Dolomite Lime’s Reaction Applied on the Surface of a Sandy Soil of the Northwest Paraná, Brazil

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ABSTRACT

Low Ca and Mg are serious limitations to crop production in sandy soils of the northwest Paraná, Brazil. Thus soil samples of an Oxisol collected in this region were packed into 30cm long columns. Dolomite lime (2.0, 0.84, 0.30, and < 0.30 mm screen) was added on soil surface, then leached with deionized water. Thereafter, the columns were dismantled and the soil cut into 5cm segments for chemical analysis. Dolomite lime increased pHCl, KCl-exchangeable Ca and Mg, and residual CO3 mostly in the top surface layers. Surface dolomite lime had no effect on pH, Ca, Mg, and CO3 in the leachate, independent on the lime particle size. These results indicated that surface dolomite lime application had no effect on subsoil composition and mostly of the calcium and magnesium carbonates are still unreacted on the soil surface.

Key words: Liming, Ca and Mg leaching, subsoil acidity

INTRODUCTION

Among the major soil limitations to crop production in northwest Paraná State, Brazil are: extremely deficient levels of Ca, Mg, and K, low cation exchange capacity (CEC) and organic matter contents, and very sandy texture. Fidalski & Auler (1997) and Fidalski et al. (1999) observed that Ca and Mg were the highest nutritional deficiencies in citrus orchards in this region. They also demonstrated that leaf Ca and Mg were positively related with citrus yields. They concluded that dolomite lime is essential for an economic citrus production in this soil.

In long-term pasture and in established perennial crops where deep soil cultivation is not possible, lime may be spread on soil surface and allowed time to leach. Even when cultivation is possible in citrus orchard, lime incorporation is unlikely with conventional machinery and leaching of the ions from the lime is relied upon for amelioration of acidity deeper in the soil. There are some evidences that surface applied lime in this soil has little effect on both surface and subsurface chemical properties (Rêgo, 1997; Fidalski, 1999). These authors reported that soil samples taken after twelve months of dolomite lime application on soil surface showed no effects on pH and KCl-exchangeable Ca, Mg, and Al. Their hypotheses are that lime was either leached below the root zone or was still unreacted with the soil acidity. To test these hypotheses, laboratory experiments were conducted with PVC columns with soil samples collected in this region to identify the main chemical reactions involved in the surface applied dolomite lime.
MATERIAL AND METHODS

Soil samples were collected from a cultivated site on the Instituto Agronômico do Paraná (IAPAR) experimental station at Paranavaí, Brazil (23°05'S, 52°26'W, 480m above sea level). This soil was selected based on following characteristics: sandy texture, low basic cations contents, and their potential use for production of citrus and pasture. The soil had an original pH CaCl$_2$ value of 5.0; exchangeable Ca, Mg and K contents of 12.8, 6.6, and 3.0 mmolc dm$^{-3}$, respectively; total acidity (H + Al) of 29.4 mmolc dm$^{-3}$, organic carbon content of 10.31 g kg$^{-1}$ and clay, silt, and sand contents of 11, 1, and 88%, respectively.

Columns (length 30cm; diameter 4cm) of rigid polyvinyl chloride (PVC) were used to conduct the experiment. Soil samples were transferred to PVC columns and compacted to a homogeneous bulk density (mean 1.4 g cm$^{-3}$). Then, the columns were brought to field capacity with distilled water. Approximately 1 kg of agricultural dolomite limestone (27.3% CaO, 20.7% MgO, and 100% neutralizer power) containing both coarse and fine materials was passed through 2.0, 0.84, 0.30, and <0.30 mm screen. A mechanical shaker was used to separate the lime particle sizes.

Lime was added on the soil surface at a rate of 3.96g per column in order to neutralize two times the total acidity (H + Al) contents. The irrigation program was 330 ml of water per column, equivalent to three porous volume, added at a rate of 2.2 ml min$^{-1}$.

At each porous volume, the drainage water was collected and analyzed for pH, Ca, Mg and CO$_3$ contents. Soil samples were collected at 0-5, 5-10, 10-15, 15-20 and 20-25 cm depth, air dried, ground to pass 2mm sieve and analyzed for pH CaCl$_2$, KCl-exchangeable Ca and Mg, and total CO$_3$. Ca and Mg were determined by ICP (Inductively Coupled Plasma) and total CO$_3$ content by conductivimetry using flow injection analysis coupled with pervaporation (Kawazaki et al., 2000, Grassi, et al., 2002). All treatments had three replicates in a completely randomized block design.

RESULTS AND DISCUSSION

Figures 1, 2, 3 and 4 show soil pH, KCl-exchangeable Ca and Mg and total CO$_3$ content at different depths as a result of surface application of dolomitic lime. For simplicity, data included only the effects of control, the coarsest (2.0 mm) and the finest limestone (<0.3 mm).
Dolomite lime increased pH, exchangeable Ca, Mg, and CO3 mostly in the top surface layers. The finest the lime the highest the increase in pH, Ca and Mg, corroborating data presented by Verlengia & Gargantini (1972). The lime particle size had not clear effect on the mobility of Ca and Mg with depth. The low Ca and Mg distribution patterns can be explained by the electrical neutrality principles which state that Ca and Mg can not move down in the soil profile unless accompanied by equivalent anions.

Figures 5, 6, 7 and 8 show the effect of surface applied dolomite lime on drainage water composition. Ca, Mg, CO3 and pH in the leachates were consistent and complementary to their distributions in the soil profile.

With increasing pH, there was an increase in pH and a decrease in Ca, Mg and CO3 concentrations as a result of dilution. Surface applied dolomite had little effect on drainage water pH, Ca, Mg and CO3. These results (soil and drainage) indicated little mobility of surface applied lime in this sandy soil and most of calcium and magnesium carbonates were still unreacted on the soil surface.

The low rate of lime movement into the soil profile was probably a result of the low solubility of the CaMg(CO3)2, weak soil acidity, and the lack of stable soluble anion as a product of the reaction of dolomite with soil acidity. The electrical neutrality principle dictates that Ca and Mg movements must be accompanied by equivalent
anions. The dolomite lime reaction is illustrated as following:

\[
\text{CaMg(CO}_3\text{)}_2 + 4\text{H}^+ \rightarrow \text{Ca}^{2+} + \text{Mg}^{2+} + 2\text{CO}_2 + 2\text{H}_2\text{O}
\]

This reaction neutralizes \(\text{H}^+\) and release \(\text{Ca}^{2+}\) and \(\text{Mg}^{2+}\), resulting an increase in soil \(\text{pH}\) and \(\text{pH}\)-dependent CEC. The \(\text{Ca}^{2+}\) and \(\text{Mg}^{2+}\) stoichiometrically replace \(\text{H}^+\) on the existing \(\text{pH}\)-dependent CEC whose magnitude changes under the influence of the increase alkalinity. As expected, it was found higher exchangeable \(\text{Ca}\) than \(\text{Mg}\) after dolomite lime reaction in this soil due to higher selectivity of the colloidal negative sites for \(\text{Ca}^{2+}\) than for \(\text{Mg}^{2+}\) (Loyola & Pavan, 1989). The above reaction will continue to the right as long as the soil supplies acidity in the form of \(\text{H}^+\) ions.

The results of this study clearly indicated that surface lime applications in this soil had little effect on the subsoil acidity and most of \(\text{Ca}\) and \(\text{Mg}\)-carbonates are still unreacted on the soil surface. Seldom would lime be applied to a topsoil with the objective of improving the subsoil fertility without any acid inputs. For this soil, without acid inputs, transfer of alkalinity to the subsoil is unexpected.

**CONCLUSION**

Agriculturally realistic liming rates are not likely to ameliorating subsurface soil acidity on the Northwest Paraná without acid inputs. The carbonates stay unreacted on the soil surface.

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**REFERENCES**


