Effect of Exogenous Carbon on Movement of Simazine and 2,4-D in Soils

L. Cox, A. Cecchi, R. Celis, M. C. Hermosin, W. C. Koskinen, and J. Cornejo*

ABSTRACT

Organic amendment (OA) addition is an agricultural practice that can greatly affect pesticide behavior in soil. The influence of three OAs, consisting of a liquid amendment (LOA), solid (SOA) humic amendment, and a solid urban waste (SUW), on sorption, persistence, and leaching potential of simazine (2-chloro-4,6-bis (ethylenimino)-s-triazine) and 2,4-D (2,4-dichlorophenoxyacetic acid) in a sandy soil (P2) has been investigated. The OAs were added to P2 at a rate of 5% (w/w). Sorption was studied using the batch equilibration technique, degradation by incubations at 20°C and −0.33 kPa moisture content, and leaching using handpacked soil columns. Simazine sorption increased in the order of P2 < P2 + LOA < P2 + SUW < P2 + SOA. The herbicide, 2,4-D, did not sorb on P2 soil, but in amended soils sorption increased in the same order as simazine. The lower sorption of both herbicides in P2 + LOA as compared with P2 + SOA and P2 + SUW is attributed to higher dissolved organic matter (DOM) content of LOA. Simazine’s half-life was reduced upon amendment. Degradation of 2,4-D followed a sigmoidal rate in P2, P2 + SOA, and P2 + SUW, but not in P2 + LOA, in which 2,4-D persistence was much higher. The higher stability of 2,4-D in P2 + LOA has been attributed to 2,4-D-dissolved organic C interactions. Leaching studies indicate that degradation affects movement to a higher extent than sorption, specially in the case of 2,4-D in P2 + SUW soil, in which the amounts leached are much higher than those in unamended soil. In the case of simazine and P2 + SUW soil, greater reduction in large-size conducting pores upon amendment, as revealed in Hg porosimetry studies, resulted in a greater reduction of leaching than that suggested from the small differences in sorption and degradation between P2 and P2 + SUW soil.

Pesticides are recognized as a source of potential adverse environmental impacts (nonpoint and point pollution) and their presence in surface and ground waters has grown considerably (Wauchope, 1978; Ritter et al., 1994). Soil applied herbicides reach surface and ground waters by the losses associated with runoff and leaching processes (Beck et al., 1993). These losses are attenuated by the natural process of sorption, since degradation, transport, and biological activity of pesticides are greatly influenced by sorption on soil constituents, in particular soil organic matter (Chiou, 1989; Koskinen and Harper, 1990).

Organic amendments, commonly used to enrich low organic matter content soils, can modify surfaces of soils and subsurface materials increasing sorption potential and reducing pesticide contamination of ground water (Zsolnay, 1992; Barriuso et al., 1996; Cox et al., 1997). The addition of OAs to the soil surface is also known to stimulate soil microbial activity, which could potentially lead to accelerated degradation, reducing the total amount of chemical available for leaching and to injure crops planted in rotation (Felsot and Dzantor, 1995; Topp et al., 1996; Cox et al., 1997). Accordingly, the study of the influence of OA on pesticide behavior in soil is of great interest, specially when it is an issue very seldom considered when making the decision of fertilizing soil or disposing organic wastes; other aspects being considered are nutritional ones, risk of nitrate leaching, or accumulation of heavy metals in soil.

The incorporation of OAs in soils introduces both solid organic matter and DOC. Dissolved organic C has been shown to reduce pesticide sorption because of DOC-pesticide interactions and competition between DOC molecules and pesticide molecules for sorption sites (Lee et al., 1990; Barriuso et al., 1992; Businelli, 1997; Celis et al., 1998a; Cox et al., 2000). This decrease in sorption would enhance transport and increase the risk of ground water contamination. Consequently, the effect of OA addition on pesticide behavior is not easy to predict, especially when liquid amendments are applied or when DOM concentrations resulting from the OA are high.

The aim of this study was to assess the influence of OA (commercial LOA, SOA, and a SUW) on sorption and degradation of pesticides, and the potential applicability of these amendments for reducing pesticide leaching in soil. We have selected two herbicides widely used in agriculture, simazine and 2,4-D. Simazine is relatively persistent and, despite its low water solubility (5 mg L⁻¹), its significant nonpoint source of water contamination is reported (Wauchope, 1978; Agbekodo et al., 1996). The anionic herbicide, 2,4-D is of relatively high water solubility (620 mg L⁻¹) and, as other anionic contaminants, its presence in ground water is a concern because anionic contaminants are weakly retained by most soil or sediment components (Goodrich et al., 1991; Hermosín and Cornejo, 1993).

MATERIALS AND METHODS

Herbicides, Soils, and Amendments

Simazine is a colorless solid with a melting point of 226°C, a vapor pressure of 810 nPa, and a water solubility (20°C) of 25 μM (Worthing and Hance, 1991). As other chloros-triazines, the hydroxylated and N-dealkylated derivatives are, respectively, major chemical and biological degradation products of simazine (Khan, 1980). High purity (>98%) simazine was purchased from Riedel-de Haën (Germany). The herb-


Table 1. Physicochemical properties of the unamended soil (P2) and soil amended with 5% (w/w) of liquid organic amendment (LOA), solid organic amendment (SOA), and solid urban waste (SUW).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>P2</th>
<th>SOA</th>
<th>SUW</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH (1:5)</td>
<td>7.1</td>
<td>6.6</td>
<td>20</td>
</tr>
<tr>
<td>Dry matter, g kg⁻¹</td>
<td>754</td>
<td>164 (600, 200, 200)</td>
<td>43</td>
</tr>
<tr>
<td>Organic C, g kg⁻¹</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dissolved organic C, mg L⁻¹</td>
<td>52</td>
<td>52</td>
<td>52</td>
</tr>
</tbody>
</table>

Table 2. Physicochemical properties of the unamended soil (P2) and soil amended with 5% (w/w) of liquid organic amendment (LOA), solid organic amendment (SOA), and solid urban waste (SUW).

<table>
<thead>
<tr>
<th>Sample</th>
<th>pH</th>
<th>Organic C</th>
<th>DOC</th>
<th>Clay (I, K, M)</th>
<th>Fe, P₂O₅ g kg⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>P2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LOA</td>
<td>7.1</td>
<td>6.6</td>
<td>20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SOA</td>
<td>7.7</td>
<td>8.8</td>
<td>100</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SUW</td>
<td>7.7</td>
<td>7.8</td>
<td>25</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Dissolved Organic Carbon Measurements and Dialysis Experiment

To determine the amount of DOC in the extracts was measured in a Shimadzu 5050 Total Organic Carbon Analyzer (Shimadzu, Columbia, MD).

Sorption Studies

Sorption studies were performed using the batch equilibration procedure. Duplicate samples of 5 g of each soil were treated with 10 mL of simazine solutions of initial concentrations (Ci) ranging from 0.5 to 10 μM, or 10 mL of 2,4-D solutions of Ci ranging from 5 to 100 μM. The suspensions were shaken at 20 ± 2°C for 24 h and then centrifuged at 31,000 × g at the same temperature. Previously, it was determined that equilibrium was reached in <24 h, and that no measurable degradation occurred during this period. Sorption controls (10 mL of solution without soil) were also shaken for 24 h and recoveries were close to 100%. Equilibrium concentrations (Ce) in the supernatants were determined by high-performance liquid chromatography (HPLC) under the following conditions: Nova-Pack column (Waters Corp., Milford, MA), 150 mm length by 3.9 mm i.d.; column packing, C18; flow rate, 1 mL min⁻¹; eluent system, 70:30 water/acetonitrile mixture for simazine and 55:45 diluted H₃PO₄ (pH = 2.0)/methanol for 2,4-D; UV detection at 230 nm. Differences between Ci and Ce were assumed to be the amounts sorbed by the solid phase (Cs). Sorption isotherms were fit to the Freundlich equation and sorption coefficients, Kf, and 1/n, calculated. Sorption coefficients, Kf, normalized to organic C content of the soils (Kfₒ) were also calculated.

Incubation Studies

Duplicate amounts (400 g) of each soil (P2, P2 + LOA, P2 + SOA, and P2 + SUW) were treated with a 100 mg L⁻¹ ethanol solution of simazine or a 100 mg L⁻¹ aqueous solution of 2,4-D to give a concentration of 5 mg kg⁻¹ and 2.3 mg kg⁻¹ of dry soil, respectively. Maximum application rates of simazine and 2,4-D are 5 and 2.3 kg ha⁻¹, respectively. Moisture content was adjusted to −0.33 kPa, and soil samples were thoroughly mixed by passing them through a sieve at least four times. Herbicide treated soil samples were transferred to kiln jars where they were incubated at 20°C for 28 d. Moisture contents were maintained at a constant level throughout the experiment by adding distilled water as necessary, soils were sampled periodically and frozen till analyzed. Herbicide residues were extracted by shaking 10 g of soil with 20 mL of DOC in the extracts was measured in a Shimadzu 5050 Total Organic Carbon Analyzer (Shimadzu, Columbia, MD).
soil, which can be considered not readily available (Cox et al., 1998).

Soil Column Studies

Herbicide leaching was studied under near-saturated conditions using 30 by 5 cm methacrylate columns made up of six 5-cm long sections sealed with silicon. The top ring was filled with sea sand and the bottom ring with sea sand plus glass wool, to minimize losses of soil, and contamination of leachate with soil particles. The other four rings were handpacked with 520 g of soil. Columns were saturated with 0.01 M CaCl₂ and allowed to drain for 24 h. After the soil columns were saturated, ~1 g of soil was removed and the distribution of pore radii of soil samples from 40 to 3.7 × 10⁻³ μm was determined in duplicates using a Carlo Erba 2000 Hg depression and intrusion porosimeter (Carlo Erba, Milan, Italy). A 0.5-g subsample of each soil was heated at 90°C for 24 h and then outgassed at room temperature for 30 min. A value for the surface tension of Hg of 0.48 N m⁻¹ and a contact angle on soils of 141.3° was used with the Laplace equation assuming cylindrical pores in the calculations.

The amounts of herbicide corresponding to an application rate of 5 (simazine) or 2.3 (2,4-D) kg ha⁻¹ were applied to the top of triplicate saturated columns as a methanol solution and left the methanol evaporate. Twenty four hours after herbicide application, columns were leached with 0.01 M CaCl₂, which was applied at a rate of 50 mL d⁻¹. Leachates were collected daily for 20 d, filtered with 0.22 μm pore-size filters, and analyzed directly by HPLC for their herbicide concentration as described earlier.

RESULTS AND DISCUSSION

Dissolved Organic Carbon of Amendments and Amended Soils

Total organic matter contents among OAs and among amended soils are very similar, although there were large differences in DOC content (Tables 1 and 2). Dissolved organic C of the OAs and amended soils decrease in the order LOA >> SOA >> SUW. The association of DOC and of OA to P2 soil was high for LOA (60%) but very low for SOA and SUW; nearly 100% of the DOC of the SOA and SUW did not associate to P2. Previous studies have indicated that LOA contains great amounts of relatively nonhumified material with high affinity for montmorillonite (Cox et al., 2000), which represents 20% of the clay fraction of the P2 soil (Table 2). In contrast, the DOC of SOA and SUW amendments contain high amounts of highly humified material (Cox et al., 2000), which appears to associate to soil to a slightly lesser extent. On the other hand, results from the dialysis experiment indicated that the DOC from LOA associates with 2,4-D but not with simazine; 2,4-D concentration inside the dialysis membrane (with DOC from LOA) was 10% higher than outside, whereas simazine concentrations inside and outside the dialysis membrane were identical.

Sorption Studies

Sorption of simazine on Soil P2 greatly increased upon amendment with SOA, whereas there was a slight increase in the case of SUW amended soil, and no significant differences between sorption on P2 + LOA and on original unamended soil (Fig. 1a). Isotherms are C-type (1/n close to 1), indicating that simazine sorption in original and amended soils is mainly a partitioning mechanism between solid organic matter and solution, without an apparent limit of sorption (Giles et al., 1960). Sorption isotherms were fit to the Freundlich equation.
Table 3. Freundlich sorption coefficients for simazine and 2,4-D in the unamended P2 soil and amended soils, P2 soil plus liquid organic amendment (P2 + LOA), P2 soil plus solid organic amendment (P2 + SOA), and P2 soil plus solid urban waste (P2 + SUW).

<table>
<thead>
<tr>
<th>Soil</th>
<th>2,4-D</th>
<th>Simazine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(K_f)</td>
<td>(1/n_f)</td>
</tr>
<tr>
<td>P2</td>
<td>0.59 (0.53–0.65)†</td>
<td>0.95 ± 0.08</td>
</tr>
<tr>
<td>P2 + LOA</td>
<td>0.61 (0.54–0.68)</td>
<td>0.93 ± 0.09</td>
</tr>
<tr>
<td>P2 + SOA</td>
<td>1.51† (1.38–1.66)</td>
<td>0.86 ± 0.08</td>
</tr>
<tr>
<td>P2 + SUW</td>
<td>0.66 (0.62–0.71)</td>
<td>0.99 ± 0.05</td>
</tr>
</tbody>
</table>

* Significant at the 0.05 probability level.
† Numbers in parentheses are standard errors about the mean.
‡ No sorption measured.

and sorption coefficients, \(K_f\), \(1/n_f\), and \(K_{oc}\) (\(K_f\) normalized to soil organic C content) were calculated (Table 3). Sorption coefficient, \(1/n_f\), was similar for the three soils, thus allowing comparison of \(K_f\) values. Simazine sorption increased by a factor of 2.5 upon amendment with 2,4-D, although sorption is higher in P2 amended soils.

The amounts of simazine extracted from P2 + SUW were slightly reduced when compared with unamended P2, whereas in the case of P2 + LOA and P2 + SOA disappearance from soil extracts is much faster, ~85% remaining after 28 d compared with P2 and P2 + SUW. Extrapolated half-lives are one-third of that in P2 soil (Fig. 2a and Table 4). The increase in organic C content upon amendment has been shown to promote biodegradation by enhancing microbial populations and activities (Felsot and Shelton, 1993; Guo et al., 1993; Topp et al., 1996; Cox et al., 1997; Gan et al., 1998). The lower amounts of simazine extracted from P2 + LOA and P2 + SOA as compared with P2 + SUW, can be attributed to the lower organic matter and DOC content of P2 + SUW (Table 2), and to some toxic effect of heavy metals, more abundant in SUW residue (Table 1) than in agricultural origin SOA or LOA, on soil microorganisms (Häni et al., 1996; Giller et al., 1999).

The lower sorption of simazine in P2 + LOA soil can be attributed to interactions between small DOC molecules and soil surfaces. In the case of simazine, the increase in soil sorption sites upon amendment would be counterbalanced by the competition between DOC molecules of LOA and simazine molecules, rendering similar sorption as the original Soil P2. In the case of 2,4-D, although sorption is higher in P2 + SOA and P2 + SUW than in P2 + LOA, the amendment with LOA generates sorption sites in soil, since no sorption was measured in the case of the unamended P2 soil. These differences can be explained considering the different sorption mechanisms of the two herbicides. Simazine, as other \(s\)-triazines, sorbs on solid organic matter and on montmorillonite but not on Fe oxides (Celis et al., 1998b). Simazine sorbs mainly by H-bonding or proton transfer to organic matter, while sorption on clay minerals takes place as protonated species or as molecular species on hydrophobic microsites of montmorillonite surfaces (Celis et al., 1997). The herbicide, 2,4-D, sorbs on organic matter, does not sorb on montmorillonite (unless solution pH is low) and does sorb on Fe oxide through electrostatic interactions between negatively charged carboxylic groups and positively charged Fe oxide surface (Celis et al., 1998b; Cox et al., 2000). Previous work has shown that DOC of LOA blocks montmorillonite sorption sites to a higher extent than Fe oxide surfaces, consequently, competing with simazine molecules for sorption sites to a greater extent than with 2,4-D (Cox et al., 2000).

**Incubation Studies**

The amounts of simazine extracted from P2 + SUW were slightly reduced when compared with unamended P2, whereas in the case of P2 + LOA and P2 + SOA disappearance from soil extracts is much faster, ~85% remaining after 28 d compared with P2 and P2 + SUW. Extrapolated half-lives are one-third of that in P2 soil (Fig. 2a and Table 4). The increase in organic C content upon amendment has been shown to promote biodegradation by enhancing microbial populations and activities (Felsot and Shelton, 1993; Guo et al., 1993; Topp et al., 1996; Cox et al., 1997; Gan et al., 1998). The lower amounts of simazine extracted from P2 + LOA and P2 + SOA as compared with P2 + SUW, can be attributed to the lower organic matter and DOC content of P2 + SUW (Table 2), and to some toxic effect of heavy metals, more abundant in SUW residue (Table 1) than in agricultural origin SOA or LOA, on soil microorganisms (Häni et al., 1996; Giller et al., 1999).

In contrast to simazine, 2,4-D followed a sigmoidal rate of degradation in Soils P2, P2 + SOA and P2 +
even after a 25-d incubation period nearly 70% of the initially applied 2,4-D was extractable from soil. In general, 2,4-D dissipation rates have been reported to be lowest on the soil on which sorption was greatest (John-son et al., 1995), which does not agree with results ob-tained in sorption studies (Table 3). This very high per-sistence of 2,4-D in P2 + LOA amended soil can be attributed to a toxic effect of the LOA on soil bacteria or to DOM-2,4-D interactions, which protects herbicide from degradation. The latter will be discussed in the section below.

Soil Column Studies

Soil Pore-size Distribution

Total cumulative soil pore volume was reduced upon OA (Figure 3), because of a reduction of large-size or conducting pores (>1 μm), as shown in Table 5. Solid OAs (SOA and SUW) reduced soil porosity to a greater extent as compared with LOA, and this reduction was more significant in the case of the SUW, of lower DOC (Table 5). These data suggest that nonsoluble organic matter cements and aggregates together soil particles, blocking large-size conducting pores (Tisdall and Oades, 1982). In contrast, soil pores in the 1- to 0.01-μm range increased in the amended soils, especially in the case of the liquid amendment LOA, which has higher DOC content. Similar results were observed in a clay soil amended with olive-mill wastewater (Cox et al., 1997).

Herbicide Leaching

Simazine was detected in leachates from original Soil P2 slightly earlier than in leachates from amended soils, and the maximum simazine concentration was much higher (Figure 4a). Lower sorption will lead to less retardation and, hence, earlier breakthrough (Gamendinger et al., 1993; Beck et al., 1993). The similar breakthrough curves (BTCs) of simazine on P2 + LOA and P2 + SOA of different sorption capacity (Table 3) and similar half-lives (Table 4), indicate that degradation affects leaching of simazine to a higher extent than sorption processes. Simazine BTC in Soil P2 + SUW was between those corresponding to P2 and P2 amended with LOA or SOA, although K_{f} values are similar to those corresponding to P2 and P2 + LOA (Table 3). The higher persistence of simazine in P2 + SUW, when compared with the other amended soils (Table 4), can explain this different leaching behavior. However, sima-zine BTCs in P2 and P2 + SUW soils of similar sorption

<table>
<thead>
<tr>
<th>Herbicide</th>
<th>P2</th>
<th>P2 + LOA</th>
<th>P2 + SOA</th>
<th>P2 + SUW</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>t_{1/2} (d)</td>
<td>R^2</td>
<td>t_{1/2} (d)</td>
<td>R^2</td>
</tr>
<tr>
<td>Simazine</td>
<td>362 (326–408)</td>
<td>0.94</td>
<td>102** (95–109)</td>
<td>0.98</td>
</tr>
<tr>
<td>2,4-D</td>
<td>–</td>
<td>43 (38–49)</td>
<td>0.96</td>
<td>–</td>
</tr>
</tbody>
</table>

* Significant at the 0.05 probability level.
** Significant at the 0.01 probability level.
† Numbers in parentheses are standard errors about the mean.
and half-life (Fig. 1a, Tables 3 and 4), were very different; simazine BTC in P2 + SUW was broader, the maximum was lower and slightly shifted to the right, and tailing was more pronounced than in P2 soil.

The processes retarding pesticide leaching occur in the transition between mobile and immobile water phases, where the interaction between soil solution and soil surfaces through diffusion, sorption, and degradation processes takes place (Doran, 1987; Walker and Welch, 1989; Beck et al., 1993). Mercury porosimetry studies have shown a greater reduction in large-size conducting pores upon SUW amendment compared with the other OA (Figure 3, Table 5). This decrease could result in an increase in contact time of the herbicide with soil surfaces, enhancing sorption and degradation processes and, consequently, rendering differences in leaching behavior greater than those suggested from the small differences in sorption and degradation observed between P2 and P2 + SUW soils (Cox et al., 1997).

The 2,4-D moved more rapidly through soil columns than simazine (Fig. 4b). Differences between 2,4-D leaching in P2 and amended Soils P2 + SOA and P2 + SUW can be attributed to higher sorption of 2,4-D on these soils, and also to a reduction in large-size conducting pores (Table 5), which increases contact time with soil surfaces favoring degradation. Leaching of 2,4-D was not reduced upon amendment with LOA. A maximum of the BTC and total amounts leached were much higher in P2 + LOA soil. As indicated earlier, 2,4-D degradation in P2 + LOA soil, with greater DOC (Table 2), is much slower than in P2 or other amended soils.

Table 5. Pore volumes for different pore-size ranges of the original soil and amended soil obtained by Hg porosimetry.

<table>
<thead>
<tr>
<th>Soil</th>
<th>&gt;1 μm</th>
<th>1–0.1 μm</th>
<th>0.1–0.01 μm</th>
<th>&lt;0.01 μm</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>P2</td>
<td>150</td>
<td>9</td>
<td>14</td>
<td>7</td>
<td>180</td>
</tr>
<tr>
<td>P2 + LOA†</td>
<td>104*</td>
<td>14</td>
<td>19</td>
<td>4</td>
<td>141</td>
</tr>
<tr>
<td>P2 + SOA‡</td>
<td>89*</td>
<td>13</td>
<td>16</td>
<td>7</td>
<td>125*</td>
</tr>
<tr>
<td>P2 + SUW§</td>
<td>70**</td>
<td>14</td>
<td>15</td>
<td>8</td>
<td>107*</td>
</tr>
</tbody>
</table>

* Significant at the 0.05 probability level.
** Significant at the 0.01 probability level.
† P2 soil plus liquid organic amendment.
‡ P2 soil plus solid organic amendment.
§ P2 soil plus solid urban waste.
Fig. 4. (a) Simazine and (b) 2,4-D breakthrough curves in unamended sandy soil (Soil P2) and amended sandy soil plus liquid organic amendment (P2 + LOA), sandy soil plus solid organic amendment (P2 + SOA), and sandy soil plus solid urban waste (P2 + SUW).

The degree to which DOM will complex with organic chemicals depends on the nature and source of the DOM (Nelson et al., 1998). The herbicide-DOM interaction might be, in part, responsible for the greater 2,4-D persistence in P2 + LOA and, consequently, for greater leaching. As discussed above, dialysis experiment results strongly indicated that the small DOM molecules of LOA are able to associate with 2,4-D molecules (probably through electrostatic forces) but not with simazine. Most likely, facilitated transport of 2,4-D in P2 + LOA took place as a consequence of the formation of stable complexes which also protected herbicide molecules from soil microorganisms.

CONCLUSIONS

Organic amendments modify leaching behavior of herbicides by affecting soil physical properties such as soil porosity, sorption, and persistence. Knowledge of the nature of the exogenous organic C is essential, in particular the amount of DOC, to determine the potential leaching of pesticides. Consequently, the general idea that organic matter reduces the leaching potential of herbicides is not always true. Although difficult to extrapolate to field, our results indicate that simultaneous application of 2,4-D and LOA can increase the risk of groundwater contamination by this herbicide because of its high stability in the P2 + LOA amended soil.

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