Retention and Release of Copper on Montmorillonite As Affected by the Presence of a Pesticide

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The effect of the cationic pesticide chlordimeform (N'-(4-chloro-2 methylphenyl)-N,N-dimethylmethanimidamide hydrochloride) on adsorption—desorption processes of copper onto the clay mineral montmorillonite has been studied. This effect was evaluated when both adsorbates were adsorbed simultaneously and when the pesticide was previously adsorbed on montmorillonite. For different concentrations of pesticide used, copper adsorption decreases (in relation to pesticide-free treatment) when both adsorbates are adsorbed simultaneously, indicating a competition between both cations for interlamellar possitions. In contrast, copper adsorption increases when the pesticide is previously adsorbed. Copper desorption isotherms show a very high hysteresis. The maximum percentage of copper desorbed is only 7%, being lower when pesticide is present and decreasing with pesticide concentration.

Keywords: Adsorption; desorption; copper; pesticide; chlordim eform; montmorillonite

INTRODUCTION

The behavior of copper in soils is very important from both an agricultural and an environmental point of view. Copper is an essential element for plants and animals. An increase in its levels in soils, however, can inhibit organic matter biodegradation, as well as its subsequent nitrification. Furthermore, some animals are particularly sensitive to copper (Baker, 1990).

The supply of sewage sludge and/or animal manure (especially that of swine) to agricultural land, as well as the use of fungicides and algaecides containing copper, can represent an important source of pollution for this heavy metal.

The metal mobility in soils and its potential hazard depend mainly on the adsorption-desorption processes that can take place on the colloidal components. From the point of view of adsorption, clay minerals are the most important inorganic components in soils (Christensen, 1989). Their importance is due not only to their abundance but also to the high specific area and exchange capacity values. The addition of pesticides and organic fertilizers can affect copper adsorption by the clay fraction for several reasons, e.g., competition with the metal, chelation in solution, etc. (Puls et al., 1991; Madrid and Díaz-Barrientos, 1994; Morillo et al., 1994, 1997).

Although many studies have been published on the adsorption of pesticides and heavy metals independently, little attention has been given to the phenomena which take place when both are present together (Khan et al., 1980; Morillo and Maqueda, 1992; Morillo et al., 1994; Maqueda et al., 1994; Undabeytia et al., 1996; Grabec et al., 1996).

The objective of this study was to characterize copper adsorption—desorption on a clay mineral in the presence and absence of the cationic pesticide chlordimeform. The

Chart 1



effect of the pesticide in Cu adsorption was studied when both adsorbates were adsorbed simultaneously and when the pesticide was previously adsorbed on montmorillonite.

MATERIALS AND METHODS

The clay mineral used was a montmorillonite from Arizona, designed SAZ-1 (Van Olphen and Fripiat, 1979), supplied by the Clay Mineral Society. Its cation exchange capacity (CEC) is 123.5 mequiv/100 g, of which 100.8 mequiv are due to Ca^{2+} , 19.2 mequiv to Mg^{2+} , 2.6 mequiv to Na^+ , and 0.9 mequiv to K^+ .

Technical grade chlordimeform (90% purity), supplied by Schering Agro, S.A., was used. This pesticide is soluble in water (50% by weight) and ionizes completely, giving the chlordimeform cation and chloride anion. The molecular structure of chlordimeform hydrochloride is shown in Chart 1.

The adsorption experiments were done in triplicate in 50mL polypropylene centrifuge tubes, by mixing 0.1 g of clay mineral with 20 mL of 0.01 M NaCl solutions containing various concentrations of chlordimeform and/or Cu(II). The concentrations used were 0.3, 0.5, and 1.0 mM for chlordimeform and 5-30 mg L⁻¹ for copper solutions. The samples were shaken for 24 h at 20 ± 1 °C. The amounts of chlordimeform and Cu(II) adsorbed were calculated as the differences between their concentrations before and after equilibrium. The concentration of chlordimeform was determined spectrophotometrically at 240 nm and Cu(II) by atomic absorption spectrometry (AAS). The inorganic cations Ca²⁺ and Mg²⁺ released during chlordimeform or/and Cu adsorption were determined by AAS.

Successive adsorptions (Cu(II) after chlordimeform) were achieved by mixing 0.1 g of clay mineral with 20 mL of 0.01 M NaCl solutions containing various concentrations of chlor-

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Figure 1. Adsorption isotherm of Cu(II) on montmorillonite.

dimeform. The samples were shaken for 24 h at 20 ± 1 °C. After reaching equilibrium, the suspensions were centrifuged, the supernatant was removed, and the solid samples were washed twice with distilled water and treated again with Cu-(II) solutions under the same conditions as previously.

The adsorption isotherms were obtained by plotting the amount of Cu adsorbed C_s (μ mol g⁻¹) versus the equilibrium concentration C_e (μ mol L⁻¹). In all cases, the pH of the equilibrium solutions from simultaneous and successive adsorptions was 6.5 ± 0.1 .

Desorption experiments were performed after equilibrium adsorption was reached, by removing half of the supernatant after centrifugation, replacing it with 10 mL of 0.01 M NaCl, allowing equilibration for an additional 24-h period, and subsequently continuing as in the adsorption experiment. This process was repeated twice more. The time-dependence (kinetic) study of the adsorption and desorption experiments indicated that the time of 24 h was long enough for reaching the equilibrium. The desorption isotherms were obtained by plotting the amount of Cu that remained adsorbed (μ mol g⁻¹) versus the final concentration (μ mol L⁻¹) for every desorption process.

RESULTS AND DISCUSSION

Adsorption of Copper in the Absence of Chlordimeform. The adsorption isotherm of Cu(II) on montmorillonite is shown in Figure 1. This isotherm corresponds to the L-type according to Giles et al. (1960). The shape of this isotherm indicates that as sites are filled it becomes increasingly more difficult for adsorbate to find a vacant site. The adsorption of metal ions by layer silicates and other adsorbents may occur in different types of surface sites, the higher energy sites being occupied within the lower range of surface coverage (Inskeep and Baham, 1983; García-Miragaya et al., 1986; Madrid et al., 1991). In the case of montmorillonite, this can be expected to occur on variable charge edge sites of high energy and through exchange with the exchangeable cations (Ca and Mg principally) saturating the permanent charge planar sites.

The species concentrations of copper in solution for our system (pH 6.5), estimated by using the equilibrium constants given by Lindsay (1979), are 91.64% Cu²⁺, 5.93% CuOH⁺, and 2.44% CuCl⁺. The species CuCl₂ and CuCl₃⁻ represent a very low proportion (below 0.01%). Although copper adsorption possibly occurs principally through the Cu²⁺ species, the possibility of surface hydrolysis should be taken into account (Harsch and Doner, 1984).

To determine if only Cu^{2+} species was adsorbed on the clay, the amount of Ca and Mg desorbed was plotted versus the amount of Cu adsorbed in microequivalents per gram, a straight line with a slope of 0.75 being obtained. This value differs from the theoretical value of 1 that would be obtained if all the copper were adsorbed as Cu^{2+} by cation exchange. This discrepancy can be attributed in part to surface hydrolysis onto the clay, to adsorption of Cu onto edge positions, and also



Figure 2. Adsorption isotherms of Cu(II) on montmorillonite from chlordimeform-free solutions (\Box) , in comparison with simultaneous (Δ) and successive (O) Cu adsorption, at various chlordimeform concentrations: 0.3 (a), 0.5 (b), and 1.0 (c) mM.

to the adsorption of other than Cu^{2+} species. Stadler and Schindler (1993) modeled Cu adsorption onto montmorillonite, observing that at pH values greater than 5.5 copper adsorption was explained taking into account the reaction of the metal with the aluminol groups, principally by the formation of the AlOCuOH species. Thus it appears that the adsorption of copper occurs principally by adsorption of the Cu^{2+} species onto interlayer positions; and as Cu^{2+} or its hydroxylated form onto edge positions.

Adsorption of Copper in the Presence of Chlor**dimeform**. Figure 2 shows the adsorption isotherm of copper on montmorillonite from chlordimeform-free solution in comparison with copper adsorption in the presence of different pesticide concentrations (0.3, 0.5, 0.5)and 1.0 mM). It is observed that copper adsorption decreases when the pesticide concentration increases, suggesting a competition with the pesticide for the adsorption sites. Only the point for the lowest copper concentration seems to be independent of the pesticide concentration. This can be explained by the fact that at low copper concentration it is adsorbed on high affinity sites on edge positions. The pesticide does not compete for these positions. Pesticide competition at the lower pesticide concentration has less effect on Cu adsorption than on Zn adsorption (Morillo and Maqueda, 1992), indicating a higher preference of this clay mineral for Cu than for Zn.

Copper adsorption by montmorillonite can be described by the Freundlich adsorption equation (r > 0.99, p < 0.01), the linear form of which is

$$\log C_{\rm s} = \log K_{\rm f} + n \log C_{\rm e}$$

where C_s is the amount of adsorbed Cu (μ mol g⁻¹) and

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 Table 1. Freundlich Constants and Correlation

 Coefficients for Cu Adsorption on Montmorillonite in the

 Absence and Presence of Chlordimeform

clordimeform, mM	$K_{ m f}$	п	r
0.0	240	0.49	0.99
0.3	204	0.46	0.99
0.5	174	0.46	0.99
1.0	155	0.43	0.99

 Table 2. Freundlich Constants and Correlation

 Coefficients for Cu Adsorption on Montmorillonite in the

 Absence of Chlordimeform and When the Clay Is

 Previously Treated with the Pesticide

_	0			
	chlordimeform, mM	$K_{ m f}$	п	r
	0.0	240	0.49	0.99
	0.3	276	0.47	0.99
	0.5	260	0.54	0.99
	1.0	246	0.53	0.99

 $C_{\rm e}$ the Cu concentration (mmol L⁻¹) in the equilibrium solution. $K_{\rm f}$ denotes the amount of Cu adsorbed at an equilibrium concentration of 1 mmol L⁻¹. This constant ($K_{\rm f}$) is a measure of the relative adsorption capacity, and n gives an idea of the adsorption intensity.

The values obtained for these two constants are listed in Table 1. The competition of the pesticide on Cu adsorption is reflected by the diminishing K_f values as pesticide concentration increases, from 240 to 155 for the systems in the absence and presence of the higher pesticide concentration.

To determine the effect of the pesticide previously adsorbed on the clay on Cu adsorption, an experiment was performed in which montmorillonite was previously treated with various chlordimeform concentrations. After reaching equilibrium, the solution was removed and the sample was washed twice with distilled water prior to subsequent treatment with Cu. It was observed that Cu adsorption was higher when the clay was previously treated with the pesticide than in the adsorption from pesticide-free solutions or when both adsorbates were added simultaneously. This may be due to the fact that a great part of the pesticide is adsorbed on planar positions in the clay, opening the layers and making easier the subsequent penetration of copper. This behavior is possible only when the amount of chlordimeform adsorbed is much lower than the cationic exchange capacity of the mineral, and so a very small fraction of interlamellar positions are occupied by the pesticide, the other positions being available for copper adsorption.

Table 2 shows the Freundlich parameters for Cu adsorption isotherms when the pesticide is previously adsorbed on the clay. In this case the pesticide had a less marked effect on Cu adsorption, showing little dependence upon the pesticide concentration. The slight differences between K_f values support these observations. As in the case of adsorption of Cd (Undabeytia et al., 1996) the greatest Cu adsorption occurs when the pesticide is previously adsorbed on the clay, in contrast to the behavior described previously when montmorillonite is treated simultaneously with copper and the pesticide.

A useful parameter to measure the adsorbantadsorbate affinity is the distribution coefficient $K_{\rm D}$, which is defined as

$$K_{\rm D} = \frac{[\rm X-clay]}{[\rm X-solution]}$$



Figure 3. Cu(II) K_D values as function of the amount of Cu adsorbed in simultaneous (a) and successive (b) adsorption, at various chlordime form concentrations: 0 (O), 0.3 (D), 0.5 (Δ), and 1.0 (\bullet) mmol L⁻¹.

where the numerator is the sorbate concentration adsorbed on the clay, and the denominator is the sorbate concentration in the equilibrium solution, both expressed in microequivalents per gram.

To calculate K_D for copper it was assumed that this heavy metal is in solution in the form of Cu²⁺, since the proportion of this species is >90%.

Figure 3 shows the K_D values for copper adsorption at various chlordimeform concentrations (0, 0.3, 0.5, and 1 mM). Figure 3a represents the values when the clay mineral is treated simultaneously with the pesticide and Figure 3b when montmorillonite is previously treated with the pesticide. In all cases there are high K_D values for low surface coverage; these decrease when the loading of the metal is higher. The affinity of Cu for the adsorbent decreases until an almost constant value is reached. This could indicate again that in all cases sites of high affinity play an important role at low surface coverage.

In simultaneous adsorption (Figure 3a), K_D values decrease as pesticide concentrations increase due to the higher competition of the pesticide especially for interlamellar positions. On the other hand, when the pesticide is previously adsorbed on montmorillonite (Figure 3b), K_D values are very similar for each pesticide concentration due to the scarce competition between both sorbates, since in successive adsorption of Cu(II) the chlordimeform concentration present in equilibrium solution is very low, because it is only a small amount of that previously adsorbed on the clay which has been desorbed. In all cases the K_D values are always higher than the corresponding values when Cu is adsorbed in the absence of the pesticide. This is in agreement with the results obtained in Figure 2, showing the higher adsorption of Cu on montmorillonite previously treated with chlordimeform, and indicates the higher affinity of Cu for montmorillonite when the pesticide has been previously adsorbed.

Desorption Processes. Figure 4 shows the adsorption isotherms of Cu from chlordimeform-free solutions in comparison with simultaneous and successive Cu adsorption when the concentration of pesticide was 1 mM and the consecutive three-step desorption isotherms.



Figure 4. Adsorption-desorption isotherms of Cu on montmorillonite from chlordimeform-free solutions (a) and in the presence of 1 mM chlordimeform: (b) simultaneous and (c) successive.

In all cases a very high hysteresis is observed; i.e., the results of desorption deviated markedly from those corresponding to the adsorption isotherms. This hysteresis indicates that the adsorption is virtually nonreversible, since the amount of copper that remains adsorbed for each desorption point is practically the same. The desorption cycles involve centrifugation, removal of half of the volume of the supernatant, and addition of a corresponding volume of 10 mL of 0.01 N NaCl, and thus the concentrations of Ca²⁺ and Mg²⁺ cations in suspension are reduced. A small increase in the magnitude of the surface potential (Undabeytia, 1997), due to a smaller sum of total concentrations of divalent cations, contributes to enhanced adsorption of Cu in desorption cycles, as occurs in Cd desorption process on montmorillonite (Undabeytia et al., 1997b).

Figure 5 shows the percentage of copper desorbed versus the initial amount of copper used. The maximum percentage desorbed after three cycles of desorption is only 7% of the Cu previously adsorbed, corresponding to the pesticide-free isotherm, being lower when the pesticide is present and decreasing with pesticide concentration. This indicates that the increasing pesticide concentration in some way impedes the metal desorption. In general, for a given concentration of pesticide the amount of Cu desorbed was smaller for the lower concentration of Cu, possibly due to the fact that when low amounts are adsorbed the copper is adsorbed principally on edge positions, from which it cannot be desorbed by cationic exchange. At higher adsorption densities the interlamellar positions have more importance and desorption from them is easier, since they are not high affinity positions. Consequently the heterogeneity of adsorption sites affects the adsorption process and hysteresis is likely to be associated with the different paths of the adsorption and desorption processes.



Figure 5. Percentages of Cu(II) desorbed versus the initial amounts of Cu used in simultaneous (a) and successive (b) adsorption, at various chlordimeform concentrations: $0 (\bigcirc)$, $0.3 (\Box)$, $0.5 (\bullet)$ and $1.0 (\blacksquare)$ mM.

Cu desorption experiments carried out with the clay mineral previously treated with the pesticide (successive adsorption) show (Figure 5b) that the amount of desorbed metal is a little lower than in the absence of the pesticide, as occurs when copper and pesticide are adsorbed simultaneously. In successive adsorptions the percentages of copper desorbed are very similar and almost independent of the pesticide concentration with which the clay was previously treated, since the cations which are desorbing copper are principally those from the background electrolite, because the chlordimeform concentration in equilibrium solution is very low (it is only a small fraction of pesticide desorbed from the clay).

For the lower metal concentrations (5 and 10 mg L⁻¹), the percentages of Cu desorption are lower in successive adsorptions than in simultaneous adsorption (Figure 5). This may be due to the greater affinity for copper adsorption in successive processes, as was shown for the higher K_D values obtained in successive than in simultaneous adsorption, for lower surface coverage (Figure 3).

The experiments carried out in this work show that, given the mutual influence between pesticides and heavy metals in adsorption—desorption processes, these phenomena must be taken into account if they are to be applied rationally.

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Received for review June 9, 1997. Revised manuscript received December 5, 1997. Accepted December 19, 1997.

JF970500K