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OF HUMIN IN A TROPICAL SOIL
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CONTRIBUTION OF HUMIC SUBSTANCES FROM DIFFERENT COMPOSTS TO THE SYNTHESIS OF HUMIN IN A TROPICAL SOIL

Claudivan Costa Lima(2), Eduardo de Sá Mendonça(3) & Asunción Roig(4)

SUMMARY

The contribution of humic substances of different composts to the synthesis of humin in a tropical soil was evaluated. Increasing doses (0, 13, 26, 52, and 104 Mg ha⁻¹) of five different composts consisting of agroindustrial residues were applied to a Red-Yellow Latosol. These composts were chemically characterized and ¹³C NMR determined and the quantity of the functional alkyl groups of humic acids applied to the soil as compost was estimated. Thirty days after application of the treatments, organic matter samples were collected for fractionation of humic acids (HA), fulvic acids (FA) and humin (HU), from which the ratios HA/FA and (HA + FA)/HU were calculated. The application of the composts based on castor cake resulted in the highest HU levels in the soil; alkyl groups of the HA fraction of the composts were predominant in the organic components added to the HU soil fraction.

Index terms: humic acids, fulvic acids, ¹³C NMR, alkyl groups, recalcitrance.

RESUMO: CONTRIBUIÇÃO DE SUBSTÂNCIAS HÚMICAS PROVENIENTES DE DIFERENTES COMPOSTOS ORGÂNICOS NA SÍNTESE DE HUMINA EM UM SOLO TROPICAL

Avaliou-se a contribuição de substâncias húmicas provenientes de diferentes compostos orgânicos na síntese de humina em um solo tropical. Para isso, foram adicionadas doses...
INTRODUCTION

In tropical and subtropical soils greatly impacted by weather events, organic matter (OM) is an essential element of the productive capacity, particularly concerning fertility (Bayer & Mielniczuk, 1999), and is believed to account for more than 90% of the cation exchange capacity (CEC) of soils in the tropical ecosystems (Melo et al., 1997). The maintenance or recovery of the productive capacity of these soils can be obtained with a management system that increases the OM contents, as by using composts.

The mineral nature of soil, as well as the quantitative and qualitative characteristics of the humic acids (HA) and fulvic acids (FA) in compost can affect both the reactivity and stability of these humic substances (HS) in the medium. (Mikutta et al., 2006, Lima et al., 2005, 2006). Carboxylic (-COOH) and phenolic (-OH) functional groups, which are usually found in the periphery of the HS (Fassbender, 1975), are related to their reactivity, contributing to the increase in CEC (Jerzykiewicz et al., 1999; Lead et al., 1999, Hayes & Malcom, 2001), whereas the hydrophobic functional groups are more directly related to the stability of HS in the soil. For being recalcitrant and adsorbed by the soil mineral matrix or occluded intra-aggregates (Mikutta et al., 2006), hydrophobic HS are more resistant to microbial degradation (Piccolo et al., 1999), and have greater residency in the soil. This stability is more effective in poorly crystallized soils containing aluminosilicates, such as allophone, imogolite (Torn et al., 1997) and also montmorillonite minerals, where adsorption apparently occurs within mineral lattices (Khan, 1946, 1950, 1959). Kaolinite, orthoclase, feldspar and microcline are adsorbed at the crystal surface and easily dissolved following alkali treatment (Khan, 1946, 1950, 1959). However, Wiseman & Pütmann (2005) found a relationship between C storage in non-allophanic soils composed of varied phyllosilicates, as well as Fe and Al oxyhydroxides.

The level of alkyl groups (paraffinic C) of HA present in the composts, which is similar to the soil fraction of soil (HU) (Stuermer et al., 1978; Hatcher et al., 1980; Preston & Riepeister, 1982), may control humin synthesis when these composts are applied, stabilizing OM in the soil. This is probably caused due to the interaction between the functional alkyl groups of HA that exist in the composts with the soil mineral matrix, which confers to these composts the characteristics of a “functional humin”. Conceptually, humin is defined as the fraction of humified organic matter insoluble in alkali or acid medium, and which is strongly adsorbed by the soil matrix (Stevenson, 1994). Humin has already been referred to as “anhydrous humic acids” by Odén (1919), as “humic clay-acid complex” by Khan (1945) and later reiterated by Kononova et al. (1966), due to the great similarity between the analytical properties and those of HA in terms of elementary composition, functional groups and infrared spectrum. Moreover, soil treatment with HF, which makes it possible to destroy clays, results in the solubilization of humin in alkali medium (Stevenson, 1994). Despite this fact, Hacher et al. (1985) found significant differences between humin and HA spectra in a 13C RMN analysis, in the same sample, suggesting that humin is not a clay-humic acid complex. Humin differs from HA in the soil in its higher polysaccharide content and lower relative proportion of carboxylic C, but mostly by the higher concentration of paraffinic compounds (alkyl groups) (Hacher et al., 1985).

The purpose of this study was to evaluate the contribution of humic acids of different composts derived from agro-inpowdersial organic residues to the synthesis of humin in a tropical soil.

MATERIAL AND METHODS

The study was carried out in a greenhouse of the Department of Soils of the Federal University of Viçosa, Viçosa, Minas Gerais State, Brazil. A sample of dystrophic medium-textured red-yellow Latosol, from the municipality of João Pinheiro, Minas Gerais, was collected in the 0–20 cm layer and sieved through...
2 mm mesh. This soil has the following chemical and physical properties: pH in water: 4.76; P-Mehlich-1: 2.1 mg dm⁻³; K: 122 mg dm⁻³; Ca: 0.18 cmol dm⁻³; Mg: 0.06 cmol dm⁻³; Al: 0.20 cmol dm⁻³; H + Al: 3.9 cmol dm⁻³; SB: 0.55 cmol dm⁻³; t: 0.75 cmol dm⁻³; T: 4.45 cmol dm⁻³; V %: 12.4; m %: 26.7; MO: 1.43 dag kg⁻¹; HA: 0.09 dag kg⁻¹; FA: 0.16 dag kg⁻¹; HU: 0.47 dag kg⁻¹; P-rem: 26.3 mg L⁻¹; soil density: 1.33 g cm⁻³; particle density: 2.68 g cm⁻³; clay content: 37 dag kg⁻¹; texture class: sandy clay loam.

Five composts were added to this soil (Tables 1 and 2). The humic substances (HA and FA contained in the referred composts were extracted and purified for characterization in a ¹³C NMR spectrum (Figure 1). To perform the ¹³C NMR analyses in solid state and "magic angle spin" (MAS) of HA and FA samples, a Varian 300 MHz equipment was used with zirconium oxide rotor operating at 75.42 MHz of C, at a rotation rate of 4 KHz, contact time 1.5 ms, acquisition time 35 ms, pulse width 6.7 μs, and the pulse angle 90º. Hexamethylbenzene was used as reference material.

The qualitative and semiquantitative interpretation of the RMN spectrum was based on Inbar et al. (1989), Kögel-Knabner (1997) and Ussiri & Johnson (2003). The spectrum was divided into five regions according to the chemical shift, as follows: C alkyl (0–110 ppm), aromatic groups (110–140 ppm), phenolic groups (140–160 ppm), carboxylic group (160–190 ppm) and carbonyl carbon of aldehyde/ketone (190–240 ppm).

Based on the HA contents of the composts (Table 2), for a dose of 13 Mg ha⁻¹, the quantitative HA levels (in kg), applied to the soil with compost were estimated. Also, based on the same criterion, the quantitative levels of alkyl functional groups of HA added to the soil were estimated, from qualitative and semiquantitative interpretations of the data obtained with ¹³C NMR spectroscopy.

The treatments consisted of five doses (0, 13, 26, 52 and 104 Mg ha⁻¹, dry matter base) of the five composts obtained, which were applied to the above soil, and this mixture was stored in 20 dm³ recipients. The treatments were arranged in a 5 x 5 factorial design, with three replications, and distributed in random blocks.

Soil samples were collected 30 days after compost application for fractioning of the organic matter in humic acids (HA), fulvic acids (FA) and humin (HU), according to the "International Humic Substances Society" - IHSS (Hayes et al., 1989), and the C content of these fractions was determined according to Yeomans & Bremner (1988). From these determinations the HA/FA and (HA + FA)/HU ratios were calculated (Benites et al., 2003).

The data obtained were subjected to descriptive statistics, variance analysis, simple linear correlation (Pearson’s Correlation), and the Tukey test at 5 % for the comparison of averages, using the SAEG (System of Statistic and Genetic Analyses) program of the Federal University of Viçosa (FUNARBE, 1993). Regression equations were also adjusted according to the compost doses applied.

### Table 2. Chemical characterization of composts developed from different materials and enriched with minerals

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>SC</th>
<th>AS</th>
<th>SM</th>
<th>FC</th>
<th>M-G</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH-CaCl₂[(1)]</td>
<td>7.24</td>
<td>6.28</td>
<td>7.50</td>
<td>5.88</td>
<td>5.76</td>
</tr>
<tr>
<td>C Total (g kg⁻¹)</td>
<td>142</td>
<td>162</td>
<td>85</td>
<td>278</td>
<td>310</td>
</tr>
<tr>
<td>N Total (g kg⁻¹)</td>
<td>8.9</td>
<td>11.2</td>
<td>10.3</td>
<td>18.7</td>
<td>27.9</td>
</tr>
<tr>
<td>Ratio C/N</td>
<td>15.9</td>
<td>14.5</td>
<td>8.3</td>
<td>14.8</td>
<td>11.1</td>
</tr>
<tr>
<td>HA (dag kg⁻¹)</td>
<td>0.98</td>
<td>0.79</td>
<td>0.92</td>
<td>1.89</td>
<td>2.87</td>
</tr>
<tr>
<td>FA (dag kg⁻¹)</td>
<td>0.68</td>
<td>0.32</td>
<td>0.49</td>
<td>1.77</td>
<td>1.22</td>
</tr>
<tr>
<td>HS (dag kg⁻¹)</td>
<td>1.66</td>
<td>1.11</td>
<td>1.41</td>
<td>3.66</td>
<td>4.09</td>
</tr>
<tr>
<td>LOM (dag kg⁻¹)</td>
<td>6.16</td>
<td>6.20</td>
<td>6.81</td>
<td>16.58</td>
<td>55.80</td>
</tr>
<tr>
<td>Ashes (dag kg⁻¹)</td>
<td>71.8</td>
<td>74.4</td>
<td>76.4</td>
<td>41.6</td>
<td>16.1</td>
</tr>
</tbody>
</table>

(1) Solid-solution relationship 1:5; HA: humic acid; FA: fulvic acid; HS: humic substances; LOM: light organic matter; SC: simple compost; AS: SC + NPK, where N: ammonium sulphate; SM: SC + powder from serpentinite and mica schist rocks; FC: filter cake + sugarcane bagasse; M-G: castor oil plant residue + sugar cane bagasse.

### Table 1. Composts formulated with different materials and enriched with minerals

<table>
<thead>
<tr>
<th>Compost</th>
<th>Composition</th>
<th>Proportion</th>
</tr>
</thead>
<tbody>
<tr>
<td>SC</td>
<td>Sugarcane bagasse + bagasse ash + manure from caged laying hen</td>
<td>3:3:2</td>
</tr>
<tr>
<td>AS</td>
<td>SC + NPK[(2)] (N: ammonium sulphate)</td>
<td>3:3:2</td>
</tr>
<tr>
<td>SM</td>
<td>SC + (micaschist + serpentinite)[(2)]</td>
<td>3:3:2</td>
</tr>
<tr>
<td>FC</td>
<td>Sugarcane bagasse + filter cake</td>
<td>2:1</td>
</tr>
<tr>
<td>M-G</td>
<td>Sugarcane bagasse + castor oil plant residue</td>
<td>2:1</td>
</tr>
</tbody>
</table>

[(1)] The compost was enriched with NPK in a way that the application of 20 Mg ha⁻¹ of the compost would contain the fertilizer dose recommended for sugarcane, which is, according to Korndörfer et al. (1999), 80 kg ha⁻¹ P₂O₅, 90 kg ha⁻¹ K₂O and 60 kg ha⁻¹ N. For this calculation, the initial volume of organic residues was considered to be reduced by 2/3 in the composted mixture.

[(2)] Addition of 62.5 kg Mg⁻¹ of the rock powder mixtures at a proportion of 1:1.
RESULTS AND DISCUSSION

Quantitative levels of alkyl functional groups of composts

The nature of the composted material and the mineral enrichment of the compost were found to have quantitative and qualitative influence on HS generation (Table 3), which corroborates results reported by Lima et al. (2005) and Bernal et al. (2009). These authors also stated that the addition of minerals contributes to accelerate the decomposition process of the organic matter of the compost, and at the same time favors the humification of organic matter.

Table 3. Estimates of the quantitative levels of humic acid fractions (HA) and fulvic acids (FA) and of alkyl groups of the HA fraction (in kg), contained in the 13 Mg ha⁻¹ dose, in the different composts

<table>
<thead>
<tr>
<th>Treatment</th>
<th>HA</th>
<th>FA</th>
<th>Alkyl groups of the HA fraction kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>SC</td>
<td>127</td>
<td>88</td>
<td>81</td>
</tr>
<tr>
<td>AS</td>
<td>103</td>
<td>42</td>
<td>58</td>
</tr>
<tr>
<td>SM</td>
<td>120</td>
<td>64</td>
<td>68</td>
</tr>
<tr>
<td>FC</td>
<td>246</td>
<td>230</td>
<td>139</td>
</tr>
<tr>
<td>M-G</td>
<td>373</td>
<td>159</td>
<td>273</td>
</tr>
</tbody>
</table>

Figure 1. ¹³C NMR spectra and carbon composition of humic acids (a) and fulvic acids (b) from composts developed from different materials and enriched with minerals. SC: simple compost; AS: SC + NPK, where N: ammonium sulphate; SM: SC + serpentineite and micaschist rock powder; FC: filter cake + sugarcane bagasse; M-G: castor oil plant residue + sugarcane bagasse.
All composts had higher HA than FA contents; the application of FC compost resulted in the highest quantitative levels of FA; and the M-G compost resulted in the highest quantitative HA levels. The composts that led to the greatest amounts of soil humic substances (HA + FA), in decreasing order, were: M + G > FC > SC > SM > AS.

The SC, AS, SM, FC and M-G composts had HA consisting of 64, 56, 57, 56 and 73 % of alkyl functional groups, respectively. Among these composts, the M-G compost is worth mentioning, in which the percentage of alkyl functional groups of HA of 237, 370, 301 was 96 % higher than in SC, AS, SM, and FC composts, respectively.

Humin fraction of the soil

The C contents of the HU fraction were higher than those of the HA and FA fractions in all treatments and tended to increase with increasing compost doses (Figure 2). As the composts do not contain humin, once the humic fraction is conceptually adsorbed in the soil mineral matrix (Stevenson, 1994), other organic fractions added to the soil through the compost may have incorporated functional characteristics of humin. This incorporation of composts to the humin fraction can be determined by the HA fraction of composts applied to the soil (Figure 1), once the HA content of the compost was found to be more correlated to the humin content (r = 0.84**) than to the HA content of the soil (r = 0.45**). This interpretation is reinforced by the observation of the 13C spectra obtained by means of NMR analysis of HA from the studied composts, which are highly similar to the humin spectra of the different soils obtained by several authors (Stuermer et al., 1978; Hatcher et al., 1980; Preston & Ripmeester, 1982).

A significant correlation was found between the estimate of the alkyl groups of HA in the composts and the humin content in the soil (r = 0.82**). The humin content of the M-G compost increased most (Figure 2). This result can be attributed to the alkyl groups of HA of the corresponding compost. The quantitative level of this functional group was found to be on average 251 % higher than of the other composts (Table 3). The mineral enrichment of AS and SM composts favored mineralization of the precursors of the alkyl groups (Figure 1), as demonstrated by the reduction of the relative abundance compared to SC (Baldock & Preston, 1995; Gressel et al., 1996; Dai et al., 2001). In the latter, preferential decomposition of other fractions, such as carbohydrates, and preservation of the alkyl groups has probably occurred, reflected in the soil humin contents.

The correlations of the HA alkyl groups of the different composts were evaluated by the estimates obtained for every studied dose. The alkyl groups of HA were positively correlated with soil humin at the doses 13 Mg (r = 0.91**), 26 Mg (r = 0.97**), 52 Mg (r = 0.88**) and 104 Mg ha⁻¹ (r = 0.59**). This synthesis was less significant when higher doses were applied, indicating that the increased level of alkyl groups of HA in the soil affects the effectiveness of the synthesis of the "functional humin". These results differ from the findings reported by Hacher et al. (1985) and are closer to the proposals of several authors, e.g., Odén (1919), Khan (1945) and Kononova et al. (1966) who considered humin to be a "humic clay-acid complex" in the beginning of the 20th century.
Compost application to the soil has generally promoted alterations in the ratios of the HA, FA and HU fractions, compared to those usually found in the soil (Table 4). These alterations may be caused not only by the quantitative characteristics of HS, but also by the quality of HA contained in the applied composts, which have incorporated the characteristics of functional humin. The HA and FA contents of the soil were found to be similar in all treatments, i.e., FA was predominant over HA (HA/FA < 1). In the FC treatment, at doses of 52 and 104 Mg ha⁻¹, this ratio was greater than in the other treatments, except in the soil without composts, indicating that the relative amounts of HA and FA in this soil type are more balanced. Although all composts have higher HA than FA contents (Table 2), a prevalence of FA was found after compost application to the soil, as observed by Canellas et al. (2001). This predominance can be attributed to the limited condensation reaction (Mendonça & Rowell, 1994). The soil texture and presence of oxyhydroxides may also interfere with the amounts of FA and HA. FA is most predominant in clay (Mendonça & Rowell, 1996) and dystrophic soils (Cunha et al., 2005).

The HA of the composts may interact with clay, forming very stable complexes (Sposito, 1989). This association may occur by means of HA hydrophobic groups, or else, the methylene groups derived from cutin and suberin (Mahieu et al., 1999; Nierop et al., 1999). This association can also be seen when the ionization of the acid functional groups of organic molecules is repressed, that is, when the pH values are in the PCZ (Sposito, 1989). In this case, the HA turns into the humin fraction, and a relative increase in the FA contents occurs, which consequently leads to a reduced HA/FA ratio.

The values of the (HA + FA)/HU ratio were lower than 1 in the SC, AS, SM and M-G treatments, indicating a greater proportion of HA adsorbed in the soil mineral fraction that became the humin fraction. This adsorption was most intense at the dose of 13 Mg ha⁻¹ in the M-G treatment. Only in the FC treatment, from the 52 Mg ha⁻¹ dose upwards, the (HA + FA)/HU ratio was greater than 1, unlike in the other treatments. This indicates that the organic fraction of FC-treated soil contains a higher proportion of the HA + FA than of the HU fraction, or that the quantitative HA and FA levels of this compost caused lower interaction of these fractions with the soil mineral fraction.

CONCLUSIONS

1. Following the application of increasing doses of different composts to the soil, the FA contents in the soil were higher than HA, even though the composts had higher HA than FA contents.

2. The incorporation of the organic compound to the humin fraction was guided by qualitative characteristics of the HA fraction present in the composts, more specifically by the alkyl group content of this fraction.

3. The application of the M-G compost resulted in a higher content of soil humin.

4. Further studies are needed to evaluate the effect of the incubation period of composts in the soil on the synthesis and stability of “functional humin”.

ACKNOWLEDGEMENTS

The authors wish to thank the CNPq and FAPEAL for the financial support for this research, as well as Dr. Ana de Godos de Francisco and Diego Martínez Pérez of SACE – Universidad de Murcia – Spain, for their help with the 13C NMR spectroscopy.

Table 4. Relation between the carbon contents in the form of fractions of humic acids (HA) and fulvic acids (FA) and between HA + FA and humin (HU), in a soil treated with increasing doses of different composts

<table>
<thead>
<tr>
<th>Treatment</th>
<th>HA/FA - Dose</th>
<th>(HA + FA)/HU - Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0(1) 13 26 52 104</td>
<td>0(1) 13 26 52 104</td>
</tr>
<tr>
<td>SC</td>
<td>0.44 b 0.35 ab 0.37 b 0.46 b</td>
<td>0.44 a 0.45 b 0.47 c 0.68 b</td>
</tr>
<tr>
<td>AS</td>
<td>0.60 a 0.32 b 0.36 b 0.40 bc</td>
<td>0.35 a 0.51 b 0.61 b 0.73 b</td>
</tr>
<tr>
<td>SM</td>
<td>0.56</td>
<td>0.44 b 0.38 a 0.38 b 0.43 bc 0.53</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.37 a 0.49 b 0.39 c 0.44 c</td>
</tr>
<tr>
<td>PC</td>
<td>0.65 a 0.36 ab 0.46 a 0.62 a</td>
<td>0.35 a 0.73 a 1.04 a 1.14 a</td>
</tr>
<tr>
<td>M-G</td>
<td>0.57 ab 0.32 b 0.30 c 0.34 c</td>
<td>0.18 b 0.54 b 0.65 b 0.58 bc</td>
</tr>
</tbody>
</table>

(1) The ratios between the HS at the 0 Mg ha⁻¹ dose were calculated from the HA, FA and HU contents originally found in the soil; averages followed by the same letter in the column did not differ by Tukey test at 5 %. SC: simple compost; AS: SC + NPK, where N: ammonium sulphate; SM: SC + powder from serpentinite and mica schist rocks; FC: filter cake + sugarcane bagasse; M-G: castor oil plant residue + sugarcane bagasse.
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