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Environmental significance

Surface precipitation of Mn^{2+} on clay minerals enhances Cd^{2+} sorption under anoxic conditions \dagger

Natacha Van Groeninge[n,](http://orcid.org/0000-0003-2587-2430) **D** Blanche Glück[,](http://orcid.org/0000-0001-9477-7774) **D** Iso Christl **D** * and Ruben Kretzschmar^D

The influence of Mn^{2+} on the sorption of metal(loid)s onto clay minerals is still unclear despite its relevance in suboxic and anoxic environments which often exhibit elevated dissolved Mn^{2+} concentrations. In this study, the effects of Mn^{2+} on Cd^{2+} sorption to two types of clay minerals, a well-crystalline natural kaolinite (KGa-1b) and a synthetic montmorillonite (Syn-1), were investigated. Batch experiments on Mn^{2+} and Cd²⁺ sorption to Ca-saturated KGa-1b and Syn-1 were conducted under anoxic conditions. At low Mn²⁺ and Cd²⁺ concentrations (1 and 5 μ M), both metals exhibited similar affinity for sorption to the clays, suggesting that elevated Mn^{2+} concentrations might effectively decrease Cd²⁺ sorption as predicted using a three-plane surface complexation model. However, competitive Mn–Cd experiments at higher concentrations (\geq 50 µM) revealed that for both clay minerals, the presence of Mn²⁺ increased $Cd²⁺$ sorption to the solid phases. Although solutions were undersaturated with respect to known Mn(II) solid phases, analysis using X-ray absorption spectroscopy (XAS) evidenced the formation of Mn(II)containing solid phases which can specifically adsorb or incorporate Cd^{2+} . This process, which was mediated by the presence of clay minerals, overcompensated the decrease in Cd^{2+} adsorption to clay surfaces due to competition with Mn^{2+} . We conclude that, contrary to predictions based on a competitive surface complexation model, elevated Mn^{2+} concentrations can contribute to decrease dissolved $Cd²⁺$ concentrations in anoxic clay-containing environments, such as contaminated sediments or flooded paddy soils. **PAPER**
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Clay minerals represent major sorbents in soils and sediments for nutrients and contaminants. In suboxic and anoxic environments, elevated concentrations of Mn^{2+} can establish as the result of reductive dissolution of Mn(m/w) solid phases, e.g., during water saturation of paddy soils. The impact of elevated Mn²⁺ concentrations on trace metal interactions with clay minerals is largely unknown, but is highly relevant for trace metal availability to organisms. In this study on Mn–Cd interactions with clay minerals under anoxic conditions, we demonstrate that elevated Mn^{2+} concentrations do not cause a net decrease in Cd²⁺ sorption due to competition, but lead to an overall increase of $Cd²⁺$ sorption due to the formation of Mn(II)-containing surface precipitates that adsorb or incorporate $Cd²⁺$.

Introduction

Elevated concentrations of cadmium (Cd) in agricultural soils are of widespread concern due to its high toxicity, long half-life in the human body, and high bioavailability in many soils. Cadmium has no known biological function and is toxic to humans,¹ causing liver and kidney damage, low bone density and increasing the risk of cancer.² It was identified by the World Health Organization (WHO) as one of the "ten leading chemicals of concern". Elevated dissolved concentrations of Cd can be found in the environment due to release from both natural sources (e.g., weathering of trace metal-enriched minerals) and anthropogenic sources (e.g., agriculture, industrial and mining activities). Cd concentrations in soils have been reported to range from geologic background concentrations, being typically below 0.10 mg kg^{-1} , up to about 10 mg kg^{-1} indicating

Institute of Biogeochemistry and Pollutant Dynamics, Department of Environmental Systems Science, ETH Zurich, Universitätstrasse 16, CHN, 8092 Zurich, Switzerland. E-mail: iso.christl@env.ethz.ch; Tel: +41 44 633 60 01

[†] Electronic supplementary information (ESI) available: Data for the characterization of the clay minerals, information about the kinetics of metal sorption, acid-base titration of the clay minerals, additional data for Cd^{2+} and Mn2+ adsorption on KGa-1b and Syn-1, detailed information about the model and the used fitting parameters, a second set of competition sorption experiments, calculation of saturation indices, protocol to ensure anoxic conditions, percentage of Cd^{2+} and Mn^{2+} sorbed in samples for solid phase analysis, concentration of Si release during sorption experiments, details on XAS data collection and reduction and additional XAS results. See DOI: 10.1039/d0em00155d

geogenically or anthropogenically elevated concentrations.³ Accumulation in plants can occur as the result of uptake from contaminated soils and contaminated water used for irrigation. Some plants like, e.g., rice, cacao, tobacco, and mushrooms are known to bioaccumulate Cd and thus are an important pathway for human exposure to Cd.⁴ Approximately 90% of the exposure to Cd for non-smokers is through diet.⁵ Herein, rice represents one of the most important pathways for Cd uptake, especially in many parts of Asia.

The retention of Cd^{2+} in soils and sediments is, as for most divalent cations, governed by sorption processes which include adsorption, polymerization, and surface precipitation.⁶ Herein, clay minerals, carbonates, as well as Fe(m)- and Mn(rw , m)-(oxy hydr)oxides are most important due to their abundance, large specific surface areas, and specific sorption properties. In addition, Cd can be retained in the solid phase when precipitating as carbonate, hydroxide, phosphate, or sulfide. In soils low in Cd, adsorption is considered the dominant mechanism controlling Cd mobility, \bar{y} whereas precipitation gains importance in soils with higher Cd activity, for example, in highly contaminated soils. In soils where exchangeable Cd represents a major fraction of total Cd,⁸ Cd is expected to be easily remobilized from mineral surfaces by competing divalent ions such as e.g., Ca^{2+} and Zn^{2+} and thus to become bioavailable to plants.⁹

Agricultural soils commonly exhibit weakly acidic to neutral pH values, at which clay minerals and $Fe(m)$ (oxyhydr)oxides are the most important mineral sorbent phases for Cd^{2+} . Several studies have investigated the adsorption of Cd to Fe (m) -(oxyhydr)oxides, showing that Cd can form inner-sphere complexes at the mineral-water interface¹⁰ and ternary mineral-organic-Cd complexes.¹¹ Similar sorption processes also occur on edge surfaces of clay minerals.¹²⁻¹⁵ To date, most studies on Cd adsorption in soils were conducted under oxic conditions applicable to upland soils, while the adsorption of Cd in suboxic to anoxic soils, such as rice paddy soils, is much less understood.

Compared to well-aerated soils, the adsorption of Cd^{2+} to mineral phases may be markedly altered in periodically anoxic soils of riparian floodplains, wetlands, and rice paddies. Even though Cd is not directly subject to redox reactions, changes in soil pore water conditions as well as reductive dissolution processes affect Cd adsorption during periods with suboxic to anoxic conditions. Oxides and oxyhydroxides of $Mn(m/w)$ become thermodynamically favorable electron acceptors for microbial respiration already at low oxygen levels still preventing $Fe(m)$ and sulfate reduction to predominate. As a result, dissolved Mn^{2+} concentrations in soil solutions increase before $Fe²⁺$ and sulfide. In a study by Kirk,¹⁶ flooding of acidic soils containing high contents of manganese and organic matter led to dissolved Mn^{2+} concentrations up to 1.7 mM after one week of submergence. For flooded alkaline soils and soils low in manganese, dissolved Mn^{2+} concentrations up to 0.2 mM were reported.¹⁶ These findings indicate that in submerged soils, Mn^{2+} becomes a major cation in soil solution and is therefore expected to compete with trace element cations (e.g., Zn^{2+} , Cd^{2+}) for adsorption to mineral surfaces influencing their fate in the

environment. During soil aeration, $Fe²⁺$ is rapidly oxidized by molecular O_2 , contrary to Mn²⁺, for which oxidation kinetics by dioxygen is much slower.¹⁷ Therefore, Mn^{2+} may persist under oxic conditions for prolonged periods and continue to act as a cation competing with trace metals for adsorption.

In a competitive situation, trace metal retention by solids may, however, be altered by metal cation-specific interactions on mineral surfaces as evidenced by Soltermann et $al.^{18}$ for Fe²⁺ and Zn^{2+} adsorption to a synthetic montmorillonite. Their results indicated the existence of specific sites which were not accessible to Zn^{2+} , but were able to induce oxidation of Fe²⁺ and enhance Fe sorption.¹⁸ These results show that ion-specific sorption can occur, leading to metal retention deviating from simple competitive adsorption. This is likely to hold true for many other systems, including Mn^{2+} and Cd^{2+} sorption to clay minerals. In addition, metal ion-specific formation of new mineral phases may be induced by the presence of clay minerals and contribute to metal sorption. If newly formed surface precipitates exhibit a lower affinity for the trace metal of interest than the clay mineral surfaces and concomitantly block the reactive clay sites, the formation of surface precipitates might lead to a decrease in trace metal sorption. However, if the newly formed phases have a similar or even higher affinity than the clay minerals, overall sorption may be increased markedly, especially in case the new phases add a signicant amount of surfaces, i.e., sorption sites. Hence, surface precipitation may considerably modify sorption patterns of trace metals. **Expiremental Science:** Process Articles. Process Articles. The model of the common on the common of the creation of the crea

At elevated concentrations, many divalent metal cations (Me^{2+}) , first-row transition series) have been shown to precipitate as $Me^{2+}Al^{3+}$ -layered double hydroxides (LDH) under alkaline conditions in presence of Al-containing minerals.¹⁹⁻²⁵ Starcher et al. demonstrated that the FeAl-LDH phase formed during Fe^{2+} sorption to an Al-bearing sorbent can sorb or incorporate Zn^{2+} when both metals were co-sorbed to the Albearing mineral.²⁶ However, Bhattacharya and Elzinga demonstrated that dissolved Mn^{2+} concentrations significantly higher than normally encountered in anoxic soils (*i.e.*, $[Mn^{2+}] > 10^{-2} M$) are required to precipitate $Mn^{2+}Al^{3+}$ -LDH, making the precipitation of this phase unlikely.²⁷ Despite the relevance of such processes affecting the mobility of metals in the environment, to date the above mentioned sorption processes have not been investigated in detail and it is difficult to predict which process would predominantly control Cd²⁺ sorption at elevated Mn²⁺ concentrations in clay-rich environments.

Therefore, the objective of this research was to assess the interactions between Mn^{2+} and Cd^{2+} during sorption to clay minerals. Kaolinite and montmorillonite are two very widespread clay minerals in the environment and were selected as the clay sorbents representing 1:1- and 2:1-type phyllosilicate minerals, respectively. $CaCl₂$ was chosen as the background electrolyte, as calcium is often the predominant divalent cation in soils.²⁸ Acid–base titrations and batch experiments on metal sorption of Mn^{2+} and Cd^{2+} onto the Ca-clays were carried out. The data were used to optimize parameters of a two-site Three-Plane Model (TPM)²⁹ to best describe H⁺, Ca²⁺, Mn²⁺, and Cd²⁺ adsorption to the clay surfaces. Predictions of competitive $Mn^{2+}-Cd^{2+}$ adsorption as calculated with model parameters

based on binary adsorption data (Mn–Ca and Cd–Ca clay systems) were compared to experimental results in ternary Mn– Cd–Ca clay systems. Additionally, solid phase speciation was carried out for Mn and Cd using synchrotron X-ray absorption spectroscopy (XAS) to identify the dominant sorption mechanisms.

Materials and methods

Clay minerals

A well-ordered kaolinite (KGa-1b) and a synthetic montmorillonite (Barasym SSM-100), referred to as KGa-1b and Syn-1 in the following, were used in this study. KGa-1b and Syn-1 were selected as the solids, as both are low in Fe in order to minimize potential oxidation of Mn^{2+} by structural Fe^{III}. Both materials were purchased from the Source Clays Repository of the Clay Minerals Society (Purdue University, West Lafayette, IN).

Particles with a hydrodynamic diameter of less than $2 \mu m$ were isolated by repeated centrifugation and dispersion steps. Briefly, clay particles were dispersed by adjusting the suspension pH to 9.0 with NaOH and subsequent ultrasonication for 3 min prior to centrifugation at 100g (600 rpm) for 5 min (Mistral 6000). The supernatant containing the particles smaller than $2 \mu m$ was collected and the procedure was repeated until the supernatant was clear. A yield of 62% for KGa-1b and 95% for Syn-1 was obtained after size fractionation. Size-fractionated particles were saturated with calcium by adding $CaCl₂$ solution to achieve a concentration of 0.1 M. The suspensions were subsequently placed on an overhead shaker overnight and then centrifuged at 200g (1200 rpm) for 5 min for both KGa-1b and Syn-1. To ensure complete Ca saturation, the Ca saturation step was repeated two more times by adding 0.1 M CaCl₂ solution and shaking the suspensions for 3–4 hours before centrifugation. After Ca saturation, excess salt was removed by repeated washing with doubly deionized (DDI) water (\geq 18.2 M Ω cm, Milli-Q, Millipore, Merck KGaA, Darmstadt, Germany) until the supernatants were found to be free of chloride using $AgNO₃$. The final products were freeze-dried and stored as dry powders.

Characterization of clay minerals

The size-fractionated and Ca-saturated KGa-1b and Syn-1 were analyzed as random-oriented powder samples by X-ray diffraction (XRD, D8 Advance, Bruker) to test for mineralogical purity. Diffractograms were recorded in Bragg–Brentano geometry from 3 to 80 $^{\circ}$ 2 θ with a step size of 0.02 $^{\circ}$ 2 θ and 4 s acquisition time per step, using Cu $K\alpha_{1,2}$ radiation. Energy-dispersive X-ray fluorescence analysis (XRF, XEPOS, Spectro) using pressed pellets was performed for elemental analysis of the clay minerals. The specific surface areas of the clay minerals were measured by multi-point N_2 -BET analyses (ASiQwin, Quantachrome).

The measured diffractogram of KGa-1b indicated trace impurities of TiO₂ and for Syn-1 trace amounts of γ -AlOOH which is in agreement with published results for KGa-1b and Syn-1 (see Fig. S1[†]).³⁰ Total iron contents as determined using XRF amounted to 1.41 g kg⁻¹ for KGa-1b and 0.26 g kg⁻¹ for Syn1 (see also Table S1†). We determined a N₂-BET surface area of 12.8 ± 0.1 m² g⁻¹ for KGa-1b and 95 \pm 2 m² g⁻¹ for Syn-1. Mermut and Cano³¹ reported iron contents of 2.5 \pm 0.3 g kg⁻¹ and 0.9 \pm 0.2 g kg⁻¹ Fe for KGa-1b and Syn-1, respectively. N₂-BET surface areas as published by Dogan et $al.^{32}$ amounted to $13.1 \pm 0.1 \text{ m}^2 \text{ g}^{-1}$ and $118 \pm 1 \text{ m}^2 \text{ g}^{-1}$, for KGa-1b and Syn-1, respectively. The minor differences in Fe contents and surface areas between our measurements and literature are likely due to the sample pretreatment applied in this study, whereas literature values given above refer to untreated source clays.

Potentiometric titrations of KGa-1b and Syn-1 suspensions were carried out at three CaCl₂ concentrations $(0.01, 0.03,$ and (0.1 M) with an automated titration setup³³ located in a thermostated room at 25 \pm 1 °C. The setup was equipped with four burettes (Dosimat 605, Metrohm) containing 0.05 M HCl, \sim 0.02 M Ca(OH)₂, 0.6 M CaCl₂, and CO₂-free DDI water, respectively. The four burettes were connected to a computer by a Microlink MF18 interface (Biodata, Manchester, UK). All solutions were prepared with $CO₂$ -free DDI water. For the acid– base titrations, 2.5 g KGa-1b and 0.75 g Syn-1, respectively, were suspended in 40 mL DDI water in a Teflon vessel, which was continuously purged with CO_2 -free, water saturated N₂-gas during the whole experiment. At the beginning, the starting ionic strength (0.01 M) was adjusted by adding CaCl₂ solution. As a preconditioning prior to the acid–base titration, the pH of the suspension was first increased to pH_1 9 to disperse the particles, then adjusted to pH 5 and kept at this pH for one hour to remove $CO₂$. During the subsequent acid–base titration, the clay suspensions were titrated in the potential range of 200 mV to -180 mV (about pH 3.8–10) with increments of 15 mV as the target step size for acid and base additions, respectively. After each titrant addition, the suspension was first stirred for 2 minutes before monitoring the pH value. The ionic strength was automatically readjusted after each acid or base addition using either CaCl₂ solution or DDI water. Reported data correspond to pH values recorded once pH drift was below 0.05 mV per minute or after a maximum equilibration time of 30 minutes. After completion of a full acid–base titration cycle, the ionic strength was increased to the next higher ionic strength (0.03 and 0.1 M). This way a set of acid–base titration data at three different ionic strengths was recorded in one experiment ensuring that there was no experimental offset bias between the data measured at the different ionic strengths. Environmental Science: Processes & Impacts

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> The proton binding by the clay surfaces was calculated as a $H⁺-OH⁻$ mass balance. The additions of acid and bases, the concentrations of H^+ and OH⁻ in the electrolyte solutions, as well as the consumption of OH $^{-}$ by the formation of aqueous Ca hydroxide complexes and proton consumption by dissolved silicate were considered in the calculations (see ESI† for species). The concentration of dissolved species was calculated for each data point with the geochemical speciation program PHREEQC (version 3) (ref. 34) using measured pH and the known total Ca and Cl concentrations as input. For silicate, the total dissolved Si concentration as determined at the end of the titration experiment was used. Hydrolysis of dissolved Al was not considered as including a correction for Al hydrolysis as based on total dissolved Al determined at the end of the

experiment led to an obvious overcorrection of proton consumption (see Fig. S5†).

Metal sorption experiments

The sorption of Mn^{2+} and Cd^{2+} on Ca-saturated KGa-1b and Syn-1 was studied in batch experiments as a function of pH (4–10) and CaCl₂ concentration $(0.1, 1, 3, 10,$ and 30 mM) at fixed total sorbate concentrations $(10^{-3}$ to 10^{-8} M) at 25 \pm 1 °C. Based on kinetic pre-experiments (see Fig. S2 and S3†), an equilibration time of 24 h was chosen for all experiments. Clay suspensions were prepared with a solid-to-solution ratio of \sim 1 g L⁻¹ in 0.1, 1.0, 3.0, 10, and 30 mM CaCl₂, respectively. All solutions were prepared with CO_2 -free DDI water to avoid the formation of carbonate solids at alkaline pH values contributing to the removal of Mn^{2+} and Cd^{2+} from solution. Before starting the sorption experiments, suspensions were equilibrated overnight at pH 5 to remove CO_2 . Subsequently, $CdCl_2$ or $MnCl_2$ solutions were added to the clay suspensions. The pH was adjusted by adding a pre-determined amount of base (0.01 or 0.1 M NaOH) to reach the target pH value. After 24 hours of equilibration on an end-over-end shaker, samples were passed through 0.22 μ m Nylon filters (VWR International), acidified, and analyzed for Ca, Al, Si, Cd, Mn, Zr, V and Fe by inductively coupled plasmaoptical emission spectrometry (ICP-OES, Vista MPX, Varian) or mass spectrometry (ICP-MS, 8800 QQQ-ICP-MS, Agilent). The pH of the clay suspensions was measured for all samples at the end of the 24 hours equilibration period. Competitive sorption experiments were performed at a $CaCl₂$ concentration of 10 mM, which was chosen to saturate non-specific cation exchange sites with calcium and more selectively observe specific sorption of Mn^{2+} and Cd^{2+} on edge-sites of the clays. The influence of various Mn^{2+} concentrations (0.05 mM and 0.5 mM MnCl₂) on the sorption of Cd^{2+} at trace concentrations $(0.001 \text{ mM}$ and (0.05 mM) was studied as function of pH $(4-10)$ for both KGa-1b and Syn-1. **Exception Exception Control Common access Article on Distribution (and Common and Control Common and Creative Common and Creative Common and Creative Common and Creative Common Common Common Common Common Common Common C**

Batches for binary Mn–Ca and ternary Mn–Cd–Ca sorption experiments were prepared in an anoxic glovebox $($ <10 ppm O_2) using O₂-free solutions to prevent catalyzed oxidation of Mn^{2+} by the clay minerals.³⁵ For equilibration, the anoxic Mn^{2+} -containing samples were sealed in two layers of gas-tight aluminum foil inside the anoxic glovebox but equilibrated on an end-overend shaker outside the glovebox for 24 hours, similarly as for the samples containing Cd^{2+} , only.

Modeling

Clay minerals exhibit permanent negative charge due to isomorphic substitution in the crystal structure,^{36,37} giving rise to cation exchange sites on face surfaces of the clay platelets. In addition, reactive surface hydroxyl groups located at edge surfaces can have pH-dependent charge. Both types of surface sites contribute to total cation binding. In the modeling approach, one type of permanently charged sites, assigned to face surface areas and denoted as Ex^- , and one type of pHdependent charged sites, assigned to edges surface areas and denoted as \equiv SOH^{-0.5}, were considered. An edge-to-face surface area ratio of 1:4 was assumed and corresponding fractions of the total N_2 -BET surface area were assigned to edge and face surfaces, respectively. This percentage of edge surface area is in line with published aspect ratio measurements for KGa-1b.³⁸⁻⁴⁰ As the permanent charge of KGa-1b was reported to be located predominantly in the tetrahedral sheet, 50 percent of the face surface were assumed to carry small amounts of permanent negative charge. Correspondingly, reactive surface areas of edge and face sites were set to 2.56 $m^2 g^{-1}$ and 5.12 $m^2 g^{-1}$ for KGa-1b and to 18.71 $m^2 g^{-1}$ and 74.84 $m^2 g^{-1}$ for Syn-1, respectively (see Tables S2 and S3†). Ion sorption on both face and edge surfaces was described with a 1 pK Three-Plane surface complexation model (TPM)²⁹ (see Fig. S4†). More detailed information about the modeling parameters can be found in the ESI.†

PHREEQC coupled with the parameter optimization code $PEST⁴¹$ was used for data fitting. The stability constants for the formation of aqueous complexes as well as solubility product constants of possible solid phases were taken from Smith and Martell.⁴² The acid–base titration data recorded in CaCl₂ background electrolyte were modeled in order to obtain the surface parameters of the clay minerals, namely the site densities of \equiv SOH^{-0.5} on edge and Ex⁻ on face surfaces and the protonation and surface complex formation constants for Ca^{2+} and Cl^- . Capacitances of both the 1- and 2-plane were fixed to 1 and 5 F m^{-2} , respectively.¹⁰ Surface complex formation of Mn²⁺ and $Cd²⁺$ on the clay minerals were optimized using the binary metal adsorption data, while fitted surface parameters were fixed according to the best fit of acid-base titration data. Finally, the competitive effect of Mn^{2+} on Cd²⁺ adsorption was predicted by running forward calculations with PHREEQC using all optimized surface complex formation parameters. Model predictions were compared to experimental data on competitive sorption.

X-ray absorption spectroscopy

A second set of samples was prepared to investigate the solid phase speciation of Mn and Cd and to analyze the effects of an elevated Mn²⁺ concentration (2.5 mM) on the sorption of Cd^{2+} (0.25 mM) to clay minerals at pH \sim 8 using XAS. The samples were prepared with a solid-to-solution ratio of \sim 5 g clay per L and a CaCl₂ concentration of 50 mM in order to maintain similar conditions to the batch experiments described above. The suspensions were equilibrated in 250 mL Nalgene® bottles for 1 day under anoxic conditions while stirred at 800 rpm. After equilibration in the anoxic glovebox, 10 mL aliquots were taken and filtered for liquid analysis using ICP-OES or ICP-MS. For XAS analysis an aliquot of the suspension was filtered through a 0.22 µm cellulose nitrate filter. The collected solid was washed with deoxygenated $CO₂$ -free DDI and subsequently dried in the glovebox in the dark.

Solid samples from these batch experiments were analyzed by Mn K-edge (6539 eV) and Cd K-edge (26 711 eV) X-ray absorption spectroscopy (XAS) at the SAMBA beamline of SOL-EIL (Saint-Aubin, France). Dried solids were homogenized and pressed into 13 mm pellets and sealed between Kapton® tape. For transport to the synchrotron, samples were sealed in two layers of gas-tight aluminum foil under N_2 gas. Two samples were additionally prepared as oriented clay films and measured with polarized-XAS ($\alpha = 10^{\circ}$, 35°, 55°, and 80°). The oriented samples were prepared by collecting the clay on a cellulose nitrate filter, cutting and stacking the filters, and sealing them between Kapton® tape. All measurements were conducted at 25 K to avoid beam damage and oxidation of $O₂$ -sensitive samples. Sample spectra were evaluated by linear combination fitting (LCF) to obtain information about the speciation and the (trans) formation of solid-phases. Shell-fit analyses of the extended Xray absorption fine structure (EXAFS) spectra were performed in order to gain information about the short range local coordination environment of Mn and Cd. The nature of the backscattering atoms in the Mn sorption samples was determined using Morlet wavelet transforms (WT) of k^3 -weighted EXAFS data using the Fortran version of the HAMA software developed by Funke et al.⁴³ Details on all measurements, data reduction, and analyses are provided in the ESI.†

Results and discussion

Sorption of Mn^{2+} and Cd^{2+} to Ca-clays

The pH-dependent sorption of Mn^{2+} and Cd²⁺ to KGa-1b and Syn-1 is shown in Fig. 1, S7, and S8† along with the best surface complexation model fits. For both clay minerals, an increase of Mn^{2+} and Cd²⁺ sorption was observed with increasing pH (Fig. 1, S7 and S8†). Mn^{2+} and Cd²⁺ sorption isotherms (Fig. S9) and S10†) did not exhibit an apparent sorption maximum for either cation when plotted on a log–log scale. This may indicate that the sorption sites had not been saturated with respect to

 Mn^{2+} or Cd²⁺ within the studied range of metal concentration in solution or that additional sorption processes aside of adsorption (e.g., surface precipitation) contributed to the overall uptake (sorption) of Mn^{2+} and Cd^{2+} (see isotherms in Fig. S9 and S10†). Furthermore, the sorption of both metal cations decreased with increasing Ca^{2+} concentration, especially in the acidic pH range. This observation points towards the presence of two distinct types of surface sites on the clay mineral surfaces, one site dominating sorption in the acidic pH range and a second site dominating sorption under neutral to alkaline conditions. In the acidic pH range, elevated Ca^{2+} concentrations competed for cation binding indicating that cation sorption was dominated by cation exchange. Under neutral to alkaline conditions, cation sorption was governed by sorption on pHdependent edge sites, for which Cd^{2+} and Mn^{2+} showed a higher sorption affinity than Ca^{2+} . A closer look at the effects of Ca^{2+} concentration furthermore reveals that the sorption of Cd^{2+} was decreased to a larger extent by increasing Ca^{2+} levels than was the sorption of Mn^{2+} . This finding implies that below pH 7, Ca^{2+} more effectively competed with Cd^{2+} than with Mn^{2+} . Fitted affinities of each divalent cation to permanently charged sites ($log K_{Ex2Me}$) were similar for both clay minerals and followed the order $Mn^{2+} > Cd^{2+} > Ca^{2+}$ (Tables S2 and S3†), which is in line with the above mentioned observations. Environmental Science: Processes & Impacts

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From neutral to alkaline pH, the sorption of cations is governed by sorption to edge sites. For both clay minerals, similar sorption of Cd^{2+} and Mn²⁺ to edge sites was observed (Fig. 1, S7 and S8†). This is in line with the fitted affinities of Cd and Mn hydroxide surface complexes (Tables S2 and S3†), which

Fig. 1 Sorption of Cd²⁺ and Mn²⁺ to (a, b) Syn-1 and (c, d) KGa-1b as a function of pH using Mn²⁺ and Cd²⁺ concentration of 1 and 5 µM at various CaCl₂ concentrations (from 1 mM to 30 mM CaCl₂, for DDI water samples CaCl₂ was of \sim 0.1 mM and \sim 0.001 mM for Syn-1 and KGa-1b respectively). The solid concentration was $\sim\!\!1$ g L $^{-1}$. Lines represent fits for Cd²⁺ and Mn²⁺ sorption described with the two-site TPM, calibrated with the protonation data.

differed negligibly. Additionally, this observation is in good agreement with the known minor differences in metal first hydrolysis between these two metal cations (Me²⁺ + H₂O \rightleftharpoons MeOH⁺ + H⁺ with $\log K_{\text{MnOH}^+} = 3.4$ and $\log K_{\text{CdOH}^+} = 3.9$) as affinities of cation adsorption to reactive surface hydroxyl groups was reported to be related to the first cation hydrolysis constant.⁴⁴–⁴⁶

Competitive sorption between Mn^{2+} and Cd^{2+} on Ca-clays

Experimental data and modeling of the Mn^{2+} and Cd^{2+} sorption data revealed that Mn^{2+} had a slightly higher affinity than Cd^{2+} for sorption to face sites, being the dominant sorption site on both clay minerals in the acidic pH range, and that the affinities for surface complexation on edge sites were similar for both metal cations (see ESI, Tables S2 and S3 and Fig. S13†). In natural systems, Mn^{2+} concentrations usually exceed Cd^{2+} concentrations in solution during anoxic conditions. Especially in submerged soils, Mn^{2+} concentrations can be highly elevated, reaching up to millimolar concentrations. A high Mn^{2+} to Cd^{2+} ratio in solution is therefore expected to decrease Cd adsorption to clay minerals as based on the findings on the binary Mn–Ca and Cd–Ca sorption experiments presented above. Predictions based on the calibrated Three-Plane Model, which considers Mn^{2+} –Cd²⁺ competition during adsorption, illustrate this effect which becomes prominent at total Mn concentrations exceeding 50 μ M (see lines in Fig. 2). However, the data of the competitive sorption experiments revealed that Cd^{2+} adsorption increased with increasing total Mn^{2+} concentration over the entire pH range (see Fig. 2). The clear difference between model predictions and experimental data indicates that the presence of Mn^{2+} triggered a process that counteracted competitive metal adsorption and led to a net increase in Cd^{2+} sorption. We therefore hypothesized that the observed increase of Cd^{2+} sorption in the presence of Mn^{2+} was caused by a newly formed, Mn-containing solid phases. This newly formed Mn-containing solid phases may serve as an additional sorbent for Cd^{2+} or incorporate Cd^{2+} in its structure. Ainsworth *et al.* found that $Cd²⁺$ could be incorporated in hydrous ferric oxide (HFO) by recrystallization.⁴⁷ Similarly, Cd^{2+} could be sorbed by a newly formed Mn-containing solid phases through more than just adsorption. Thermodynamically, elevated Mn^{2+} concentrations are expected to form pyrochroite $(Mn(OH)_2)$ at alkaline pH (see Fig. S14†). But the studied suspensions were undersaturated with respect to the solubility of known Mn- and Cd-phases (Table S5†). Alternatively, Me (n) Al-LDH phases, which have already been evidenced in literature for alkaline conditions,¹⁹–²⁵ may have formed. It has been demonstrated that LDH phases are able to sorb or incorporate contaminants.²⁶ However, it is unknown to what extent Mn-containing LDH phases are able to sorb or incorporate Cd^{2+} .

X-ray absorption spectroscopy

To clarify the predominant mechanisms leading to the observed deviations from the model predictions in competitive systems, the solid phase speciation of both Cd and Mn was investigated by synchrotron XAS. As observed for the competitive sorption

Fig. 2 Sorption of Cd^{2+} (50 μ M) to (a) Syn-1 and (b) KGa-1b in 10 mM CaCl₂ with various Mn²⁺ concentrations (0 mM, 50 μ M and 500 μ M) as a function of pH. The solid concentration was ~ 1 g L⁻¹. Lines show the predicted effect of Mn^{2+} on Cd^{2+} sorption assuming that competition for adsorption sites is the dominant type of interaction in a competitive system.

experiments, the XAS samples displayed an increase of Cd^{2+} sorption upon addition of Mn²⁺ to the clay suspension at pH \sim 8 (see Fig. 3). The marginally higher suspension pH for samples with Mn^{2+} is insufficient to explain the pronounced increase of $Cd²⁺$ sorption in KGa-1b samples. Thus, XAS samples furthermore point towards the formation of a newly formed Mncontaining solid phases, which adsorbed or incorporated $Cd²⁺$.

The Mn K-edge XANES spectra for selected Mn^{2+} sorption samples are shown in Fig. 4 together with relevant Mn reference compounds. The position of the white line between 6553 and 6557 eV in the Mn XANES spectra for all the sorption sample suggests a valence state close to 2+ for the sorbed Mn.

Dissolved Si released from Syn-1 and KGa-1b during 1 day equilibration between pH 6 and 10 amounted on average to \sim 0.016 mmol Si per g clay and 0.0016 mmol Si per g clay for Syn-1 and KGa-1b (see Fig. S16†), respectively, when Mn was absent. In the presence of Mn, up to 2 times lower dissolved Si concentrations were observed (see Fig. S17†). Likewise, the clay mineral samples equilibrated with Cd displayed a decrease in dissolved Si concentration, but to a smaller extent. These findings suggest that the released Si was adsorbed or incorporated

into newly formed phases. Alternatively, the addition of Mn and Cd to the clay suspension may have limited the dissolution of the clay minerals and thereby decreased the Si release. It has been shown by Soltermann et al.⁴⁸ that Fe^{2+} can be taken up by the clay minerals, leading to the formation of Fe-bearing phyllosilicate structures. Similarly, the observed decrease of the concentration of dissolved Si upon addition of Mn^{2+} may be due to the incorporation of Mn and Si into a newly crystallized

error bars indicate the standard deviation of duplicates.

phyllosilicate phases. To investigate the possible solid phases Mn species formed, linear combination fits (LCF) of Mn K-edge XANES spectra were performed (Fig. S19 and Table S7†). A two-component LCF model was selected as the addition of more components did not decrease the reduced χ^2 -values by at least a factor two.⁴⁹ LCF results showed that the sorbed Mn remained as Mn^{2+} (Table S7†), as was already indicated by the white line position. The XANES spectra of sorption samples were adequately fitted by including Mn(II)Al-layered double hydroxide (LDH) (with generic composition $[M_{1-x}^{2+}M_x^{3+}({\rm OH})_2]({\rm Cl}^-)_{x/n}$ yH₂O, 40–66%) and $Mn(\pi)$ -hydroxide $(Mn(OH)_2, 34-38%)$ or rhodonite $(MnSiO₃, 42–60%)$. To fit the Mn spectra of the Syn-1 samples, the reference compound rhodonite (42–60%) was required, which suggests the incorporation of the released Si into a newly formed Mn-silicate phases. This is in line with the observed higher dissolved Si concentration for Syn-1 compared KGa-1b, which would promote the formation of Mn-phyllosilicates upon Mn^{2+} addition to Syn-1 compared to KGa-1b were $Mn(OH)$ ₂ formation was promoted.

It has been shown in many studies that at elevated concentrations of divalent metal cations (Me^{2^+}) , first-row transition series), $Me(II)Al(III)$ -LDH can form under alkaline conditions in the presence of Al-containing minerals.^{19-22,24-27,50-52} The formation of Mn(II)Al-LDH formation is favored over $Mn(OH)_{2}$ in environments where Al^{3+} is present, because MnAl-LDH phases

X-ray energy (eV)

Fig. 4 Normalized XANES spectra of the Mn K-edge of reference materials (black lines) and clay mineral samples reacted with 2.5 mM Mn^{2+} in absence (blue line) or presence (green line) of Cd²⁺ (0.25 mM $Cd²⁺$). Clay mineral samples were equilibrated for 1 day under anoxic conditions at \sim pH 8 with a solid concentration of \sim 5 g L⁻¹ KGa-1b or Syn-1. Vertical dashed lines correspond to peak maxima of $Mn(OH)_{2}$ and manganite representing Mn(II)- and Mn(III)-containing solid phases, respectively.

are by two orders of magnitude less soluble than is $Mn(OH)_2$ ²⁷ Based on solubility calculations by Bhattacharya and Elzinga, the formation of $Mn(u)$ Al-LDH did not appear likely to occur in our samples, as very high Mn^{2+} concentrations (>10⁻² M) are required to induce the precipitation of such phases at nearneutral pH.²⁷ However, LCF of XANES spectra suggested that a large fraction of the sorbed Mn^{2+} was present as MnAl-LDH. Potentially, this may be explained by the ability of Si to be incorporated in LDH phases, which increases the stability of LDH precipitates.⁵³ Because including a reference spectrum for Mn^{2+} sorbed to illite (IMt from the Source Clay Repository of the

Clay Minerals Society) did not improve the LCF signicantly, we concluded that surface adsorption of Mn^{2+} to the clay mineral, under these conditions, was a minor contribution and the majority of Mn^{2+} was sorbed by formation of secondary phases.

In order to get more detailed information about the binding environment of Mn and Cd on Syn-1 and KGa-1b, shell-by-shell fitting of the Fourier transforms (FT) of k^3 -weighted Mn and Cd K-edge EXAFS spectra (see Fig. 5) were performed. For all Mncontaining samples, a first shell was observed at \sim 1.7 Å (uncorrected for phase shift), representing the first oxygen shell surrounding the Mn atoms. A second shell with $R + \Delta R$ at 2.8 Å was observed, which can be attributed to Mn and Al or Si backscatterers. Due to the similar or even identical crystallographic distances of Mn²⁺ and Al³⁺ in LDH phases (3.318 Å),⁵⁴ it is difficult to unequivocally differentiate these two atoms. Additionally, constructive interferences from $Si⁴⁺$ atoms could occur when present at a slightly larger distance of 3.393 \AA like e.g., in rhodonite.⁵⁵ Paper

City Minerals Society) did no improve the LCF significantly, we will a conclination number buceven 2.4-5.0. The finite common comm

For the Mn-containing samples (see Table 1 and Fig. 5a), the first shell was fitted with 4.0–5.5 O atoms at a distance of 2.18 \AA , which is characteristic for an octahedral arrangement of O around Mn(m).⁵⁴ Fitting of the FT of the $k^3\chi(k)$ EXAFS spectra was unsatisfactory when including a Mn–Si or Mn–Al backscattering path for the second shell. Hence, the second shell was fitted with a Mn atom located at radial distance of $3.27-3.29$ Å and

with a coordination number between $2.4-5.0$. The fitted distance of the second shell corresponds to a Mn atom around $Mn(\pi)$, confirming that a Mn-containing solid phases formed under anoxic conditions.⁵⁴ The higher second-shell CN found for KGa-1b samples indicate that the formation of a $Mn(\mathbf{I})$ containing solid phases was more favored in presence of KGa-1b than in presence of Syn-1. This difference may be due to sorption of Mn^{2+} to KGa-1b compared to Syn-1 (Table S7 \dagger), which resulted in higher dissolved Mn^{2+} concentration and thus enhanced the formation of $Mn(OH)_2$ in the KGa-1b suspensions.

The k^3 -weighted EXAFS spectra of the Syn-1 samples showed a truncated oscillation at 7–8 \AA^{-1} similar to that of the MnAl-LDH reference (Fig. S20†). This feature has been identified as a signature feature for the presence of Al in the octahedral $Me^{II}(OH)_{2}$ sheets formed by divalent cations such as Ni, Zn, Co, and Fe.⁵² To gain more information about the possible incorporation of Al into the Mn-containing solid phases, wavelet transforms (WT) of EXAFS spectra were performed for the samples and compared to the references $Mn(OH)_2$, MnAl-LDH, and rhodonite. WT complements the FT by resolving the kdependence of the absorption signal, allowing better distinction between heavier and lighter backscattering atoms. The WT of the second coordination shell of the Mn-containing samples and references are shown in Fig. S22 and S23.† The WT plots of

Fig. 5 Fourier transform real parts and magnitudes of the Mn and Cd K-edge EXAFS spectra of selected samples with corresponding shell fit. Points indicate experimental data and solid lines show the model fits. The fits were performed over a R-range of 1.2–3.5 Å (k -weight = 3) except for the samples equilibrated with Cd^{2+} which were fitted over the R-range of 1.2–2.5 Å. The corresponding fitting parameters are reported in Tables 1 and 2.

Table 1 Shell-fitting parameters for Mn K-edge EXAFS spectra of Mn-containing samples which were equilibrated for 1 day under anoxic conditions in absence or presence of Cd^{2+} .^{a} Parameter uncertainties are given in parentheses

		$Mn-O$			Mn–Mn					
Syn-1	pН	CN^b	$R^{c}(\AA)$	$\sigma^{2d}(\AA^2)$	CN^b	$R^{c}(\AA)$	$\sigma^{2d}(\AA^2)$	ΔE_0^e	Red. χ^{2f}	R -Factor ^{β}
No Cd^{2+} With Cd^{2+}	8.1 8.4	4.0(0.2) 4.6(0.3)	2.18(0.01) 2.18(0.01)	0.006(0.006) 0.007(0.001)	2.4(0.3) 2.8(0.5)	3.29(0.01) 3.27(0.01)	0.006(0.001) 0.008(0.001)	3.13(0.50) 2.13(0.68)	10 7	0.0014 0.0025
		$Mn-O$			Mn-Mn					
KGa-1b	pН	CN^b	$R^{c}(\AA)$	$\sigma^{2d}(\AA^2)$	CN^b	$R^{c}(\AA)$	σ^{2d} (Å ²)	ΔE_0^e	Red. χ^{2f}	R -Factor ^g
No \mathbf{Cd}^{2+} With Cd^{2+}	8.7 8.2	5.5(0.5)	2.18(0.01)	0.005(0.001)	5.0(0.6)	3.27(0.01)	0.006(0.001)	1.68(0.74)	43 42	0.0028 0.0041
equilibration period for each sample.		5.1(0.5)	2.18(0.01) (coordination number). ^c Mean half	0.005(0.001) ^a The amplitude reduction factor S_0^2 , was set to 0.7 based on first shell optimization for all sorption samples (<i>R</i> -range 1.1-2.5 Å). ^b Path degeneracy path length. reduced $\chi^2 = (N_{\text{idp}}/N_{\text{pts}})\sum((\text{data}_i - \text{fit}_i)/\varepsilon_i)^2(N_{\text{idp}} - N_{\text{var}})^{-1}$. N_{idp} , N_{pts} and N_{var} are, respectively, the number of independent points in the model fit (7), the total number of data points (141), and the number of fit variables (7). ε_i is the uncertainty of the i^{th} data point. ^g R-Factor, normalized sum of squared residuals $\left(\sum (data_i - fit_i)^2 / \sum data^2\right)$. Note: displayed pH values correspond to the pH measured at the end of the	4.3(0.7)	3.30(0.01)	0.006(0.001) d Debye-Waller parameter. e Energy-shift parameter.	1.63(0.89)	$\frac{f}{f}$ Fit	accuracy;

Table 2 Shell-fitting parameters for Cd K-edge EXAFS spectra of Cd sorption samples which were equilibrated for 1 day under anoxic conditions in absence or presence of Mn²⁺.^{a} Parameter uncertainties are given in parentheses

$Syn-1$	pН	$Cd-O$			$Cd-Mn$			ΔE_0^e e	Red. χ^{2f}	R -Factor ^g
		CN^b	$R^{c}(\overset{\circ}{\mathbf{A}})$	σ^{2d} (\AA^2)	CN^b	$R^{c}(\AA)$	$\sigma^{2d}(\AA^2)$			
No Mn^{2+}	7.8	3.9(0.7)	2.33(0.03)	0.006				11.89(3.65)	1318	0.0271
With Mn^{2+}	8.4	3.9(0.2)	2.28(0.01)	0.006	1.0(0.2)	3.33(0.02)	0.004	6.03(1.39)	80	0.0067
		$Cd-O$			$Cd-Mn$					
KGa-1b	pH	CN^b	$R^{c}(\mathring{A})$	σ^{2d} (\AA^2)	CN^b	$R^{c}(\AA)$	σ^{2d} (A^2)	ΔE_0^e e	Red. χ^{2f}	R -Factor ^g
No Mn^{2+}	8.1	3.7(0.5)	2.28(0.03)	0.006				7.05(2.82)	3	0.0134
With Mn^{2+}	8.2	4.2(0.2)	2.26(0.01)	0.006	1.5(0.2)	3.34(0.01)	0.004	3.26(0.91)	24	0.0025

 a The amplitude reduction factor ${S_0}^2$, was set to 0.7 based on first shell optimization for all sorption samples (R-range 1.1–2.5 Å). b Path degeneracy (coordination number). ^c Mean half path length. ^d Debye–Waller parameter. Debye–Waller parameter was fixed to 0.006 Å² for Cd–O based on the σ^2 obtained for first shell fitting of all sorption samples. σ^2 (Mn) was fixed to 0.004 Å² based on σ^2 fitted for Mn(OH)₂ reference. ^{*e*} Energy-shift parameter. ^{*f*} Fit accuracy; reduced $\chi^2 = (N_{\text{idp}}/N_{\text{pts}})\sum ((\text{data}_i - \text{fit}_i)/\varepsilon_i)^2 (N_{\text{idp}} - N_{\text{var}})^{-1}$. N_{idp} , N_{pts} and N_{var} are, respectively, the number of independent points in the model fit (4–7), the total number of data points (101), and the number of fit variables (3–5). ε_i is the uncertainty of the *i*th data point. ^g R-Factor; normalized sum of squared residuals $\left(\sum\limits_i (\text{data}_i - \text{fit}_i)^2 / \sum\limits_i \text{data}^2\right)$. Note: displayed pH values correspond to the pH measured at the end of the equilibration period for each sample.

the newly formed Mn-containing solid phases or the incorporation of Cd in the formed Mn-containing solid phases.

Fitting the first shell of Cd-containing samples required a Cd–O backscattering path at a radial distance between 2.26– 2.33 Å and coordination number (CN) between 3.7-4.2. The minor changes in CN or radial distance of the first shell suggest that the type of surface complex Cd formed did not change substantially upon the addition of Mn^{2+} . Shell fitting of the minor second shell for the Cd^{2+} -containing samples in absence of Mn^{2+} was not possible, which may indicate that only small amounts of Cd precipitates were formed. The second shell observed in the FT of the Cd K-edge EXAFS spectra of Mn and Cd-containing samples was successfully fitted with a Cd-Mn backscattering path. This backscattering path was constructed by replacing one Mn atom in a pyrochroite structure by a Cd atom.⁵⁴ Correspondingly, the second shell was fitted with a radial distance of 3.33–3.34 \AA and a CN between 1.0 and 1.5. This radial distance is in agreement with the crystallographic distance of Mn in pyrochroite (3.318 Å) .⁵⁴ **Example the model of the transmission of the incompare article in the communite such as** $\frac{1}{2}$ **and the the formed on 4/2 in the formed under** α **Creative of the income of the income of the income of the income of the**

The Mn K-edge P-EXAFS spectra of the oriented sample recorded at $\alpha = 10^{\circ}$, 35°, 55° and 80° are presented in Fig. S24.† The presence of isosbestic points in the spectra provided evidence that differences in the measured spectra were due to orientation effects alone.⁵⁶⁻⁵⁹ The pronounced dependence upon orientation in the regions between the isosbestic points confirmed the successful preparation of a clay film with preferred orientation.⁵⁷⁻⁶⁰ The angle dependence of the spectra also confirmed the anisotropic formation of the secondary Mncontaining solid phases in KGa-1b samples. However, no clear angle dependency was observed in the Cd K-edge P-EXAFS spectra (Fig. S25†).

Conclusion

Modeling metal sorption data showed that both Cd^{2+} and Mn^{2+} had similar affinities for surface complexation on edge sites, suggesting that high Mn^{2+} concentration would lead to less sorption of Cd^{2+} at alkaline pH as indicated by the model prediction for competitive systems. Yet, in competitive experiments containing both Mn^{2+} and Cd^{2+} , Mn^{2+} enhanced Cd^{2+} sorption, demonstrating that Mn^{2+} induced a process that led to a pronounced net increase of Cd^{2+} sorption. Mn K-edge XAS data analysis revealed that Mn^{2+} sorption was partly through the formation of $Mn(n)$ -containing solid phases, although sorption experiments were performed under anoxic conditions and the solution were undersaturated with respect to common Mnphases. Shell fitting of the FT of Cd K-edge EXAFS spectra indicated that Cd was sorbed by forming inner sphere surface complexes or by being incorporated into the formed $Mn(n)$ containing solid phases. This way, the Mn^{2+} addition increased the retention of Cd in the presence of clay minerals. Our findings imply that in anoxic submerged soils, where pH values typically are circumneutral, elevated Mn^{2+} concentration may contribute to increased retention of Cd by adsorption or incorporation into newly formed $Mn(\pi)$ -containing solid phases on clay minerals. Consequently, the dynamics of dissolved Mn^{2+} may play an important role in the cycling of Cd in redox variable environments such as e.g., contaminated paddy fields used for rice production.

Conflicts of interest

The authors declare no competing financial interest.

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