Humic Acid Formation in Artificial Soils Amended with Compost at Different Stages of Organic Matter Evolution

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A composting process was conducted under optimal conditions for 150 d, obtaining three biomasses at different levels of maturity: raw material (RM), fresh compost obtained after 11 d of composting (FC), and evolved compost (EC) obtained after 150 d of composting. During the composting process, HAs were extracted and fully characterized by mass balance, DRIFT, and 1H and 13C-nuclear magnetic resonance spectroscopy. Each compost sample was incubated for 180 d in an artificial soil, after which HA extraction was repeated and characterized. To compare composts containing different amounts of labile organic matter (OM), an equal amount of unhydrolyzable OM was added to the soils. Our results indicated that compost HAs consist of a biologically and chemically stable fraction (i.e., the unhydrolyzable HA [U-HA]) and a labile fraction, whose relative contents depended on the composting duration. Humic acid from more EC contained a higher amount of recalcitrant fraction (aromatic carbon) and a lesser amount of labile fraction (aliphatic carbon) than HA from RM and FC. These results suggest that the humification process during composting preserves the more recalcitrant fraction of the compost-alkali soluble/acid insoluble fraction (HA-fraction). Incubation of composts in soil showed that due to the higher labile fraction content, HAs from raw material were more degraded than those from EC. The abundance of labile carbon of soil amended with less-evolved compost (RM and FC) allowed the more recalcitrant fractions of U-HA to be more preserved than in EC. These results suggest that less-evolved compost could contribute more than well evolved compost to the stable soil OM.

Soil quality depends on the soil organic matter (SOM) quantity, quality, and dynamics (Lal, 2000). Up to 70 to 80% of SOM in mineral soil is composed of humic substances (HS) (Piccolo, 2002). Compost can be beneficial for soil because it improves soil properties and contributes to the soil humus balance (Giusquiani et al., 1995; Quédraogo et al., 2001; Leifeld et al., 2002). The definition of HS is only used for operational purposes because it is based on the solubility in the aqueous solutions used as extractants (Piccolo, 2002), and the term humic acid (HA) is used to indicate the HS soluble in dilute alkali but insoluble in dilute acid (pH < 1) (Piccolo, 2002). Humic acid is a negatively charged colloid, is recalcitrant to biodegradation, and can be stored in soil for long periods (Qualls, 2004). Because of these properties, HA plays an important role in determining soil characteristics by influencing its chemical, physical, and biological properties. Consequently, compost HA is considered an indicator of the ability of compost to contribute to SOM balance (Adani et al., 2006a).

Humification during composting has been the subject of extensive study (Sugahara and Inoko, 1981; Inbar et al., 1990; Chen and Inbar, 1993; Chefetz et al., 1999; Genevini et al., 2002a). However, the nature of the humification process during composting remains unclear. Furthermore, the end products of compost humification differ from those of soils (González-Vila et al., 1999; Almendros et al., 2000).

It was previously reported that HA is formed ex novo during composting (Chen and Inbar, 1993). This report was based on the findings that the relative content of the HA fraction increased during composting. This interpretation of the humification process refers to the classical route of HA formation, which hypothesizes that bio-macromolecules are broken up into small constituents and then recombined to form chemically complex “geopolymers” (Senesi and Loffredo, 1999). However, when considering the absolute content of HA, there is evidence that during composting this content does not increase notably (Adani et al., 1999; González-Vila et al., 1999).

More recently, it was shown that humification during the composting process should be interpreted as the concentrating...
of pre-existing nonhydrolyzable HA molecules (proto-HA) (Chefetz et al., 1999; Genevini et al., 2002a, b). Also, during composting, modifications of nonhydrolyzable HA, such as decarboxylation, demethoxylation, and a general removal of O-alkyl and alkyl carbons, occur (González-Vila et al., 1999; Chefetz et al., 1999; Genevini et al., 2002a).

This view of the humification process does not conflict with more recent studies on the nature of humic substances that indicate that HA is composed of low-molecular-weight hydrophobic and amphiphilic compounds (lignin, polysaccharides, lipids, and peptides) derived from the degradation and decomposition of dead biological material, arranged in supramolecular structures (Piccolo, 2002; Simpson, 2002; Sutton and Sposito, 2005). Therefore, based on our previous work (Genevini et al., 2002a, 2002b; Chefetz et al., 1999; Adani et al., 2006b) and taking into consideration the new interpretation of the HA structure, it can be concluded that compost HAs are represented by a mix of low-molecular-weight molecules structured in a supramolecular association and whose composition depends on the duration of the degradation processes (the composting process removes more labile fraction).

The production of fresh compost by high-rate composting—compost produced without the curing phase—represents a new trend in composting (Scaglia et al., 2000). This composting strategy was devised to reduce composting time and produce a hygienic, stable, and safe product.

It can be argued that if humification of OM takes place during the curing phase (Chen and Inbar, 1993), then the humus content of fresh compost is limited compared with mature compost. However, if humification is considered as the conservation and modification of more recalcitrant biomolecules, then the differences in the contribution to the soil–humus balance depend only on the relative concentration of these molecules.

The aim of this work was to study the effects of the composting process on the concentration and preservation of pre-existing HA (nonhydrolyzable HA) in compost. This study investigated the effects of compost, obtained at different degrees of evolution (fresh compost [FC] obtained after 11 d of composting and evolved compost [EC] obtained after 150 d of composting), on the quantity and quality of soil HA.

**Materials and Methods**

**Compost Preparation**

The raw material (RM) used was an organic mix consisting of municipal solid waste and green waste collected from the city of Milano (North Italy) mixed together in a ratio of 2.5:1 (wet weight) or 1:1 (dry weight). Raw material (RM) was composted in a laboratory adiabatic reactor (Adani et al., 2004). Fresh compost was obtained at the end of the high-rate composting phase (11 composting days) when compost attained the medium biological stability as required by the regional rules of Lombardia (North Italy) (Scaglia et al., 2000). This product was characterized by an oxygen uptake rate expressed as a dynamic respiration index (DRI) of 1000 mg O₂ kg⁻¹ h⁻¹ (Adani et al., 2004). More EC was obtained after a total of 150 composting days (high-rate composting plus a curing phase). This product exhibited a DRI of 200 mg O₂ kg⁻¹ h⁻¹, which was lower than the DRI that defines mature compost (500 mg O₂ kg⁻¹ h⁻¹) (Adani et al., 2004).

All matrices (RM, FC, and EC) were sampled during composting, following procedures described by The U.S. Composting Council (1997). Samples were vacuum dried at 60°C, ground to a Ø <0.5 mm, and stored for subsequent use.

**Incubation Test**

The soil used for the incubation tests was a sandy mineral substrate consisting of sand (particle size Ø = 0.5–0.8 mm, pH 7) and clay (bentonite–montmorillonite–like mineral sieved at Ø < 1 mm, pH 7, CEC = 6.5 cmol+ kg⁻¹) (Adani and Ricca, 2004). Sand and clay were mixed in a ratio of 9:1 (w/w). An artificial medium was chosen to avoid contamination from SOM (Almendros and Dorado, 1999) and does not represent a natural soil (Adani and Ricca, 2004) and needs future comparison with natural soil. Organic amendments were added to the soil by normalizing each biomass to the nonhydrolyzable fraction of the total organic carbon (TOC) (see below for analytical details) and named the total unhydrolyzable carbon (U-TOC).

An equal amount of this fraction (3 g kg⁻¹ dm⁻³) was added to each sample amended with different composts. This choice was based on previous research (Adani and Ricca, 2004) indicating that the U-TOC was the stable OM fraction remaining after a medium-term degradation process (6 mo). Moreover, the nonhydrolyzable OM fraction (the biologically protected OM) was shown to be the recalcitrant fraction that contributes, together with physically and chemically stabilized OM, to the stable soil OM (Six et al., 2002; Mikutta et al., 2006). Therefore, the ability of different organic matrices characterized by different carbon contents available to contribute to the stable OM of soil could be compared using this method.

Using 5-L pots, incubation tests were replicated twice for each sample. Each pot was filled with approximately 2.5 kg of soil (dry matter). Soils were wetted with water and kept at 60% (w/w) of the maximum water-holding capacity. Water content was corrected gravimetrically every 3 d. Pots were incubated for 180 d in a chamber at 26 ± 2°C under dark conditions. During the incubation tests, soils amended with composts at different levels of maturity—SRM (soil amended with RM), soil amended with fresh compost (SFC), and soil amended with evolved compost (SEC)—were sampled at time 0 (T0), 60 (T2), 90 (T3), 120 (T4), and 180 d (T6), respectively. Each sample weighed approximately 100 g, except for the last sample, which weighed 150 g and was formed by three subsamples taken from each replicate. Soils were vacuum-dried at 65°C and then used for analytical determination. Analyses were performed in triplicate.

**Organic Matter and Organic Carbon Fractionation**

The OM was fractionated into its macromolecules using the wood analysis method adapted for soil (Adani and Ricca, 2004). Three consecutive extractions were performed on soil samples. The fraction soluble in benzene/ethanol mixture (2:1 v/v) and 16.87 mol L⁻¹ ethanol (soluble lipids, waxes, soluble tannins, etc.) was determined. Analyses were performed in triplicate.
proteins, and part of fulvic acids) was solubilized by the first extraction step (fraction I); protein, hemicelluloses, and part of fulvic acids (fraction II) were solubilized in hot 0.94 mol L⁻¹ H₂SO₄ under reflux for 2 h (second step); and cellulose (fraction III) was solubilized in 13.50 mol L⁻¹ H₂SO₄ at 4°C for 24 h (third step). The lingo-humic complex (fraction IV), insoluble in any solvent, was left in the residual fraction (lingo-humic complex plus acid-insoluble ashes) and quantified by ignition at 650°C of an aliquot of this fraction.

Isolation of the lingo-humic complex was performed for RM, composts, and soils. Volatile solids and ash content for the three organic matrices were determined by ignition loss at 650°C (APHA, 1992).

Humic acid (HA) and unhydrolyzable humic acid (U-HA) extractions were performed as described previously. Humic acid was directly extracted from RM, compost, and soil samples, whereas U-HA was extracted for RM, compost, and soil samples from the lingo-humic complex obtained as described previously.

In brief, 2 g of dried compost (10 g for soils) was extracted in a 250-mL Erlenmeyer flask with 100 mL of 0.1 mol L⁻¹ NaOH and 0.1 mol L⁻¹ Na₃P₂O₇, solution at 65°C for 24 h under N₂ in a Dubnoff thermostatic bath at 100 oscillations min⁻¹ (Adani and Ricca, 2004). The sample was cooled to room temperature in a 150-mL centrifuge tube and centrifuged at 13,000 rpm for 20 min. The extraction was repeated by adding distilled water until the supernatant was clear. The supernatant solution (alkaline extract) (TE or U-TE) was filtered through a 0.45-μm Millipore filter. The insoluble humic fraction (HU or U-HU) was retained and washed until a neutral pH was obtained and then stored. The alkaline extracts were then acidified to below pH 1.5 and centrifuged at 5000 rpm for 20 min. The insoluble fractions (HA or U-HA) were washed with distilled water until a neutral pH was obtained and followed by an extraction with 2.82 mol L⁻¹ HF for 24 h at room temperature to remove inorganic constituents (silicates). Then, HA (or U-HA) was washed to remove excess of HF with distilled water until a neutral pH was obtained, vacuum dried at 60°C, and weighed. The ash content of all extracted fractions was less than 1%. The residual alkali-soluble/acid-soluble fraction was purified by chromatography using a polyvinylpyrrolidone column, obtaining two fractions: fulvic acid (FA) and nonhumified material (NH).

The alkali-soluble/acid-soluble fraction obtained starting from the lingo-humic fraction represented the U-NH, as purified by chromatography using a polyvinylpyrrolidone was omitted because FA is solubilized during OM fractionation (see Fraction I and II). All fractions obtained during the HA and U-HA extractions were purified by organic carbon determinations (Cia-vatta et al., 1990) obtaining total organic carbon (TOC), total extractable carbon (TEC), humic acid carbon (HAC), fulvic acid carbon (FAC), humin carbon (HUC), and corresponding unhydrolyzable-C. Nonhumified carbon (NHC) was calculated as [TEC – (FA + HAF)] and U-NHC as (U-TEC – U-HA).

Variations in the content of each single fraction during composting and soil incubation were calculated from the start and end absolute fraction contents (g) and reported as the relative increase or decrease (g kg⁻¹).

Diffuse Reflectance Infrared Fourier Transformed, ¹H–Nuclear Magnetic Resonance, and ¹³C–Nuclear Magnetic Resonance Spectroscopy

Diffuse reflectance infrared fourier transformed (DRIFT) spectra were recorded with a FT/IR-300 E JASCO spectrometer equipped with a Diffuse Reflectance attachment (Pike Technologies, Inc, Madison, WI). The samples, a 7-mg sample and 700 mg potassium bromide (FT grade; Aldrich Chemical Co, St. Louis, MO), were finely ground using a Wing-L-Bug agate ball mill for 10 min (Specamill-Gresby-Specac, Kent, UK). The spectra were acquired in the 4000 to 600 cm⁻¹ range with 4 cm⁻¹ resolution, and 100 scans were performed on each acquisition. A background spectrum of finely powdered potassium bromide was recorded using the same instrument settings. Peak assignments were made according to Adani and Ricca (2004). In brief, the bands at 2920 to 2850 cm⁻¹ were attributed to aliphatic chains and peaks at 1710 cm⁻¹ to the stretching of C=O of ester, ketons, and acid carboxylic. Bands at 1666 cm⁻¹ and 1542 cm⁻¹ were attributed to the stretching of C=O of amide (amide I) and to deformation of N–H bond and stretching of C=N in amide (amide II). The bands at 1600 cm⁻¹ and 1420 cm⁻¹ were due to the C=O stretching of aromatic rings. The peak at 1516 cm⁻¹ was typical of an aromatic structure (e.g., lignin) (C=C stretching). The peak at 1456 cm⁻¹ was attributed to C–H bond deformation in aliphatic chains. The bands at 1273 cm⁻¹ and 1215 cm⁻¹ were due to C–O bonds of phenol. The band at 1032 cm⁻¹ was attributed to C–O stretching in the polysaccharide, and the band at 840 cm⁻¹ was due to C–H deformation out of the plane (lignin structures).

¹H-NMR spectra were recorded using a Bruker AC 300 spectrometer at 300 MHz under homogated decoupling conditions, the HOD peak produced by water impurities and proton exchange being irradiated. The radio frequency level of HOD irradiation was optimized for each sample. The NMR samples were prepared by dissolving 10 mg of sample in 0.5 mL solutions of NaOD 0.5 mol L⁻¹. ¹H-NMR gave semi-quantitative data, using calculations of the typical area corresponding to three regions. These regions were 0.5 to 3.3 ppm (alkyl protons, protons attached to carbon by an α bond to an aromatic ring or carboxylic groups, Hₐ), 4.8 to 3.3 ppm (protons attached to carbon bearing oxygen or nitrogen), and 7.8 to 6.5 ppm (aromatic and olefinic protons, Hₐ).

¹³C NMR spectra were recorded using a Bruker AC 300 spectrometer at 75.43 MHz. The NMR samples were prepared by dissolving approximately 50 mg of HA-like or unhydrolyzed-alkali-soluble HA-like in 0.5 mL of NaOD 1.5 mol L⁻¹. A quantitative intensity distribution was achieved by using the inversion-gated decoupling method: 0.23 s acquisition time, 45° pulse, 2 s relaxation delay, line broadening at 20 Hz, and a total acquisition time of 48 h. All chemical shifts were quoted with reference to internal 4,4-dimethyl-4-silapentane sulfonic acid (Na-salt).

For a semi-quantitative approach, the ¹³C-NMR spectra were subdivided into six regions. The signals ranging between 190 and 165 ppm were attributed to carboxyl and amide carbons. Peaks between 165 and 145 ppm arose from C=O of phenols. Peaks between 145 and 110 ppm were due to aromatic
The decrease of volatile solids during composting indicated that degradation occurred during the whole composting process (Fig. 1). The composting process proceeded through the decomposition of the readily degradable organic fraction (cellulose, hemicellulose, proteins, and lipids; i.e., fraction I, II and III) (Almendros et al., 2000), whereas the ligno-humic complex (fraction IV) was concentrated, and, as expected, the ash content remained constant (Fig. 1). Composting was also found to affect the C fractions because these all decreased during composting (Table 1). In contrast, U-TOC was relatively stable during composting, decreasing by only 136 g kg\(^{-1}\) (Table 1). Because U-TOC represented the sum of all U-C fractions, the large decrease of the alkali-soluble fraction (U-TEC = U-HAC + U-NHC) registered during composting could only be explained by an increase of the alkali-insoluble fraction (i.e., U-humin C) (Table 1). These data contrasted with our previous findings that indicated during the composting process, soluble U-HA is formed from the insoluble U-HU (Genevini et al., 2003). A similar observation supporting our most recent findings has previously been reported (González-Vila et al., 1999). As demonstrated by our DRIFT spectra data (Fig. 2), composting also qualitatively modifies HA. When compared with HA extracted from FC and raw material (RM), the HA DRIFT spectra of EC indicated a decrease of aliphatic chains content and an increase of N-containing and lignin-derivates molecules (González-Vila et al., 1999). The increase of N-molecules was most likely due to molecules of a microbial origin (González-Vila et al., 1999).

Purification of the parent material before HA extraction removed the HA labile fractions and simulated the biological process. This confirmed our previous findings (Chefetz et al., 1999; Genevini et al., 2002a, 2002b; Adani et al., 2006b) (i.e., a reduction of aliphatic chains and N-containing molecules and an increase of lignin-derived molecules as revealed by the comparison of DRIFT spectra of HA and corresponding U-HA) (Fig. 2). The decrease of N-molecules seems to confirm their microbial origin in HA because this fraction could only be removed by a chemical treatment. Also, it could not be removed during the biological process due to the presence of OM-degrading microorganisms that continuously contributed through their biomass to enrich HA with N-containing molecules (González-Vila et al., 1999).

### Results

#### Composting Process

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### Table 1. Absolute organic carbon and unhydrolyzable organic carbon contents for the different carbon fractions during the composting process.

<table>
<thead>
<tr>
<th>Compost</th>
<th>TOC†</th>
<th>TEC</th>
<th>HAC</th>
<th>FAC</th>
<th>NHC</th>
<th>HUC</th>
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<tbody>
<tr>
<td>RM†</td>
<td>600.9 ± 11.5§</td>
<td>301.7 ± 3.7c</td>
<td>107.6 ± 1.9c</td>
<td>9.2 ± 0.2b</td>
<td>184.9 ± 4.2c</td>
<td>299.3 ± 12.1b</td>
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<tr>
<td>FC</td>
<td>414.1 ± 17.4b</td>
<td>133.9 ± 4.3b</td>
<td>56.4 ± 1.9b</td>
<td>2.9 ± 0.2a</td>
<td>74.5 ± 4.8b</td>
<td>277.5 ± 17.9b</td>
</tr>
<tr>
<td>EC</td>
<td>227.8 ± 16.4a</td>
<td>77.0 ± 3.7a</td>
<td>33.6 ± 0.1a</td>
<td>2.3 ± 0.5a</td>
<td>41.1 ± 3.8a</td>
<td>150.8 ± 16.8a</td>
</tr>
</tbody>
</table>

† FAC, fulvic acid carbon; HAC, humic acid carbon; HUC, humic carbon; NHC, nonhumified carbon; TEC, total extractable carbon; TOC, total organic carbon.

‡ EC, evolved compost; FC, fresh compost; RM, raw material.

§ Means followed in the same column by the same letter are not statistically different (p < 0.05) according to Tukey test.

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al., 1999). Unhydrolyzable HA extracted from the final stage of composting (U-HA from EC) was more aromatic and less aliphatic than U-HA from RM and FC (Table 3), indicating that U-HA was modified during composting. From $^{13}$C-NMR it can be seen that this modification was due to the loss of the major part of the alkyl-C (a decrease of 56%), passing from RM to EC. This was confirmed by calculating the absolute content of the different C-types by using data reported in Table 3 and Table 1. From this calculation it could be shown that alkyl-C accounted for more than 50% of the total C loss, followed by an aryl-C loss of 16.7%, a carboxyl-C loss of 16.3%, an O-methoxyl-C loss of 10.2%, and a N/O-alkyl-C loss of 3.4%. Phenol-C remained unchanged.

Consequently, the relative concentrations of phenol and aryl-C increased by 79.7% and 34.7%, respectively. This trend was further confirmed by the good correlation found between aryl-C and phenol-C vs. alkyl-C (alkyl-C vs. aryl-C: $r = -0.99$, $P < 0.01$; alkyl-C vs. phenol-C: $r = -0.97$, $P < 0.01$). These results suggest that a fraction formed mainly by the alkyl-C of the U-HA becomes more hydrolyzable as the biological process (composting) proceeds. The loss of this fraction from the U-HAC could also explain the decrease of U-HAC content in EC as previously shown by mass balance (Table 1).

### Soil Incubation

As a consequence of microbial activity, soil incubated with raw material (SRM) exhibited a TOC decrease after 6 mo of incubation (Fig. 3). A general decrease of all C fractions composing the TOC occurred during the incubation period (Fig. 4). In particular, TEC, HAC, FAC, NHC, and HUC decreased by 622, 707, 807, 577, and 497 g kg$^{-1}$, respectively. In contrast, recalcitrant carbon (U-TOC) remained relatively constant during the 6-mo incubation period (Table 4). This is important because it justifies the approach used in this experiment whereby we added an equal amount of U-TOC (the stable fraction of carbon over a medium-term incubation period), independent of the degree of compost evolution (Adani and Ricca, 2004), to each sample. We noted decreases in U-TEC, U-HAC, and U-NHC of 581, 607, and 539 g kg$^{-1}$, respectively, whereas U-HUC increased by 745 g kg$^{-1}$.

Soil amended with fresh compost exhibited a similar trend to SRM (Fig. 3) in that all carbon fractions decreased during incubation, but less than was the case in the other samples.
**Table 3. Integrated area (%) of different carbon types of humic acid (HA) and unhydrolyzable humic acid (U-HA) extracted from compost at different stages and incubated soil with composts at the start and end of the trials (13C-nuclear magnetic resonance).**

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<tr>
<td>HA-RM†</td>
<td>9.8</td>
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<td>16.5</td>
<td>5.9</td>
<td>7.2</td>
<td>57.2</td>
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<tr>
<td>HA-FC</td>
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<td>12.2</td>
<td>7.3</td>
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<td>HA-EC</td>
<td>12.4</td>
<td>10.1</td>
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<td>U-HA-RM‡</td>
<td>14.6</td>
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<td>U-HA-EC</td>
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<td>U-HA-EC-SRM</td>
<td>14.6</td>
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<td>29.4</td>
<td>5.1</td>
<td>10.6</td>
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<tr>
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<td>32.6</td>
<td>8.8</td>
<td>12.8</td>
<td>18.4</td>
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<td>U-HA-EC-FC</td>
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<td>10.4</td>
<td>11.7</td>
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<tr>
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<tr>
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<td>41.1</td>
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<td>12.2</td>
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</table>
| † EC, evolved compost; FC, fresh compost; RM, raw material.  
‡ Prefix U indicates unhydrolyzable fraction.  
§ Soil incubated with raw material (SRM), fresh compost (SFC), and evolved compost (SEC).  
¶ T0, T6: soil sampled after 0 and 6 mo of incubation.

Total organic carbon decreased by 484 g kg⁻¹, whereas TEC, HAC, FAC, NHC, and HUC decreased by 689, 577, 608, 373, and 358 g kg⁻¹, respectively. Unhydrolyzable TOC remained constant during the 6-mo incubation period (Table 4). In contrast, U-TEC, U-HAC, and U-NHC decreased by 611, 590, and 645 g kg⁻¹, respectively, whereas U-HUC increased by 471 g kg⁻¹ (Table 4).

Soil amended with EC showed a different trend with respect to SRM and SFC. Carbon fractions decreased, but this decrease was less pronounced than observed in SFC and SRM. Total organic carbon, TEC, HAC, and HUC decreased by 276, 55, 560, and 335 g kg⁻¹, respectively (Fig. 3), whereas NHC increased by 302 g kg⁻¹ (Fig. 3). Unlike the other two samples, U-TOC decreased significantly (~211 g kg⁻¹). Similarly, U-TEC and U-HAC decreased significantly by 351 and 500 g kg⁻¹, respectively, whereas U-NHC increased (428 g kg⁻¹) (Table 4).

After 6 mo of incubation of the amendments in artificial soils (sample T6), HA composition exhibited a modified DRIFT spectrum (Fig. 4). This indicated that incubation removed the labile fraction (i.e., lipids from HA for SRM and SFC and concentrated N-containing and lignin-derived molecules). In contrast, the DRIFT spectrum of HA extracted from the EC-amended soils (SEC) after 6 mo of incubation did not show an appreciable difference to that of HA extracted at time zero (T0).

These observations were confirmed by ¹H-NMR and ¹³C-NMR spectra of HA that changed significantly for SRM and SFC after 6 mo of incubation. The alkyl fraction decreased, and the O/N-alkyl and aryl/phenol fractions increased (Tables 2 and 3). On the other hand, SEC did not exhibit much variation (Table 3).

As demonstrated by the composting process and the DRIFT spectra (Fig. 4), chemical pretreatment of soils before HA extraction had an effect similar to biological processes (Fig. 3). The NMR data demonstrated that during composting of SRM and SFC, the alkyl fraction became more hydrolyzable, and phenol-C increased with prolonged incubation times (Table 3).

**Discussion**

Degradation processes during composting affect the HA content and its quality. The HA content decreases due to the degradation of labile organic molecules forming more recalcitrant (i.e., the unhydrolyzable) HAs (Chefetz et al., 1999). Therefore, HAs were formed by a labile fraction and a recalcitrant fraction that was preserved (Binachessi da Chuna-Santino and Bianchini Júnior, 2002). We have found that HA from more evolved compost contains more recalcitrant fraction (aromatic carbon) and less labile fraction (aliphatic carbon) than HA from RM and FC, indicating that HA composition is dependant on the composting duration.

Labile fractions consist of a free and a hydrolyzable-lipid fraction (free fatty acids and esterifies lipids) that account for approximately 67% (¹³C-NMR data) of total HA lost during composting (Genevini et al., 2002a). O/N-alkyl C contributed modestly to the composition of the labile fraction as its content increased during composting (see ¹³C-NMR data in the Results section). The degradation of HA-carbohydrates during composting is well documented in the literature (González-Vila et al., 1999; Almendros and Dorado, 1999). It is likely that the O/N-alkyl fraction of HA, for example the carbohydrates and proteins that are hydrolyzable, also contribute to the labile-HA fraction. However, this fact was probably masked by the production of newly formed alkali-soluble molecules of microbial origin that contributed to HA molecules (González-Vila et al., 1999). In addition, the strong decrease of the alkyl fraction determining a strong relative increase of the other
fractions may have contributed to this apparent nondegradation of the O/N-alkyl fraction. This may have also occurred for the aromatic fraction (aryl-C + Phenol-C) but is unlikely because the increase of this fraction was double that of O-alkyl-C, indicating a degradation of recalcitrant fraction.

The unhydrolyzable recalcitrant fraction of HA, the U-HA, is mainly composed of highly polymerized aromatic lignin-derived molecules (acid-resistant lignin) (Leary et al., 1986) and carbohydrates (Knöker and Lüdemam, 1995), probably forming an unhydrolyzable cross-linked network (e.g., alkali soluble part of the cell wall of plant material) (Adani and Ricca, 2004; Adani et al., 2006c; Adani et al., 2007). The U-HA was not stable during the composting processes. Our results suggested that the alkyl fraction contributed above all to the loss of the U-HA. This fact contributed to the relative increase of the aromatic-C content of the U-HA.

Composts in soils were degraded at a rate depending on their degree of evolution (Pascual et al., 1999), as RM was degraded to a greater extent than EC. The use of fresh material allowed for higher TOC and, more significantly, higher U-TOC contents at the end of the incubation. These data are not surprising because TOC dosed with RM and FC was much higher than that dosed with EC. On the other hand, the U-TOC results are unexpected because it was dosed with the same amount for the different samples. This suggested that the biological stability of the biochemically protected carbon (the U-TOC) depended on the amount of available-C for microorganisms. Therefore, the higher amounts of labile-C from samples with RM and FC allowed for the preservation of the more stable-C.

Humic acids were degraded at different rates depending on their content in the labile fractions (Almendros and Dorado, 1999). As composting removed the HA labile fraction, HA from EC showed the lowest rate of degradation when incubated in soil. Nevertheless, because of the higher level of HA dosed, soil amended with RM showed a higher HA content at the end of the incubation period. However, the final content of HA from FC and EC was similar. Humic acid degradation of SFC proceeded mainly by degradation of the alkyl fraction, whereas for SEC, the aromatic fraction was mainly degraded.

When the U-HA was considered, the two samples showed interesting results. Although at the end of the incubation period U-HA contents were the same, the joint decrease of U-HA and U-TOC registered for SEC suggested that U-HA was probably degraded. In contrast, SFC showed constant U-TOC values, suggesting that the U-HA fraction was not degraded but probably was insolubilized.

Humic acid from EC was expected to be more recalcitrant to degradation. However, we found that a prolonged stabilization/humification process influenced its degradability, as confirmed by the reduced presence of the labile fraction with respect to the other samples. Although this in itself is not a new finding because it was previously shown that HAs were less degraded when copious nutrients were present in the system (Filip and Kubát, 2001; Filip and Tesařová, 2004), the common belief has been that compost should be obtained after long composting periods because the formation of evolved humified material takes a long time (Chen and Inbar, 1993). However, our results indicate that the contribution of compost to soil humus depends on the compost U-HA content and on the presence of the labile fraction that is able to modulate...
the preservation of the recalcitrant fraction. Therefore, RM and FC contribute to soil humus better than EC due to the presence of the labile fraction. In addition, the labile fraction plays an important role in soil structure formation and as nutrient (Swift, 1991).

References


