MEASURING SURFACE CHEMICAL PROPERTIES OF SOIL USING FLOW CALORIMETRY¹

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Flow calorimetry, which is ideally suited for measuring reactions occurring at the liquid/solid interface, has been used to study the surface chemistry of many types of solids, but little use of it has been made in the study of surface reactions of soils. The purpose of this study was to demonstrate the application of flow calorimetry to the study of two fundamental soil chemical processes, namely cation exchange and phosphate sorption. Surface horizon samples of a Typic Acrorthox and a Typic Tropohumult from Puerto Rico, a strong acid cation exchange resin (Dowex 50W-8), and an amorphous Al(OH)₃ were used. Heats for K/Ca exchange on the Dowex resin and the Oxisol, and K/Na exchange on the Ultisol, were consistent with literature values that were obtained using conventional batch calorimetry or derived from the temperature dependence of the exchange constant. Although peak areas associated with a given pair of exchange reactions were equal, peak shapes were generally not equivalent, indicating differences in the rate at which the two reactions occurred. For example, Ca displacing exchangeable K occurred more rapidly than the reverse reaction on the Dowex resin. The reaction of phosphate with the Ultisol and amorphous Al(OH)3 was exothermic. Exposure of the soil to several cycles of phosphate was sufficient to saturate the sorption sites, as evidenced by the loss of a detectable heat signal. However, phosphate reactive sites were regenerated by flushing the column with a salt solution at pH 10. Precipitation of Al-phosphate was shown to be endothermic, confirming that precipitation was not the primary mechanism for phosphate sorption in this study. The results of this study show that flow calorimetry can provide valuable information about surface chemical reactions in soils that cannot be obtained readily by other methods. (Soil Science 2002;167:782-790)

Key words: Phosphate sorption, ion-exchange, heat of adsorption.

CALORIMETRY provides a direct, quantitative measure of the heat involved in a reaction. This measured heat is related to a change in en thalpy, a fundamental property of the system that can provide information about the chemical processes taking place.

Flow calorimetry is ideally suited for mea

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suring interactions occurring at the liquid/solid interface. A typical flow calorimeter consists of a column holding 50 to 100 mg of a solid through which a carrier liquid is passed. A chemical or physical interaction between the solid and a component added to the carrier liquid results in a temperature change that can be related quanti tatively to the heat of the reaction. Flow calo rimetry has the following advantages over con ventional batch calorimetry (Steinberg, 1981): i) flow calorimetry can resolve a complex series of reactions that occur more or less simultaneously but at different rates; ii) multiple adsorption/ desorption cycles can be applied to the same sample, allowing reversible and irreversible pro cesses to be distinguished; iii) changes that occur in the surface properties of the solid as a result of specific treatments or aging effects can be quanti fied; and iv) when both the amount of adsorption and its associated heat are measured, information about surface heterogeneity can be obtained.

Flow calorimetry has been used to study the surface chemistry of many types of solids (Can ham and Groszek, 1992; Steinberg, 1981; Schnei der et al., 1997; Groszek and Partyka, 1993; Noll, 1987; Meziani et al., 1997; Taraba, 1990), but lit tle use has been made of it in the study of surface reactions of soils. Questions abound about some of the basic chemical processes that are involved in soils. While sorption isotherms have tradition ally been used to obtain information about the rate and amount of solute sorbed over time, they give no insight into the mechanism by which sorption occurs (Sposito, 1984). Although en thalpies of adsorption on soil have been derived from temperature dependence of the sorption isotherm, in some instances the conclusions have been contradictory. The purpose of this paper is to summarize briefly the literature for two fun damental soil chemical processes, ion exchange and phosphate sorption, and to demonstrate the application of flow calorimetry to the study of these two processes in soils.

METHODS AND MATERIALS

Instrumentation

Several inexpensive flow calorimeters for measuring heats of adsorption from solution onto solids were constructed in our lab. Approximately 50 mg of soil were placed inside a small column (Fig. 1), and solutions containing reactive species were forced through the column using a total pressure drop of about 100 cm of water. Flow rates were controlled with a precision needle valve at

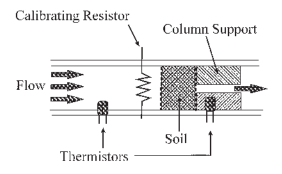


Fig. 1. Schematic of column, thermistors, and calibrating resistor used in the flow calorimeter.

the outlet side of the calorimeter and were gener ally constant to within a few μL min⁻¹ during a day's run. Typical flow rates were in the range of 0.25 to 0.35 mL min⁻¹. Assuming that water saturated soil requires a gravimetric water content of about 0.20, 50 mg of soil would have a pore volume (PV) equal to about 0.01 mL. Thus, typi cal flow rates for this study corresponded to about 25 to 35 PVs min⁻¹. Run times varied between 20 min and 1 h, depending on the time required for the signal to return to the baseline. Thus, en dothermic and exothermic peaks corresponded to between 500 and 2000 PVs.

A pair of thermistors, one upstream and the other downstream from the soil column, formed one half of an electronic bridge and sensed tem perature changes in the solution as it passed through the column. A change in solution tem perature produced a differential output voltage from the bridge. This differential voltage was fed into an instrumentation amplifier, and the amplified signal was fed into a computer for process ing. The system possessed high sensitivity, low thermal drift, and a good signal to noise ratio.

Peak areas were obtained by integrating the signal (volts) numerically over time. This time averaged peak area (V min) was converted to a flow rate averaged peak area (V mL) by multiply ing by the average flow rate. This was measured for each peak by collecting the effluent volume and dividing by the time over which the volume was collected. Peak areas were converted to en ergy units (Joules) by comparison with peaks generated with a calibrating resistor located within the flow stream and immediately up stream from the column (Fig. 1). Voltage and cur rent for the heat pulses were measured, and the heat input was then calculated from the relation $Q ext{ (Joules)} = V A t$, where $V ext{ is voltage, } A ext{ is am}$ perage, and t is time, in seconds, that the resistor was energized.

The column assembly with thermistors and calibrating resistor was sealed inside a 500 mL polyethylene bottle, and the bottle was placed in a 50 L insulated container, filled with water, at room temperature. The water bath provided good thermal stability against ambient tempera ture changes and generally resulted in baselines with negligible drift.

Soils

The 0 to 15 cm depths of an Oxisol and an Ultisol from Puerto Rico were air dried and sieved to an aggregate size between 0.25 and 0.50 mm before being used in this study. The Oxisol

TABLE 1

Mineralogical and chemical properties of the two Puerto Rican soils

Soil	Organic matter		CEC	Total Fe + Al	Clay	Texture ²	Surface area
	$\rm g~kg^{-1}$	pН	$\operatorname{cmol}_{\operatorname{c}} \operatorname{kg}^{-1}$	$\rm g~kg^{-1}$	mineralogy ¹	$\rm g~kg^{-1}$	$(m^2 g^{-1})$
Oxisol	40.5	4.9	3.1	287	k>go>gi=q	100 340 560	41.9
Ultisol	18.6	4.7	11.0	86	k>q>s>go>m	120 290 590	37.8

¹k - kaolinite, gi - gibbsite, go - goethite, q - quartz, s - smectite, m - mica ²Sand-Silt-Clay

(clayey, oxidic, isohyperthermic Typic Acror thox) was found near Mayaguez, and the Ultisol (clayey, mixed, isohyperthermic Typic Tropohu mult) was taken from the central mountainous area near Corozal. Mineralogical and chemical properties of these soils were obtained using stan dard methods and are presented in Table 1 (Appel and Ma, 2002).

Cation Exchange Resin

A 50 to 100 mesh Dowex 50W 8 strong acid cation exchange resin was also used. It was initially in the $\rm H^+$ form and had an exchange ca pacity of 5.0 mmol_c $\rm g^{-1}$ dry weight.

$Al(OH)_3$

This material was obtained by titrating a so lution of $AlCl_3$ with NaOH to pH 6 and allow ing the suspension to stand overnight. The solid was separated from the solution phase by cen trifugation and dried in an oven overnight at 70 °C. X ray diffraction of the dried residue showed it to be noncrystalline.

RESULTS AND DISCUSSION

Sensitivity and Precision of Heat Measurements

A typical peak to peak noise level for the flow calorimeters used in this study was 5 mV or less. The thermistors had a nominal resistance of 10 k Ω at 25 °C and a temperature coefficient of resistence of about -400Ω °C⁻¹. Assuming an acceptable signal to noise ratio for peak detection to be about 5, and using the measured gain for the instrumentation amplifier, the calculated sensitivity for the flow calorimeter corresponded to a temperature change of about 10^{-4} °C. This is higher than that for a true microcalorimeter, which can detect a temperature change on the order of about 10^{-5} °C (Steinberg, 1981).

Figure 2 shows peak areas that were gener ated by 30 mJ heat pulses and plotted as a function of flow rate. Peak areas obviously depended on flow rate. This dependence was taken into account when comparing heat data obtained at dif

ferent flow rates by applying a correction factor that was based on the linear relationship in Fig. 2.

Heat pulses for calibrating the instrument generally ranged from about 5 mJ to more than 100 mJ in size, corresponding to times of 2 to 45 s for energizing the calibrating resistor. Calibrating resistors were 22 k Ω and were energized at about 7.5 V. Figure 3 shows a series of heat pulses and the associated calibration curve, the latter corrected for differences in flow rate. Lin ear regression of heat pulse peak areas versus energy input consistently gave $R^2 > 0.99$.

Precision for replicated heat pulses generally resulted in coefficients of variation that were less than 5%. Precision for exothermic and endother mic peaks obtained on a single soil column were of a magnitude similar to that for heat pulses. Pre

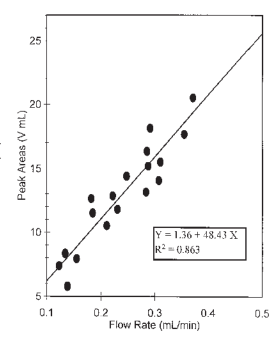


Fig. 2. Relationship between peak area and solution flow rate.

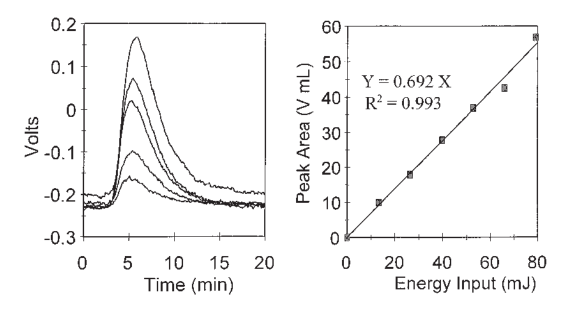


Fig. 3. Peaks obtained with various size heat pulses and the associated calibration curve.

cision obtained with replicated samples over periods of several months using different flow calorimeters and obtained with different opera tors was generally less than 15%. Because of their small size, reproducibility from sample to sample will depend strongly on sample homogeneity. In the case of the two Puerto Rican soils, use of 0.25 to 0.50 mm aggregates apparently resulted in a reasonably homogeneous material.

Ion Exchange

Using a crude calorimeter fitted with ther mometers, Coleman (1952) observed heats of ex change for alkali and alkaline earth cations on ex change resins that were typically in the range of 4 to 13 kJ mol_c⁻¹. Gaines and Thomas (1953) de scribed a method for deriving thermodynamic exchange constants from selectivity coefficients. Using the temperature dependence of the ex change constant, the standard enthalpy of ex change, ΔH° , can be calculated using the relation, ([pdiff]ln K/[pdiff]T) = $\Delta H^{o}/RT^{2}$. In several instances, enthalpies derived using this relation have compared well with calorimetric values, al though the standard enthalpy change, ΔH° , is not measured directly by calorimetry. Laudelout et al. (1968) corrected their calorimetric values to standard state conditions and found that they agreed well with values derived from the tem perature dependence of ln K. In the case of NH₄/Ca exchange on Camp Berteau montmo

rillonite, the corrected calorimetric value was 10.5 kJ mol_c⁻¹, and the one derived from tem perature dependence was 11.3 kJ mol_c⁻¹. Heats of exchange on soils and clay minerals generally follow the lyotropic series: Li < Na < K < Rb < Cs; Mg < Ca < Sr < Ba (Gast, 1972; Gast et al., 1969; Laudelout et al., 1968). Because the forces of attraction are predominantly electrostatic, the preference of the clay for cations of the same charge increases as their hydrated radii decrease. As a result of its carboxylate content, soil organic matter can exhibit properties of a strong field exchanger, reversing the selectivity for some cations as given by the lyotropic series (Juo and Barber, 1969).

Figure 4 shows the results for K/Ca exchange that we obtained on Dowex resin using flow cal orimetry. The resin was initially K saturated us ing 50 mM KCl. When the solution was changed to 25 mM CaCl₂, an endotherm corresponding to the displacement of exchangeable K by Ca was observed. When the solution was switched back to KCl, an exotherm corresponding to the dis placement of exchangeable Ca was observed. The two peaks in Fig. 4 are equal in area, as expected for a reversible ion exchange reaction. The quan tity of heat associated with the peaks was 26 mJ mg⁻¹, which is equivalent to 10.4 kJ mol_c⁻¹ for K/Ca exchange on this Dowex material. Cole man (1952) reported values of 10.5 and 11.3 kJ mol_c^{−1} for K/Ca exchange on Amberlite cation

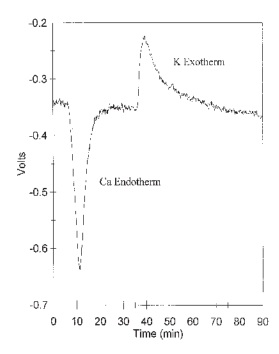


Fig. 4. Heats of K/Ca exchange obtained using $6.2~\mathrm{mg}$ of Dowex 50W-8 exchange resin.

exchange resins IR120 and IR100, respectively. He also found that Ca replacing exchangeable K was endothermic.

Figure 5 shows K/Ca exchange on a sample of the Oxisol. The two peak areas each measured 0.47 mJ mg⁻¹ of soil, which corresponds to a heat of exchange of 7.8 kJ mol₆⁻¹. Figure 6 shows K/Na exchange on the Ultisol. These two peaks measured 0.82 mJ mg⁻¹, which corre sponds to 7.5 kJ mol_c⁻¹. These heats of exchange are similar to values reported for soils and clays that were obtained using conventional batch cal orimeters or derived from temperature depen dence of the exchange constant. For example, Gast (1972) reported calorimetric heats of ex change for alkali metal cations on Chambers montmorillonite ranging from a high of 10.4 kJ mol_c⁻¹ for Li/Cs exchange, to a low of 0.4 kJ mol_c⁻¹ for Li/Na exchange. Deist and Talibu deen (1967) reported standard state enthalpies for K/Ca exchange on five soils in which the clay fraction was dominated by 2:1 layer silicates. Their ΔH^o values ranged from 3.8 to 35 kJ mol_c⁻¹. Udo (1978) derived the standard state enthalpy for K/Ca exchange on a kaolinitic soil clay and reported the value of 54 kJ mol_c⁻¹. Sparks and Jardine (1981) used a kinetic approach to derive K/Ca exchange constants at several

temperatures for a soil whose clay fraction was dominated by vermiculite, chloritized vermiculite, and mica. The temperature dependence of these exchange constants yielded standard en thalpies of exchange of 7.1 and 5.6 kJ mol_c⁻¹ for samples from the A and B horizons, respectively. Thus results obtained from flow calorimetry for ion exchange are consistent with those obtained using other methods.

Although the peak areas in Fig. 4 are equal, the peak shapes are different, indicating that the rates of exchange were not the same for the two displacement reactions. The lower peak maxi mum and the greater tailing for the K exotherm compared with the Ca endotherm in Fig. 4 indi cate that K displaced exchangeable Ca at a slower rate than the reverse reaction. In contrast, K/Ca exchange on the Oxisol (Fig. 5) showed no evi dence that the rates of exchange for these two cations were different. Exchange of K/Na on the Ultisol (Fig. 6) exhibited considerable disparity in peak shapes, indicating that Na had difficulty replacing exchangeable K. Sparks and Jardine (1981) observed a slower rate of K desorption from soil than that for K adsorption (K/Ca ex change) and attributed this to greater activation energy for desorption. Their soil contained ver

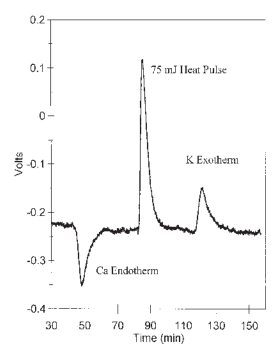


Fig. 5. Heats of $\mbox{K/Ca}$ exchange obtained using 68.4 mg of Oxisol.

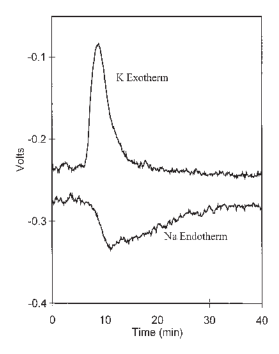


Fig. 6. Heats of K/Na exchange obtained using 57.2 mg of Ultisol.

miculite, and the rate limitation was ascribed to partial collapse of the 2:1 layer silicate following K saturation. Calorimetric studies of K/Na ex change on a high charge smectite in our lab showed peaks of considerably different shape as a result of kinetics. The peaks were also unequal in area because of some irreversibility in the ion ex change process (data not shown). The calorimet ric data suggested that Na was unable to displace exchangeable K completely, presumably because of its inability to expand the lattice spacing fol lowing K saturation. This conclusion was con firmed by X ray diffraction analysis (data not shown), which showed that the K saturated smectite did not fully expand on subsequent treatment with NaCl.

Phosphate Sorption

Phosphate sorption by soils typically exhibits an initial, rapid uptake phase, thought to be a li gand exchange reaction, followed by a much slower rate of uptake that may continue for weeks or months. This slow uptake is thought to be the result of one or more of the following processes: i) diffusion to less accessible sites within pores of solid aggregates (Willet et al., 1988; Cabrera et al., 1981; Parfitt, 1989; Madrid and de Arambarri, 1985; Lookman et al., 1994); ii) penetration into

the amorphous Fe and/or Al oxides by solid state diffusion (van Riemsdijk and de Haan, 1981; van Riemsdijk et al., 1984; Barrow, 1983); and iii) pre cipitation with metals derived by dissolution of the soil matrix (Aulenbach and Meisheng, 1988; Lookman et al., 1994; Martin et al., 1988; Nanzyo, 1984, 1986; Pierzynski et al., 1990).

Malati et al. (1993) reported that phosphate sorption on kaolinite, mica, and anatase at pH < 5 was endothermic and generally in the range of +10 to +15 kJ mol⁻¹. However, at pH 7, phos phate sorption by kaolinite was exothermic and similar in energy to hydrogen bonding. The en dothermic heats were derived from the energy terms in the Langmuir equations used to describe their sorption isotherms. Hundal (1988) reported that phosphate sorption by a clay loam soil was endothermic based on results obtained by apply ing the Clapeyron equation to liquid/solid phase equilibrium. The isosteric heat of adsorption var ied with surface coverage, ranging from about 9.2 kJ mol⁻¹ to about 18 kJ mol⁻¹. Muljadi et al. (1966a) found that an increase in temperature in creased phosphate sorption by kaolinite, gibbsite, and pseudoboehmite but that this increase was largely the result of an irreversible increase in the number of sorption sites. The heat of reaction between 200 mL of 0.1 MKH₂PO₄ and 20 g of K saturated kaolinite was less than the sensitivity of their calorimeter, 0.2 kcal mol⁻¹, leading them to conclude that the driving force for phosphate sorption by these materials was essentially en tropic. Contrary to the above, Barrow (1983) and Froelich (1988) presented data that indicated that phosphate sorption was exothermic since an in crease in temperature decreased phosphate sorp tion. Halter and Pfeifer (2001) reported that sorption of arsenate, an oxyanion chemically sim ilar to phosphate, by α Al₂O₃ also decreased with increasing temperature.

Figure 7 shows calorimetric results obtained for a sample of the Ultisol that was initially equil ibrated with a solution of 50 mM NaCl. Curve 1 resulted when this solution was changed to one in which 3 mmol L⁻¹ of the NaCl had been re placed with NaH₂PO₄ (i.e., keeping the total Na concentration at 50 mM). The reaction of phos phate with the soil was exothermic, 0.42 mJ mg⁻¹. Returning to the original NaCl solution after the NaH₂PO₄ treatment produced no heat signal (data not shown), indicating that the peak observed with NaH₂PO₄ was not caused by re versible Cl/H₂PO₄ exchange. Curve 2 in Fig. 7 was obtained during the second cycle of NaH₂PO₄ treatment and yielded 0.21 mJ mg⁻¹.

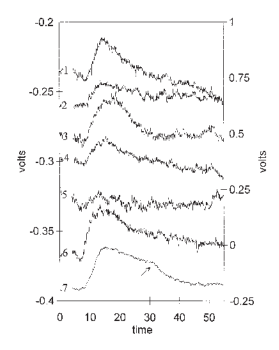


Fig. 7. Heats of reaction of phosphate on Ultisol (Curves 1 through 6) and amorphous Al(OH)₃ (Curve 7). Curves 1 through 6 are read on the left-hand scale; Curve 7 on the right-hand scale. The arrow on Curve 7 indicates a change from phosphate solution back to NaCl.

The NaCl/NaH2PO4 solution had been ad justed to the same pH (5.8) as the NaCl. Re versibility of phosphate sorption with pH has been observed (Muljadi et al., 1966b). To deter mine whether the phosphate reactive sites on the Ultisol could be regenerated, the soil was treated with 50 mM NaCl to which sufficient Na₂CO₃ had been added to obtain a pH of 10. The reac tion of the pH 10 solution with soil was en dothermic (data not shown). Changing the solu tion back to NaCl, pH 5.8, produced no detectable heat signal. When the soil was then ex posed to phosphate, an exotherm larger than that obtained in the first phosphate cycle was ob served, 0.64 mJ mg⁻¹ (Curve 3). Curve 4 shows the heat of reaction obtained in the fourth phos phate treatment cycle, which measured 0.21 mJ mg^{-1} . By the fifth cycle (Curve 5), the phosphate reaction was essentially undetectable, indicating that phosphate had reacted with all of the avail able sorption sites. The soil was then given a sec ond pH 10 treatment followed by NaCl pH 5.8, and this was followed by the sixth phosphate treatment cycle. The heat generated during the sixth phosphate cycle was identical to that fol

lowing the first pH 10 treatment, 0.64 mJ mg⁻¹ (Curve 6), suggesting that the second pH 10 treatment had regenerated the same number of phosphate reactive sites as the first pH 10 treat ment. The reaction of phosphate with amor phous Al(OH)₃ (Curve 7) was also exothermic, suggesting that amorphous Al(OH)₃ could have been responsible, at least in part, for the exother mic phosphate reaction in the Ultisol.

The reaction between phosphate and Al(OH)₃ can be described as Al(OH)_{3(crystalline)} + $PO_4^{-3} = AlPO_{4(crystalline)} + 3OH^{-}$ and, under standard state conditions, is endothermic, +129 kJ mol⁻¹ (Wagman et al., 1982). The direct pre cipitation of Al+3 with phosphate is also en dothermic under standard state conditions, +75 kJ mol⁻¹ (Wagman et al., 1982). To verify that precipitation of Al phosphate was endothermic under our column conditions, the heat of reac tion was measured when exchangeable Al was displaced from the Dowex resin, first by 50 mM KCl (exchange only) and then by 50 mM KH₂PO₄ (exchange plus precipitation). Curve 1 in Fig. 8 shows the exotherm that resulted when KCl displaced exchangeable Al from the Dowex resin. Curve 2 was obtained when the same

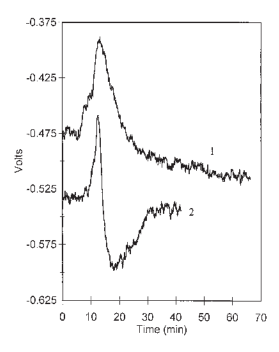


Fig. 8. Heat obtained by displacing exchangeable AI on 5.2 mg Dowex 50W-8 resin with KCI (Curve 1) and that obtained by displacing the AI with KH_0PO_4 (Curve 2).

Dowex column was resaturated with Al (by flushing with 10 mM AlCl₃, pH 4) and the ex changeable Al displaced using KH₂PO₄. The re action was initially exothermic, consistent with K/Al exchange. However, the reaction shifted to endothermic, consistent with a secondary Al phosphate precipitation. The delayed start of the endothermic reaction would be expected since the Al concentration in solution must increase sufficiently after the onset of K/Al exchange to induce Al phosphate precipitation. However, once precipitation began, it proceeded rapidly and continued until all of the Al had been ex changed from the resin. The fact that precipita tion of Al phosphate was endothermic whereas the reaction of phosphate with soil and Al(OH)₃ was exothermic shows that precipitation was not the primary mechanism for phosphate sorption in this study.

An alternative to a precipitation mechanism is ligand exchange. If the exothermic reaction of phosphate with soil is the result of ligand ex change only, then ligand exchange is a slower reaction than ion exchange. For example, K dis placement of exchangeable Ca and Na was com plete in both soils in less than 20 min (Figs. 5 and 6). Phosphate sorption on the Ulitsol took three 30 to 40 min cycles for a total of more than 100 min (Fig. 7). In the case of Al(OH)₃, the reaction was still going strong after 20 min and would have required considerably longer exposure to saturate the sorption capacity (Fig. 7). It is possi ble that phosphate sorption begins as a fairly rapid ligand exchange reaction but is followed by a slower secondary reaction. This secondary reac tion could be a diffusion limited process involv ing sites of limited accessibility on the surface or penetration into the interior of amorphous oxides/hydroxides. Either explanation could ex plain the slow evolution of heat accompanying phosphate sorption. A definitive explanation for the exothermic phosphate reaction will require additional information such as the amount of phosphate sorbed, pH and compositional changes in the effluent during sorption, and chemical and physical analysis of the solid phase after reacting with phosphate.

SUMMARY AND CONCLUSIONS

A simple, inexpensive flow calorimeter has been used to measure heats of reaction in soil. It uses a pair of thermistors to monitor changes in temperature of a solution as it flows through a column containing 50 to 100 mg of soil. A change in solution temperature generates a signal

that is amplified and fed into a computer for processing. The calorimeter exhibits high sensitivity, low thermal drift, and good signal to noise ratio.

The flow calorimeter was used to study two important soil chemical processes, cation ex change and phosphate sorption. Heats for K/Ca exchange measured by flow calorimetry were consistent with values that have been reported for other soils, clays, and cation exchange resins. In addition to heats of exchange, flow calorimetry provided insight into the kinetics of the exchange process. K displacing exchangeable Ca occurred more slowly than the reverse reaction on Dowex resin, whereas the rates were similar for K/Ca ex change on the Oxisol. Na experienced consider able difficulty in replacing exchangeable K from the Ultisol, which contained some expandible 2:1 layer silicate clay.

The reaction of phosphate with Ultisol and amorphous Al(OH)₃ was exothermic. In the Ul tisol, the phosphate exotherm decreased in mag nitude with consecutive phosphate cycles and was essentially gone after three cycles. However, reactive sites were regenerated by treating the soil with a solution at pH 10, showing that the phos phate reaction was reversible with pH. The pre cipitation of Al phosphate was endothermic, showing that precipitation was not the primary mechanism for phosphate sorption in this study. The results of this investigation show that flow calorimetry can provide useful information about surface chemical reactions in soils that cannot readily be obtained by other methods. Used in conjunction with other techniques, it should help improve our understanding of many important soil chemical processes.

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