Transformation of Pb(II) from Cerrusite to Chloropyromorphite
in the Presence of Hydroxyapatite under Varying Conditions of pH

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Abstract

Cerrusite (PbCO₃) is soluble under acidic conditions and considered to be a highly bioavailable soil Pb species. In this study, synthetic cerrusite and hydroxyapatite [Ca₅(PO₄)₃OH] were reacted under constant and dynamic pH conditions with various P/Pb molar ratios in an attempt to evaluate the effect of reaction kinetics on the formation of chloropyromorphite (Pb₅(PO₄)₃Cl) and solubilization of Pb. Under constant pH conditions, dissolution rates of both cerrusite and apatite were rapid when pH was low. Complete conversion of Pb from cerrusite to chloropyromorphite occurred within 60 min at pH 4 and below when the amount of phosphate in the added apatite was stoichiometrically equal to that needed to transform all added Pb into chloropyromorphite. The concentration of soluble Pb depended upon the solubility of chloropyromorphite. The dissolution rates of apatite and cerrusite decreased with increasing pH, and the transformation was incomplete at pH 5 and above in the 60 min reaction period. The soluble Pb level, therefore, was determined by the solubility of cerrusite. In the dynamic pH system which simulated the gastrointestinal tract (GI tract) system, a complete transformation of Pb from cerrusite to chloropyromorphite was achieved due to the complete dissolution of apatite and cerrusite at the initial low pHs. Chloropyromorphite was the exclusive reaction product in both constant and dynamic pH systems as indicated by XRD analysis. The differences in transformation rate and the control of Pb solubility between the reactions occurring in constant and dynamic pH systems indicate the significance of kinetics in controlling the bioavailability of Pb and the potential for the reaction to occur during ingestion.
Introduction

Cost effectiveness and environmental friendliness have made the addition of phosphate minerals to Pb-contaminated soil a promising technology to remediate Pb-contaminated soils. The recent research results have demonstrated that the reduction in soluble Pb levels of soil solutions by mixing phosphate minerals with Pb contaminated soils (1-9), the formation of pyromorphite in aqueous Pb solution, on exchange resin surface and from geothite adsorbed Pb by reaction with hydroxyapatite (2, 3, 8, 10). The existence of pyromorphite in the contaminated soils, road-side soils and mine-wastes has been documented (11, 12). Furthermore, pyromorphite formation in contaminated soils amended with phosphate and hydroxyapatite (9, 13, 14) have also been reported.

It is generally agreed that the risks of soil Pb movement into groundwater aquifers (mobility) and being absorbed by plants (phytoavailability), animals and/or human beings (bioavailability) are associated with the Pb species rather than the total Pb content in the soil. The reported soil Pb bioaccessibility results were consistent with Pb speciation results. For example, a greater fraction of bioaccessible Pb was obtained in the soil containing more soluble Pb phases such as lead oxide and cerussite than the soil containing galena, anglesite, lead phosphate and encapsulated Pb phases (15, 16). Solubility, therefore, is used in reference to the stability of a specific Pb species. However, solubility is a thermodynamic parameter and only defined when the reaction reaches equilibrium. In a dynamic system like soil, the kinetics of a reaction need to be considered in order to determine the possibility and the extent of a reaction occurring under a given condition and time scale. For example, in a kinetic study, the significant difference in the amount of Pb leached from soil and waste by acid was attributed to differences in dissolution rates of the two types of Pb solid phases (17). Therefore, in addition to the solubility, the
dissolution rates of a solid Pb-bearing form and the possible reaction products should be considered when evaluating the stability of the Pb form reacting in a dynamic system.

An accepted paradigm is that the free metal species, e.g., the bare metal cation (M\textsuperscript{n+}), is the most bioactive and the most toxic. The formation of ion-pairs, complex ions, polymers and microparticulates (e.g., colloidal iron oxide or humic substances) of a bare metal cation will reduce its potential for uptake by living organisms and toxicity (18). The free Pb ion concentration should be the critical parameter to monitor in soil solution or gastrointestinal fluid when evaluating the bioavailability of soil Pb species. Thermodynamically, in the aqueous suspension of a group of Pb minerals the one with the least solubility will determine the soluble Pb level as long as there is a source of ligand to precipitate the soluble Pb as the least soluble Pb mineral. Phosphate minerals, when they are added to Pb-contaminated soils, establish a phosphate source that can maintain the soluble phosphate at a level sufficient to precipitate the dissolved Pb as pyromorphite. Thus, the soluble Pb level will be controlled by the least soluble pyromorphite instead of the original soil Pb species. However, even with sufficient phosphate mineral to stoichiometrically precipitate the total soil Pb into pyromorphite, the soluble Pb level may not necessarily be controlled by pyromorphite if the dissolution rate of the added phosphate mineral is not sufficient to deliver the soluble phosphate needed to precipitate the dissolved Pb. Thus, kinetic concerns may become more important than thermodynamics in a dynamic system like soil. This kinetic concern becomes critical if the added phosphate minerals and soil Pb are simultaneously placed into a living organism where the reactions are assumed to take place, such as in a gastrointestinal system.

The dominant mechanism of pyromorphite formation in the Pb-P-H\textsubscript{2}O system has been reported as precipitation of soluble lead and phosphate (1, 8, 19), although substitution of Pb for
Ca in apatite may be possible (12). It has been found that formation of pyromorphite from soluble phosphate and Pb is a rapid process; soluble Pb can be depleted from solution within a few seconds if a stoichiometric amount of phosphate is provided and the system is well mixed as observed in our laboratory. Thus the soluble Pb concentration in the aqueous system containing apatite and Pb-bearing solids will be determined by the dissolution rate of apatite and the overall Pb release rate (desorption, dissociation and dissolution) of the Pb species. Therefore, knowledge of reaction kinetics in the Pb-bearing solids and phosphate mineral provides additional information needed in the evaluation of the technology of immobilization of soil Pb and offers the basis to improve the technology.

To demonstrate this, a series of kinetic experiments were conducted in systems containing hydroxyapatite and cerrusite to determine the effects of dissolution rates of apatite and cerrusite on: 1) the soluble Pb level during the reactions; and 2) formation of pyromorphite. Cerrusite is a secondary Pb mineral commonly found in soils (20, 21). It is also a weathering product of other secondary Pb minerals such as anglesite (PbSO₄) under appropriate condition of CO₂ partial pressure or activity (22). Cerrusite is not stable and dissolves under acid pH conditions, e.g., acidified rain (23). It has been demonstrated that the bioavailability of Pb carbonate was higher than other soil Pb species such as anglesite and Fe-Mn associated Pb. Further, the Pb content in the plants grown on the Pb contaminated soils was primarily associated with soil Pb carbonate content rather than the total soil Pb concentrations (24, 25). Thus, cerrusite can be an important mineral form of Pb in soil in controlling the soil Pb mobility and reactivity, it also can be a highly bioavailable form of soil Pb.

Materials and Methods
Materials Analytical reagent lead carbonate (PbCO₃) used in this study was from Fisher Scientific. Although an amorphous morphology was observed under a scanning electronic microscope, the powder X-ray diffraction analysis illustrated a characteristic pattern of crystalline cerrusite mineral. The mineral had a specific surface area of 2.61 m² g⁻¹ measured by BET nitrogen absorption. The synthetic hydroxyapatite [Ca₅(PO₄)₃OH] used was obtained from Bio-Rad and had a specific surface area of 67.3 m² g⁻¹ and the P/Ca molar ratio was 1.62 which is comparable to the ideal ratio of 1.67 of hydroxyapatite.

Dissolution of cerrusite and apatite Determination of dissolution rates for cerrusite and hydroxyapatite were conducted in a 1.1 L glass beaker placed in a water bath at 37°C, analogous to that of human body. A 1.0 L solution of 0.001 M NaCl and 0.1 M NaNO₃ was measured into the beaker and adjusted to the designed pH with 0.1 M HNO₃ or 0.1 M NaOH. After cerrusite or apatite was added, the solution pH was maintained by an autotitrator (Mettler, DL70ES). The suspension was continuously stirred with a plastic paddle and sampled after 1, 3, 5, 10, 15, 25, 40, 60 and 120 min of reaction. Samples were immediately filtered through a 0.2 µm filter and the filtrate was acidified and refrigerated until analyzed. The concentrations of Pb, Ca and PO₄ in the solution as a function of time were used to determine the dissolution rate for cerrusite and hydroxyapatite.

Reactions between cerrusite and apatite The effects of reaction kinetics of cerrusite and apatite on transformation of Pb from cerrusite into pyromorphite, and on the soluble Pb levels during the reactions, were determined in the systems in which the pH was maintained constant or the pH was adjusted with reaction time. The later was to simulate a gastric situation in terms of pH range and time scale.
The average pediatric gastric pH was reported to be 1.7-1.8 in a fasting stomach (16). Following ingestion of food, the pH rises to 4-6 and then returns to basal values within 2 hours. A child's stomach empties in about 60 min (16). About 20 min after feeding, stomach emptying starts and the chyme enters the small intestine where the pH values reach 7 and Pb absorption occurs. Accordingly, a pH range of 2 to 7 was selected to simulate the GI tract pH values. In addition to the measurements conducted in constant pH suspensions (pH-Constant study), a reaction was carried out in the suspension in which the pH was adjusted vs. time from 2 to 7 within 20 min (pH-Dynamic study). The pH-Dynamic study allows determination of the potential transformation rate of Pb from cerrusite to pyromorphite under simulated GI tract pH conditions assuming cerrusite and apatite were ingested simultaneously.

**pH-Constant study** To measure the reactions between cerrusite and hydroxyapatite, 1.0 g of cerrusite was mixed with apatite at three elemental ratios of P/Pb (3/5, 6/5 and 9/5) providing 1, 2 and 3 times the stoichiometric P/Pb ratio of chloropyromorphite, [Pb₅(PO₄)₃Cl]. The cerrusite and apatite mixture was added into 1.0 L of 0.001 M NaCl and 0.1 M NaN₃ solution with a previously adjusted pH. The pH of the suspension was maintained by the autotitrator and the suspension was sampled periodically while the system was continuously stirred. Sampled suspension was filtered through a 0.2 μm membrane and the filtrate was analyzed for soluble PO₄, Pb and Ca. At the end of a 60 min reaction, the suspension was filtered with a 0.45 μm membrane and the solid collected and dried for XRD analysis. From each of the P/Pb ratios, the reaction at pH 6 was allowed to continue for 48 h before the solid was collected to observe the reaction over a relatively long period.

**pH-Dynamic study** In a separate experiment, cerrusite was reacted with apatite (0.40 g apatite/1.00 g cerrusite, P/Pb = 3.2/5) while the solution pH was adjusted in increments of one
pH unit from 2 to 7 over a 20 min time interval and then maintained at 7 until the end of the 60 min reaction. The suspension was sampled twice at each pH of 2, 3, 4, 5, 6 and 7. One sample was collected 1 min after the pH was established and the second sample was taken before the pH was adjusted to next level. Sample suspension was filtered through a 0.2 μm membrane and the soluble species concentrations of PO₄, Ca and Pb were determined to assess mineral dissolution rate and Pb bioaccessibility during the reaction as well as the transformation rate of Pb from cerrusite into pyromorphite. A separate experiment, with the same procedure, was conducted with 1.00 g of cerrusite but without addition of hydroxyapatite to serve as experiment control.

**Analytical Procedures** An inductively coupled plasma spectrophotometer (ICP-AES, Trace 61 E, Thermal Jerrel Ash) was employed to analyze PO₄, Ca, and Pb concentrations. The detection limit for these 3 elements was 2 μg L⁻¹. The US EPA Water Supply Performance Evaluation Study solution (WS033) was used to calibrate and verify the analytical standard solutions used for ICP analysis. Experimental blanks, standards and spiked samples were used for analytical quality control. Solution and suspension pH was measured using a glass pH electrode which was connected to the Mettler titrator. With this set-up, the variation of pH could be maintained within 0.01 pH unit.

A X-ray diffractionmeter (Scintag, XDS 2000) with Cu K-α radiation at 35 KV and 30 mA was used. Solids used for XRD analysis were air dried, crushed and mounted on a glass sample holder. XRD patterns were collected using a step-scanning rate of 0.04° 2θ/sec.

To quantify the mineral composition of the reacted solid collected at the end of reaction, a portion (0.010 g) of dried solid was dissolved in 100.0 ml of a 10% HNO₃ and analyzed for PO₄, Pb and Ca contents using ICP. As no Ca mineral other than the added apatite was found in the
solid by XRD, the Ca content in the solid was used to compute the amount of undissolved hydroxyapatite. The amount of PO₄ in the solids was attributed to both apatite and newly formed chloropyromorphite. Lead in the solid was attributed to both cerrusite and chloropyromorphite. From a mass balance analysis, the percentage of cerrusite-Pb transformed into pyromorphite-Pb was estimated. The XRD analysis results were used to evaluate and confirm the assumption of only the three minerals; cerrusite, chloropyromorphite and apatite, in the collected solids.

Results

Dissolution of Cerrusite and Hydroxyapatite The dissolution of cerrusite expressed as the function of soluble Pb concentration vs. time at each of the 5 pHs illustrates that the dissolution process is pH dependent (Figure 1). Dissolution did not reach a steady state during the 120 min reaction period. Because of experimental difficulty in maintaining a stable pH 2 in the presence of cerrusite, no measurements were made at pH 2.

There was no attempt in this study to develop a comprehensive dissolution rate equation for cerrusite because of the difficulty in accurately determining the activities of dissolved carbonate ions. Redistribution of carbonate species, i.e., CO₃²⁻, HCO₃⁻, H₂CO₃ and CO₂(aq), occur when CO₃²⁻ is released into solution from cerrusite dissolution and may take minutes to reach steady state (26). Therefore, in a kinetic study using a time scale of minutes as the sampling interval, the activity of individual species of the carbonate species can not be easily determined.

Dissolution of hydroxyapatite in the pH range of 4 to 7 as a function of time is presented in Figure 2. The dissolution was inversely related to pH. Below pH 4, the dissolution was too rapid to be accurately determined under these experimental conditions. At higher pHs, e.g., pH 6 and 7, the dissolved Ca and PO₄ reached steady state and did not change after 60 min reaction. From these measurements, approximately 1 and 4% of the total added apatite was dissolved at
pH 7 and 6, respectively, within 60 min. In comparison approximately 20 and 40% of the added apatite was dissolved at pH 5 and 4, respectively, after only 25 min. The soluble species compositions at pH 4, 5 and even 6 were stoichiometric to the elemental composition of apatite with a P/Ca molar ratio of 3/5 (Figure 3). However, the PO₄ and Ca concentrations at pH 7 were essentially the same. This may be attributed to adsorption of Ca²⁺ because the high pH and high total surface area of the undissolved apatite are favorable for removal of the cation from solution.

**Reaction of Cerrusite with Hydroxyapatite** The precipitation of pyromorphite in a saturated solution is a rapid process and is completed within seconds. Therefore, the overall reaction in the cerrusite and apatite system should be kinetically controlled by the dissolution rates of cerrusite or apatite:

\[ 5\text{PbCO}_3(s) + \text{Ca}_3\text{(PO}_4\text{)}_3\text{OH}_2(s) + \text{Cl}^{-}(aq) = \text{Pb}_5(\text{PO}_4)_3\text{Cl}_3(s) + 5\text{Ca}^{2+}(aq) + 5\text{CO}_3^{2-}(aq) + \text{OH}^{-}(aq) \]  

(1)

From Eq. (1), the soluble Ca can be used as the indicator for the dissolution of apatite because under these conditions there was no reaction to remove the dissolved Ca from the aqueous phase. When the amount of apatite dissolved and the soluble PO₄ concentration is determined, the formation of chloropyromorphite can be estimated from the PO₄ mass balance assuming chloropyromorphite formation is the only processes to remove the soluble PO₄. From the results of the XRD analysis of this study as well as previous studies (10), the formation of chloropyromorphite is the only process in the system which removes soluble Pb and PO₄. It is also apparent from Eq.(1) that the concentrations of soluble Pb or PO₄ were determined by either the originally added minerals, e.g., apatite and cerrusite, or the newly formed mineral during the transformation process. If the relative rate of dissolution of apatite is lower than that of dissolution of cerrusite, the soluble Pb will be dependent upon the solubility of cerrusite \([-\log K_{sp} = 13.11, (27)]\). As stated before, the soluble Pb level is of interest for its high reactivity and
bioavailability. As both cerrusite and apatite dissolution are pH dependent, the reaction rate of cerrusite and hydroxyapatite should be directly associated with pH.

**pH-Constant Study** In the experiment with a P/Pb ratio of 3/5, 2.3 x 10^{-3} mol of PO_4 as hydroxyapatite, together with 1.00 g of cerrusite (3.74 x 10^{-3} mol Pb), was placed into a solution and reacted for 60 min under constant pH conditions. The amount of PO_4 added was stoichiometrically equal to that needed to transform the added cerrusite-Pb into chloropyromorphite.

The soluble Ca concentration indicates that nearly complete dissolution of added hydroxyapatite was achieved at pH 2 and 3 within 5 min (Figure 3a). At pH 4, about 70% of the apatite was dissolved in 15 min and 90% of added apatite was dissolved at 60 min. However, apatite dissolution was much lower when the pH value in the solution was above 4, only 55, 15 and 10% of the added apatite was dissolved in the 60 min reaction at pH 5, 6 and 7, respectively (Figure 3a).

The formation rate of chloropyromorphite in the suspensions was proportional to the dissolution of apatite. The XRD patterns indicate that at pH 2 and 3 the only mineral in the solid collected after 60 min reaction was chloropyromorphite (Figure 4). At pH 4, a finger print of cerrusite was observed in the XRD pattern, but the dominant mineral was chloropyromorphite. The solid collected at pH 5 after 60 min was a mixture of cerrusite and newly precipitated chloropyromorphite as identified by XRD (Figure 4). The undissolved cerrusite was the dominant mineral in the solid of pH 7, but characteristic peaks of chloropyromorphite in the XRD patterns were also observed. Thus, the added cerrusite was partially transformed into chloropyromorphite at pH 7 (Figure 4).
To quantify the transformation of cerrusite-Pb into chloropyromorphite-Pb, the total PO₄, Ca and Pb contents of the collected solids were determined and the mineral composition in these solids were calculated. At pH 2, 3 and 4, there was essentially no cerrusite in the solids (Figure 5a). At pH 5 and above, the cerrusite was a significant constituent of the solid (Figure 5a). These results were consistent with the XRD analysis which illustrates that the transformation of cerrusite-Pb to chloropyromorphite decreased with increasing pH (Figure 4 and 5a). In terms of the transformation rate, as illustrated in Figure 5b, nearly 100% conversion of cerrusite-Pb to chloropyromorphite-Pb was achieved at pH 2, 3 and 4 regardless of the P/Pb ratio because of the rapid dissolution of cerrusite and apatite at low pH conditions. In contrast, when the solution pH was greater than 5, the transformation rate was increased with increasing P/Pb ratio (Figure 5b). This may be attributed to the rate of apatite dissolution which is proportional to the total surface area of solid apatite in the suspension.

The estimated soluble Pb levels from the solubility of chloropyromorphite \([-\log K_{sp} = 84.4, (28)]\), \(10^{-3.50}\) and \(10^{-4.34}\) M, at pH 2 and 3, respectively, were in agreement with the measured soluble Pb concentrations, \(10^{-3.1}\) and \(10^{-4.15}\) M (Figure 3b), implying that chloropyromorphite was controlling Pb concentration in solution. This indicates that only chloropyromorphite-Pb was present and is in agreement with the XRD analysis of the solids (Figure 4).

Considering the soluble PO₄ concentration of \(10^{-5.28}\) M at pH 4 the concentration of soluble Pb should be \(10^{-4.8}\) M if it was controlled by dissolution of chloropyromorphite. However, the measured soluble Pb at pH 4 was \(10^{-3.62}\) M (Figure 3b) which was one order of magnitude higher than that estimated from the solubility of chloropyromorphite, but about one order of magnitude lower than that of the suspension containing only cerrusite (Figure 1). This
observation implies that the trace amount of cerrusite that remained in the suspension of pH 4 still played a role in determining the soluble Pb concentration even though nearly all of the added Pb had been converted into chloropyromorphite. At pH 5, the soluble Pb was $10^{-3.04}$ M (Figure 3b) which was comparable with the concentration obtained from the suspension containing only cerrusite, $10^{-2.95}$ M (Figure 1). As illustrated in Figure 4, approximately 55% of the added cerrusite-Pb was transformed into chloropyromorphite at pH 5, the soluble Pb level was determined by cerrusite, implying there was insufficient PO$_4$ ($10^{-6}$ M) to react with Pb to form chloropyromorphite. From Figure 1 and 2, it can be seen that decrease in dissolution rate of cerrusite from pH 4 to 5 was not as significant as that of the decrease in apatite dissolution rate. This indicates that the relative dissolution rate of apatite is lower than that of cerrusite at pH 5 and thus the supply of soluble PO$_4$ is insufficient to precipitate dissolved Pb at this pH. The soluble Pb concentrations at pH 6 and 7 were $10^{-5}$ and $10^{-5.5}$ M, at 60 min, respectively, about one order of magnitude lower than that without apatite, $10^{-3.6}$ and $10^{-4.5}$ M for pH 6 and 7 (Figure 1). This reduction in the soluble Pb concentration at these pHs could be attributed to the removal of dissolved Pb by the ongoing reactions, including dissolution of apatite and cerrusite and precipitation of chloropyromorphite. Because the dissolution rate of cerrusite at pH 6 and above decreased dramatically (Figure 1), the rate of PO$_4$ release from apatite dissolution maintained a soluble PO$_4$ level of $10^{-6}$ M (Figure 3c), which provided sufficient phosphate to precipitate the dissolved Pb at this pH level. Thus, the concentrations of soluble Pb were determined by both cerrusite and chloropyromorphite in the suspensions of pH 6 and 7.

Although not shown, similar phenomena were observed in the reactions with P/Pb ratios of 6/5 and 9/5. The amount of chloropyromorphite formed in the solid after 60 min reaction decreased with an increase in pH. The total transformation of Pb from cerrusite to
chloropyromorphite for the three P/Pb ratios after 60 min under constant pH illustrates that complete transformation is obtained at pH 2, 3 and 4 (Figure 5b). At pH 5 and above, transformation decreased with increasing pH. Increasing the P/Pb ratio caused an increase in the percent of Pb transformed into chloropyromorphite at pH 5 and 7 (Figure 5b).

**Effect of Reaction Time** To test the effect of reaction time on the transformation of Pb from cerrusite to chloropyromorphite, the reactions at pH 6 for each of the three P/Pb ratio experiments were allowed to continue for 48 hours instead of 60 min as reported above. The mineral composition of the solids collected after 48 h demonstrated that 83(±4), 94(±4) and 96(±6)% of cerrusite Pb was converted into chloropyromorphite when the P/Pb molar ratios were 3/5, 6/5 and 9/5, respectively. If we assume that the transformation rate at pH 6 should be between those obtained at pH 5 and pH 7 for a 60 min reaction time (Figure 5b), then the higher transformation rate may be attributed to the longer time of reaction. The XRD patterns for the solids collected for the three P/Pb ratio after 48 h reaction indicate some of the added cerrusite still remained at P/Pb ratio of 3/5 and 6/5, but it was not shown in the solids collected when P/Pb ratio was 9/5 (Figure 6). Furthermore, the XRD patterns in Figure 6 also demonstrated that chloropyromorphite mineral was the only reaction product.

**pH-Dynamic Study** As demonstrated by the soluble Ca concentration (Figure 7), under the condition of variable pH system (initial pH of 2.1 - 2.4), the added apatite was completely dissolved during the initial 5 min reaction time. The soluble Ca concentration of 3.4 x 10^{-3} M was the result of complete dissolution of the added apatite (0.38 g) that contained 3.78 x 10^{-3} moles of Ca. However, the dissolved PO_4 concentration was 3.5 x 10^{-4} M, about 15% of the total PO_4 from apatite dissolution. The soluble Pb concentration shown in Figure 7 was 2.4 x 10^{-4} M, about 13% of the Pb added as cerrusite. Considering full dissolution of both apatite and cerrusite at
these low pH values and the nearly stoichiometric removal of Pb and PO₄, it can be surmised that chloropyromorphite formation was occurring. This formation of chloropyromorphite was confirmed by the XRD analysis of the solid collected at the end of the experiment (not shown). The transformation of Pb from cerrusite to chloropyromorphite as a function of time was calculated based on the mass balance of PO₄, Ca and Pb in the system. In the pH-Dynamic system, more than 95% of added Pb was converted from cerrusite into chloropyromorphite within 10 min and remained in the suspension regardless of the reaction time and increase in pH.

The soluble PO₄ level was maintained at 1.0 - 2.7 x 10⁻⁴ M after the first 5 min reaction, and this relatively high soluble PO₄ level resulted in a low soluble Pb concentration during the reaction (Figure 7). The soluble Pb level was 10⁻⁵ M after 7 min reaction at pH 2.8. This is compatible with that obtained in the pH-Constant study (Figure 3b). When the pH was raised to 4.1, after 10 min reaction, the soluble Pb concentration was 10⁻⁵.⁵⁷ M which is about two orders of magnitude lower than that from the pH-Constant study and agreed with the value, 10⁻⁵.⁵ M, calculated from solubility product of chloropyromorphite. The soluble Pb concentration was further decreased to 10⁻⁶.²⁷ M at pH 5, compared to 10⁻³.⁰⁴ M in the pH-Constant study. When the suspension pH was raised to 6 and 7, the soluble Pb ranged from 10⁻⁷.²³ - 10⁻⁷.⁷ M, or 10 - 4 µg Pb L⁻¹, which is consistent with the solubility of chloropyromorphite and more than two orders of magnitude lower than that of the pH-Constant study (Figure 3b). Thus, in the pH-Dynamic system the solubility of chloropyromorphite controlled the soluble Pb level during the entire reaction process.

Unlike the pH-Constant system in which the transformation of Pb from cerrusite to chloropyromorphite was incomplete for the rapid decrease in apatite dissolution rate at pH 4 and above (Figure 2, 4 and 5), in the pH-Dynamic suspension the cerrusite-Pb was rapidly and fully
converted into chloropyromorphite as long as the amount of phosphate as apatite was stoichiometrically equal to that needed to transform the added Pb in cerrusite into chloropyromorphite.

In the constant pH system, both the undissolved cerrusite and newly formed chloropyromorphite controlled the soluble Pb levels at pH 4 and above. However, in the pH-Dynamic system, the soluble Pb level was determined by chloropyromorphite during the entire reaction period. The complete dissolution of apatite and cerrusite in the initial acidic condition resulted in a high soluble PO$_4$ level which precipitated the soluble Pb from the cerrusite dissolution. This resulted in a rapid precipitation of chloropyromorphite as well as a dramatic reduction of soluble Pb level during the reaction. At pH 4, 5, 6 and 7, the soluble Pb concentrations were 2.4 x 10$^{-4}$, 9.5 x 10$^{-4}$, 1.0 x 10$^{-5}$ and 3.1 x 10$^{-6}$ M in the pH-Constant suspensions, as compared to 2.7 x 10$^{-6}$, 5.3 x 10$^{-7}$, 5.0 x 10$^{-8}$, and 1.9 x 10$^{-8}$ M in the pH-Dynamic system. Thus, the soluble Pb was at 10 µg L$^{-1}$ or lower when pH 6 was reached, which took about 16 min in the pH-Dynamic system.

These data demonstrated that the formation of chloropyromorphite from cerrusite and apatite can take place in the pH conditions of the aqueous system of the GI tract. Apparently, under the pH condition of the GI tract, the formation of chloropyromorphite is kinetically favored. Furthermore, the results from constant pH systems can not predict the Pb-PO$_4$ reaction products obtained in a dynamic pH system because of the reaction kinetics.

In the case where cerrusite is ingested with contaminated soil in the presence of stoichiometric amounts of phosphate(s), it is logical to assume that the soluble Pb concentration will be controlled by the solubility of chloropyromorphite, which is lower than 10 µg L$^{-1}$ when
pH 7 is reached. Thus, one can predict that the bioavailability of Pb from ingested cerrusite would be significantly reduced in the presence of apatite.

Literature Cited

(9) Zhang, P.-C.; Ryan, J. A., Soil Science Society of America Annual Meeting, St. Louis, MO, 1995; ASA SSSA; 41.
FIGURE CAPTIONS

Figure 1. Effect of pH on dissolution rate of cerussite as a function of time; Initial Cerussite = 1.00 g, NaNO₃ = 0.100 M and NaCl = 0.001 M.

Figure 2. Dissolution of hydroxyapatite as a function of time and pH. The filled and unfilled symbols represent the phosphate and calcium concentrations, respectively.

Figure 3. Soluble Ca (a), Pb (b) and PO₄ (c) concentrations in the cerussite-apatite suspensions at various pHs. The P/Pb ratio = 3/5 and I = 0.100 M.

Figure 4. Effect of pH on the transformation of Pb from cerussite to chloropyromorphite, Reaction time = 60 min, P/Pb ratio = 3/5.

Figure 5. Mineral composition of the solid collected after 60 min reaction (a) and the transformation of cerussite-Pb to chloropyromorphite-Pb (b).

Figure 6. Effect of P/Pb ratios on the transformation of Pb from cerussite to chloropyromorphite, Reaction time = 48 hours, pH = 6.

Figure 7. Concentrations of soluble species in cerussite-apatite suspension as a function of pH and time. The suspension pH was adjusted from 2 to 7 within 20 min and the total reaction time was 60 min.
CI-pyromorphite

$\frac{P}{Pb} = 3/5$

$\frac{P}{Pb} = 6/5$

$\frac{P}{Pb} = 9/5$

$2^\circ \theta$