67.6</td>
<td>17.0</td>
<td>1.27</td>
<td>0.12</td>
<td>10.6</td>
<td>5.6</td>
<td>10.8</td>
</tr>
<tr>
<td>Field</td>
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<td>loamic Cambisol</td>
<td>Sandy loam</td>
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<td>37.1</td>
<td>5.9</td>
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<td>0.13</td>
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<td>6.4</td>
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<tr>
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<td>Loamy sand</td>
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<td>7.3</td>
<td>1.13</td>
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<td>Volkmarsof</td>
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<td>Sandy loam</td>
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<td>21.7</td>
<td>11.2</td>
<td>1.16</td>
<td>0.12</td>
<td>9.9</td>
<td>6.5</td>
<td>n.a.</td>
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</table>
Table 3. Ion concentrations of the nutrient solution and relative sorption rates of the two control soils (soil without application of biochar) at the six applied concentration levels.

<table>
<thead>
<tr>
<th>Ion</th>
<th>Nutrient solution NO$_3^-$N [mg L$^{-1}$]</th>
<th>P1</th>
<th>P2</th>
<th>P3</th>
<th>P4</th>
<th>P5</th>
<th>P6</th>
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<td></td>
<td></td>
<td>5</td>
<td>10</td>
<td>20</td>
<td>30</td>
<td>40</td>
<td>60</td>
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<tr>
<td></td>
<td>NH$_4^+$N [mg L$^{-1}$]</td>
<td>5</td>
<td>10</td>
<td>20</td>
<td>30</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td>PO$_4^{3-}$-P [mg L$^{-1}$]</td>
<td>1.25</td>
<td>2.5</td>
<td>5</td>
<td>7.5</td>
<td>10</td>
<td>15</td>
</tr>
<tr>
<td>Sandy loam</td>
<td>NO$_3^-$N [%]</td>
<td>-6</td>
<td>0.1</td>
<td>3</td>
<td>0</td>
<td>0.1</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>NH$_4^+$N [%]</td>
<td>15</td>
<td>15</td>
<td>16</td>
<td>15</td>
<td>16</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>PO$_4^{3-}$-P [%]</td>
<td>-78</td>
<td>6</td>
<td>50</td>
<td>59</td>
<td>57</td>
<td>65</td>
</tr>
<tr>
<td>Silty loam</td>
<td>NO$_3^-$N [%]</td>
<td>-58</td>
<td>-28</td>
<td>-16</td>
<td>-8</td>
<td>-9</td>
<td>-5</td>
</tr>
<tr>
<td></td>
<td>NH$_4^+$N [%]</td>
<td>54</td>
<td>52</td>
<td>49</td>
<td>39</td>
<td>36</td>
<td>33</td>
</tr>
<tr>
<td></td>
<td>PO$_4^{3-}$-P [%]</td>
<td>10</td>
<td>45</td>
<td>75</td>
<td>73</td>
<td>69</td>
<td>81</td>
</tr>
</tbody>
</table>
Figure 1. Mean $\text{NO}_3^-$-N removal rates in soil-char composites relative to the control [%] (the respective soil with no char added) for pyrochars (Pyro750) (a–b) and hydrochars derived at 200°C (Hydro200) and 250°C (Hydro250) (c–d) from Miscanthus (M), woodchips (W), and digestates (D) mixed with the sandy and silty loam soil at the six nutrient-solution levels ($n = 3$).
Figure 2. Mean NH$_4^+$-N removal rates in soil-char composites relative to the control [%] (the respective soil with no char added) for pyrochars (Pyro750) (a–b) and hydrochars derived at 200°C (Hydro200) and 250°C (Hydro250) (c–d) from Miscanthus (M), woodchips (W), and digestates (D) mixed with the sandy and silty loam soil at the six nutrient-solution levels ($n = 3$).
Figure 3. Mean PO$_4^{3-}$-P removal rates in soil-char composites relative to the control [%] (the respective soil with no char added) for pyrochars (Pyro750) (a–b) and hydrochars derived at 200°C (Hydro200) and 250°C (Hydro250) (c–d) from Miscanthus (M), woodchips (W), and digestates (D) mixed with the sandy and silty loam soil at the six nutrient-solution levels ($n = 3$).
Figure 4. Mean NO$_3^-$-N (a–c), NH$_4^+$-N (d–f), and PO$_4^{3-}$-P (g–i) removal rate relative to the control for fresh ($T_0$) and degraded ($T_1$) pyrochars and hydrochars and relative removal rate of control to blind nutrient solution of the field experiment (For all Treatments $n = 3$). Test statistics can be found in Tables S5, S7, and S8.
Figure 5. (a) $\text{NO}_3^-$, (b) $\text{NH}_4^+$, and (c) $\text{PO}_4^{3-}$ removal rates in soil-char composites relative to the control (silt loam without char) for washed and unwashed pyrochars (Pyro750) and hydrochars derived at 200°C (Hydro200) and 250°C (Hydro250) from Miscanthus (M), woodchips (W), and digestates (D). Significant differences between washed and unwashed biochars were tested with the unpaired $t$ test. $P$ values are indicating by *** < 0.01; ** < 0.05; * < 0.1 (for each treatment $n = 3$, means ± SE).