Source and biological response of biochar organic compounds released into water. Relationships with bio-oil composition and carbonization degree

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ABSTRACT

Water-soluble organic compounds (WSOCs) were extracted from corn stalk biochar produced at increasing pyrolysis temperatures (350-650°C) and from the corresponding vapors, collected as bio-oil. WSOCs were characterized by gas chromatography (semi-volatile fraction), negative
electron spray ionization high resolution mass spectrometry (hydrophilic fraction) and fluorescence spectroscopy. The pattern of semi-volatile WSOCs in bio-oil was dominated by aromatic products from lignocellulose, while in biochar was featured by saturated carboxylic acids from hemi/cellulose and lipids with concentrations decreased with decreasing H/C ratios. Hydrophilic species in poorly carbonized biochar resembled those in bio-oil, but the increasing charring intensity caused a marked reduction in the molecular complexity and degree of aromaticity. Differences in the fluorescence spectra were attributed to the predominance of fulvic acid-like structures in biochar and lignin-like moieties in bio-oil. The divergence between pyrolysis vapors and biochar in the distribution of WSOCs with increasing carbonization was explained by the hydrophobic carbonaceous matrix acting like a filter favoring the release into water of carboxylic and fulvic acid-like components. The formation of these structures was confirmed in biochar produced by pilot plant pyrolysis units. Biochar affected differently shoot and root length of cress seedlings in germination tests highlighting its complex role on plant growth.

**INTRODUCTION**

Biochar (BC) research has made consistent progress since it was proposed as a sustainable strategy for the abatement of greenhouse gases in terrestrial ecosystems. However, the ameliorating effect of BC in soil applications is highly dependent on its physical and chemical properties, in turn affected by production technology and biomass feedstocks. The definition of BC quality is therefore fundamental, and different criteria were proposed for its classification, like carbon content, aromaticity, and the presence of harmful chemical species such as heavy metals or polyaromatic hydrocarbons (PAHs). Apart from priority contaminants, the role of BC mobile organic compounds is being evaluated as potentially affecting its performance in soil. Volatile
organic compounds (VOCs) and water soluble organic compounds (WSOCs) were deemed responsible for the positive\textsuperscript{3} and negative\textsuperscript{4,5} effects on plants, microorganisms\textsuperscript{6} and aquatic organisms\textsuperscript{7,8}. BC labile carbon structures could also affect the composition of soil derived dissolved organic matter, in turn influencing soil ecosystem processes. The presence of organic species leached from BC was confirmed in soil application. Riedel et al.\textsuperscript{9} evidenced a compositional change in the molecular fingerprint of the organic matter released from a soil mixed with BC compared to untreated soil in column experiments. The amendment caused a marked reduction of the organic matter mobilization from the soil, but a net increase in the intensities of black carbon-type and lignin-type compounds was observed. Uchimiya et al.\textsuperscript{10} demonstrated the existence of polyaromatic moieties in the dissolved organic carbon (DOC) extracted from a BC-amended soil, attributed to unique structures of pyrogenic DOC. Vapors re-condensation and pyrolysis temperature were found to be of primary importance for the production of BC suitable for soil application\textsuperscript{5,8,11}. The contact of the pyrolysis vapors with the carbonized biomass inside the reactor and their removal is critical to prevent BC contamination\textsuperscript{5,8}. Temperature and residence time are crucial process parameters. At temperatures higher than 400°C a sharp decrease in VOCs adsorbed on BC was evidenced\textsuperscript{8,11}. High nitrogen flow can reduce the content of PAHs\textsuperscript{12,13}. The effect of process conditions on the composition of WSOCs has not been widely investigated, nonetheless WSOCs may play an important role in BC environmental impact due to their mobility in water. WSOCs were investigated by means of two dimensional GC\textsuperscript{8} and liquid chromatography\textsuperscript{3,14,15}, while ultrahigh resolution mass spectrometry (Fourier Transform Ion Cyclotron Resonance Mass Spectrometry FT-ICR-MS) revealed the presence of thousands hydrophilic species, non-detectable with other techniques\textsuperscript{8,9,16}. Fluorescence spectroscopy with Parallel Factor Analysis (PARAFAC) was used as rapid and sensitive technique to investigate its
aromatic fraction\textsuperscript{10,17–19}. These studies have noticeably increased our understanding on the chemical composition of BC WSOCs, however, the relationship with production parameters and especially with the composition of the pyrolysis vapors are poorly known. The present study was primary focused on the comprehensive characterization of BC WSOCs with spectroscopic, chromatographic and mass spectrometry techniques in relation to the composition of the water-soluble fraction of the pyrolysis vapors, condensed (bio-oils) during the pyrolyses for BC production. The linkage between WSOC patterns, BC bulk properties and their implications on seeds germination, could eventually shed light on the role of BC mobile organic compounds in the determination of its quality for environmental applications, possibly leading to the proposal of threshold levels.

**MATERIALS AND METHODS**

**Samples**

BC were produced from pelletized corn stalks at 350, 400, 450, 500, 550, 600, 650°C and characterized in a previous study.\textsuperscript{11} The vapors carried by the nitrogen flow at the outlet of the reactor presented temperatures ranging from 195°C (pyrolysis at 350°C) to 310°C (pyrolysis at 650°C), measured with a thermocouple. These values were considered sufficiently high to minimize vapors re-condensation in the part of the reactor where the biochar was synthesized. Vapors leaving the reactor were condensed in two ice/salt cold traps at -14°C to collect the pyrolysis liquids. The content of the two traps was merged to produce one bio-oil (OL) sample per temperature. In this study, the samples produced at the temperature $XXX$ °C are named $BCXXX$ and $OLXXX$.

**Lipid extraction from corn stalk biomass**
The total lipid fraction of the corn stalk biomass used in the pyrolysis experiments was determined by sequentially extracting the feedstock with CHCl$_3$-MeOH 2:1 (v/v) at 50°C for 1.5 hours (triplicate analysis). The profile of fatty acids was determined by GC-MS after methanolysis followed by the production of fatty acid methyl esters (FAME)\textsuperscript{20}.

**Extraction of biochar WSOCs**

An amount of BC was weighed (1 g ± 0.01 mg) into 20 ml vials and 10 ml of deionized water (DW, HPLC grade) was added. The vials were sealed with aluminum crimp seals with PTFE/rubber septa. The sealed vials were then placed on a mechanical shaker (IKA KS 260) covered with an aluminum foil, and left shaking at 150 rpm for 72 hours at ambient temperature. The resulting solutions were centrifuged at 3800 rpm for 10 min (ALC4232 centrifuge) to separate the solid material and filtered with PTFE syringe filters 0.45µm (Sartorius Minisart SRP) thereafter.

**Analysis of BC WSOCs by direct immersion (DI)-SPME-GC-MS**

Each BC extract (1ml) was added with 0.5 ml of 2M phosphate buffer (KH$_2$PO$_4$/Na$_2$HPO$_4$) at pH 5.7 in 1.5 ml vials. Carboxen-PDMS (Car-PDMS) SPME fiber was exposed to the solution under magnetic stirring for 30 minutes.\textsuperscript{4} The thermal desorption of the analytes and GC-MS analysis were performed with the method developed in a previous study\textsuperscript{11}. The amounts of WSOCs were expressed as peak area counts normalized by the sample weight (NA). Blank analyses of phosphate buffer and DW were performed to check procedural contaminations. A calibration curve of volatile fatty acids (VFA) was performed with a standard VFA solution (0.1% Sigma-Aldrich) containing acetic acid, propanoic acid, methyl propanoic acid, butanoic acid, 3-methyl butanoic acid and pentanoic acid in DW. Serial dilutions were prepared at 10, 5, 1 and 0.1 mg/l in phosphate buffer,
spiked with 2-ethyl butyric acid 5mg/l in DW (internal standard) and analyzed in triplicate. The concentration of each VFA was calculated using the response factors from the calibration curve and expressed as (µg/gBC).

**Analysis of BC WSOCs by ESI(-)FT-ICR-MS**

BC extracts were diluted 1:10 in methanol and analyzed by negative electrospray ionization (capillary voltage 4kV) on a Bruker solariX 12T FT-ICR-MS. 500 scans were acquired for each spectrum (syringe infusion, 200µL·h⁻¹) using an 8 MW acquisition size (broadband). Ion accumulation time was set at 0.8sec. Samples were spiked with ES tuning mix (Agilent) and a starting calibration list was developed from single point correction on the m/z 301.998139. Mass spectra were analyzed using Data Analysis software (Bruker Daltonics). Peaks were assigned with a signal to noise threshold of 4 and absolute intensity threshold of 2·10⁶. Calibration lists were developed over the m/z range 100-600 by Kendrick mass analysis. Mass spectra without the calibrant were recalibrated with a quadratic equation with a standard deviation < 100ppb (67 points). Calibrated mass lists were processed with PetroOrg software. Peaks were assigned with a threshold of 100ppb and molecular formulas within the range: C₁-100, H₄-200, O₁-20, N₀-4.

**Analysis of BC WSOCs by fluorescence spectroscopy and PARAFAC**

BC extracts were diluted in DW until the absorbance in the UV-Vis wavelength range 200-800 nm was <0.1²¹, recorded with a PerkinElmer λ650 spectrophotometer, using quartz cells with 1.0 cm optical path. Fluorescence excitation/emission matrices (EEMs) were acquired (duplicate analysis) on an Edinburgh Instrument F900 with excitation and emission wavelengths in the range of 220-500 and 280-600 nm respectively, both at 5nm intervals. Solutions of 16 EPA PAHs (Sigma-Aldrich, 1µg/ml), IHSS Suwanee River Fulvic Acid (SRFA, 1mg/ml), o-cresol and o-eugenol (0.1
mg/ml) in DW were analyzed under the same conditions. PARAFAC was performed on the EEMs corrected for instrument bias and non-trilinear signals, with N-way toolbox\textsuperscript{22}, drEEM tool for Matlab \textsuperscript{23}. The number of PARAFAC components was selected considering the Stoke’s shift, leverage values, analysis of residuals and core consistency diagnostic\textsuperscript{23,24}.

**Analysis of bio-oil WSOCs**

The OL samples were diluted 1:10 in DW and centrifuged (3800 rpm for 15 min) to precipitate the water-insoluble part, while the WSOCs were analyzed by DI-SPME-GC-MS, FT-ICR-MS and fluorescence-PARAFAC. An aliquot of 250\(\mu\)l was spiked with 150 \(\mu\)l of \(o\)-eugenol 10 \(\mu\)g/ml in DW, phosphate buffer and DW to a final volume of 1.5 ml. DI-SPME and GC-MS conditions were those used for BC. FT-ICR-MS was performed on solutions further diluted 1:100 in methanol. 500 scans were acquired for each spectrum using an 8 MW acquisition size (broadband). Ion accumulation time was set at 0.5sec. Peaks were assigned with a signal to noise threshold of 4 and absolute intensity threshold of \(2 \cdot 10^6\). Mass spectra without the calibrant were recalibrated with a quadratic equation with a standard deviation < 100ppb (89 points). Calibrated mass lists were processed with PetroOrg software. Peaks were assigned with a threshold of 100ppb and molecular formulas within the range: \(C_{1-100}, H_{4-200}, O_{1-20}, N_{0-4}, S_{0-2}\). Fluorescence-PARAFAC conditions were the same used for BC.

**Germination tests**

Seed germination tests on cress (*Lepidium sativum L.*) were conducted with BC in water suspensions at 40g/l (4 replicates) according to Rombolà et al.\textsuperscript{4}. A solution of acetic, propanoic, butanoic and 3-methyl butanoic acid in DW (98, 18, 7.6, 1.4 mg/l respectively) was also tested, alone and with BC650 at 40g/l. Fifteen seeds per Petri dish were sampled and seedlings elongation was measured (root and shoot lengths in cm). The following statistics were performed with the
software PAST (Paleontological Statistic vers. 2.16): Kruskal-Wallis test (non-parametric), one-way ANOVA (after data transformation with Box-Cox, to achieve normality of the distributions and homogeneity of the variance), post-hoc tests (Mann-Whitney and Tukey test)

**WSOCs of biochar from pilot plant pyrolysis units**

A set of five BC samples produced with pilot plant pyrolysis units were extracted with DW and WSOCs characterized by DI-SPME and Fluorescence-PARAFAC. Two reference BC from miscanthus straw (MSP550) and softwood pellets (SWP550) produced at 550°C with a 50 kg/h capacity unit were purchased from the UK Biochar Research Centre, University of Edinburgh. Three BC samples produced with PYREG and characterized in an interlaboratory ring trial (EU-COST Action TD1107)² were also investigated. These samples were produced from residues of wood chips production (BC1), a blend of paper sludge and wheat husks (BC2), and sewage sludge (BC3) at 620, 500 and 600°C respectively.²

**RESULTS AND DISCUSSION**

**Semi-volatile WSOCs (DI-SPME-GC-MS)**

The corn stalk BC and the corresponding OL presented noticeable dissimilarities in the patterns of semi-volatile WSOCs. Representative examples are reported in the chromatograms of Figure 1, while all the compounds detected in BC and OL are listed in Table S1 and S2 respectively. The series of peaks in BC350 extracts were predominantly associated to carboxylic acids, which were the main components of the low molecular weight fraction of BC WSOCs, with C₁-₁₂ straight-chain and branched, saturated and unsaturated aliphatic acids, and aromatic acids like benzoic acid and its C₁-₂ alkylated derivatives. The composition of the OL was more complex (Figure1), with 124 tentatively identified compounds in contrast with the 36 of BC350. OL profiles included primarily lignin markers (2-methoxy-, 2,6-dimethoxy- and C₁-₃ alkyl substituted phenols) and
typical degradation products of the cellulose and hemicellulose fractions of the parent corn stalk
(5-6 membered rings heterocyclic aldehydes ketones and diketones, C_{1-3} alkyl substituted and
hydroxyl substituted cyclopentenones, and furans). Only 5 lignin markers characterized BC
WSOCs out of the 15 of OL (phenol, C_{1-2} phenols, guaiacol and 4-methyl guaiacol). Their signals
became negligible in the BC produced above 500°C. However, traces of alkylated phenols were
observed in all the BC WSOCs, possibly indicating their stronger interaction with the aromatic
structure of the BC compared to the methoxylated homologues. Furthermore, WSOCs of BC
produced below 450°C featured 8 proxies of hemicellulose (furfural and methyl furfural,
benzaldehyde and hydroxyl benzaldehyde, 2-acetyl furan and C_{1-3} cyclopentenones) compared to
the 34 of OL. A series of compounds generated by the progressive carbonization of the biomass
inside the reactor during the pyrolysis were detected in all the OL, like monoaromatic
hydrocarbons (benzene, toluene and C_{1-5} alkylated derivatives), and low molecular weight PAHs.
None of these species characterized the BC WSOCs, but minor contribution of some of these
compounds was evidenced in other studies. Low molecular weight aliphatic aldehydes (C_{3-4}),
ketones and diketones (C_{4-6}) were detected in the OL, but not in BC WSOCs, indicating that, if
retained by the BC after their production, they could be released preferentially as VOCs. OL
composition included also nitrogen containing aromatic compounds deriving from the protein
fraction of the biomass feedstock (pyridines, pyrazines, aromatic nitriles quinolines and indoles).
Interestingly, BC WSOCs did not present any of these species, indicating an effective removal as
pyrolysis vapors or a stronger interaction with BC. Finally, VFA, C_{3-5} unsaturated and higher
molecular weight aromatic acids were present in the OL as free carboxylic acids but also in the
form of methyl esters, probably originated from the reactivity with methylating products (e.g.
methanol) at low pH. OL WSOCs lacked in the C_{4-12} and the methyl substituted homologues of
carboxylic acids, indicating their possible formation and preferential adsorption onto the BC surface during pyrolysis. However, it cannot be excluded that the mass spectra of the missing aliphatic acids, were covered by the dominance of other more intense signals from lignin. The formation of low molecular weight fatty acids during pyrolysis (acetic and propanoic) is associated with the thermal decomposition of the hemi/cellulose fraction. Nevertheless, the higher molecular weight fatty acids could form from the fragmentation of the parent corn stalk lipid fraction. The lipids accounted for 7.0 ± 0.7 % of the biomass dry weight. The pattern of FAME by GC-MS revealed a total of 14 compounds (Table S3), ranging from saturated (8:0-30:0) to unsaturated species (16:1, 18:1 and 18:2). Palmitic, stearic, linoleic and oleic acids were the principal constituents of the FAME in corn seeds, whose residues left in the field could contribute to the composition of the collected corn stalk. A net decrease of the fatty acids was observed in the WSOCs of BC with increasing carbonization degree, measured by the H/C atomic molar ratio. Trace amounts were released even by highly carbonized BC (H/C 0.32), while branched and C3-7 unsaturated homologues were typical of less carbonized ones (H/C 0.80-0.59). In accordance with their presence in the water extracts, fatty acids were also volatilized by BC in the form of methyl esters. Rombolà et al. evidenced the inhibiting activity of WSOCs of poultry litter BC on the germination of cress and VFA were the potential cause. Due to their mobility in air and solubility in water, VFA could play a considerable role in the agronomic/environmental performance of BC application to soil and their quantification could be useful for the determination of its quality. Total VFA concentrations decreased with the increasing BC production temperature from 3.0 ± 0.3 mg/g of CS350 to 35 ± 14 µg/g of CS650 and statistically significant correlations (r> 0.9, p<0.01) were observed between the values of each single and total VFA (Figure S1/TableS4) and the decreasing H/C values of the BC. This correlation is in line with the decreasing amount of VOCs. In
summary, the great majority of species detected in the OL WSOCs were not found in the water extracts of BC. While OL WSOCs featured mostly lignocellulosic derived pyrolysis products, BC was dominated by carboxylic acids from hemi/cellulose and lipids.

*Hydrophilic WSOCs (ESI-FT-ICR-MS)*

BC contamination could occur if pyrolysis vapors are not correctly swept from the reactor during its production.\(^5,8\) Given the divergent patterns of BC and OL WSOCs discussed in the previous section, the comparison was extended to the less-volatile components that could be detected by ESI(-)FT-ICR-MS. Because ionization with ESI is suitable for polar compounds with both acidic and basic functionalities,\(^26\) the fraction investigated was categorized as hydrophilic. The mass spectra of the OL WSOCs confirmed the complex composition evidenced in other studies on similar feedstocks,\(^27\) as molecular formula assignment allowed to identify up to 4000 peaks (Table S5 and S6). Oxygenated (C\(_n\)H\(_m\)O\(_x\)) and nitrogen (C\(_n\)H\(_m\)N\(_y\)O\(_x\)) species together accounted for more than 60% of the total intensity. Trace contribution of sulfur was observed. The C\(_n\)H\(_m\)O\(_x\) distributions were similar in all the OL, encompassing oxygen atoms in the range O\(_{1-16}\), with O\(_5\) and O\(_6\) as most abundant classes (Figure S2). Interestingly, the N\(_1\)O\(_x\) class, followed the same pattern with NO\(_6\) as most abundant group, while for the minor N\(_2\)O\(_x\) and N\(_3\)O\(_x\) classes, O\(_4\) and O\(_3\) species had the highest abundance (Figure S3). The number of identifiable peaks in the WSOCs of BC350 and BC400 was comparable to that of the OL (about 2000) but sharply decreased to 40 in BC650 (Table S7 and S8). In contrast to the dissimilarities evidenced in the GC detectable fraction, the distribution of WSOCs in BC and OL pictured by ESI presented common features, with C\(_n\)H\(_m\)O\(_x\) compounds as most abundant, followed by the C\(_n\)H\(_m\)N\(_y\)O\(_x\) distributions (Table S5 and S7). The same range of oxygen atoms characterized BC350 and BC400 with O\(_5\) as most abundant group, but from BC450 the distributions progressively shifted to the prevalence of O\(_2\) species.
The oxygenated species of BC WSOCs revealed a high bioactivity, as some carboxyl and hydroxyl functionalities were the main source of toxicity on algal growth\(^7,16\). Given their prevalence in both the BC WSOCs and the OL, the attention was focused primarily on C\(_{\text{cH}}\)H\(_x\)O\(_x\) compounds. Van Krevelen diagrams are useful to understand the nature of these species as the molecular formula assigned in the mass spectra can be compared to the major biochemical classes of compounds\(^28\). To highlight the considerable changes occurring in the WSOCs of OL and BC due to the pyrolysis temperature, Van Krevelen plots of the samples produced at 350, 450, 650°C are reported in Figure 2. The patterns of OL350, 450 and 650, suggest that the pyrolysis temperature did not affect OL composition. Contrarily, those of BC450 and BC650 were distinctly different compared to the corresponding OL, and the increasing BC production temperature caused a net decrease in the number of WSOCs. However, BC350 and OL350 were highly similar, with the series of O\(_x\) classes shifting towards higher values of O/C, as consequence to the increasing number of oxygen atoms. Linear regression of the data points in Figure 2 revealed two main pathways: series with an intercept of 2 and those aligning along the equation \(y=2x\). The first one is associated to species differing by units of CH\(_2\)\(_2\). Coherently, alkyl chain elongation was observed also in all the principal compound classes of the volatile and semi-volatile fractions of BC and OL WSOCs (organic acids class, aldehydes, ketones, phenols and mono-aromatic hydrocarbons), as evidenced in Figure S4, where Van Krevelen plots of BC350 and OL350 mass spectra by GC-MS were produced. The latter pathway is indicative of dehydration reactions. Noteworthily, BC350, BC400 and all the OL displayed a point with coordinates (1,2) and molecular formula C\(_6\)H\(_{12}\)O\(_6\), that could be tentatively attributed to glucose or one of its isomers. The dehydration pathway could play an important role in the formation of BC and OL WSOCs from the pyrolysis products of the cellulose/ hemicellulose, as many data points in Figure 2 fall in
the region conservatively attributed to carbohydrates (0.67<\text{O:C}<1.2, 1.5<\text{H:C}<2.4)^{29}. Differently, Kendrick mass defect analysis, revealed chain elongation of molecular formulas ascribed to guaiacols and syringols, which were confirmed by DI-SPME analysis. Therefore, the O_2 and O_3 species appearing in the region attributed to lignin structures 0.1<\text{O:C}<0.7, 0.7<\text{H:C}<1.5^{30}, could be assigned to higher molecular weight phenolic functionalities originated from the pyrolysis of the corn stalk lignin fraction. Similarly, those with higher oxygen content within the same range of H:C and O:C values could represent dimers, trimers or higher molecular weight homologues. Their presence in the BC WSOCs could led to the release of lighter monomers in water by photochemical degradation^{31}. Several peaks fell in the region indicative of lipids (1.6<\text{H:C}<2, 0<\text{O:C}<0.2^{28}), and especially the O_2 species can be correlated to analogues of the fatty acids composing the lower molecular weight WSOCs. Generally, the carbon numbers of all the OL WSOCs were comparable to that observed by Hertzog et al in the OL of a lignocellulosic material^{32}. Likewise, the values ranged from 5 to 35 in BC350 and BC400 (Figure 3), corroborating the similarity between the WSOCs of poorly carbonized BC and those of OL. Double bond equivalent (DBE), or degree of unsaturation is the number of rings and double bonds and can provide information on the aromaticity of the WSOC species. The DBE values ranged from 1 to 18 in BC350, BC400 and all the OL. For DBE>2 an increase in the carbon number was associated to an increase of the number of oxygens (Figure 3). In summary, BC350 and BC400 WSOCs resembled those of the OL, but for BC>450 the cellulose and hemi-cellulose derivatives disappeared (Figure 2 and S4), while lignin degradation products could be detected until 550°C (BC550). The trend was associated to a sharp decrease of the more aromatic species with DBE>10 (Figure 3). At higher pyrolysis temperatures (> BC550), WSOCs tended to have increased H:C and low DBE values
(<5) ascribable to organic acids, still detectable at the highest pyrolysis temperature (BC650) (Figures 2 and 3).

**Aromatic structures of biochar WSOCs (Fluorescence-PARAFAC)**

All the aqueous extracts of BC and OL exhibited fluorescence indicative of the occurrence of aromatic functionalities. Figure S5 reports the EEMs of the BC and OL WSOCs, and those of the standard compounds, that were acquired to qualitatively compare known chemical species with the aromatic structures recurring in biochar WSOCs. PAHs were selected for their high fluorescence even though detected only in traces in the chromatograms of the OL (Table S2), alkylated and methoxylated phenols (α-cresol and α-eugenol) as lignin derivatives, and IHSS-SRFA as model humic substance. A PARAFAC model with 4 components (C1-4) suitably represented the dataset (95% of the variance explained) and is reported in Figure 4. C1 and C2 presented excitation/emission maxima at 320/405 and 350/470 nm respectively. Fluorophores of the natural organic matter (NOM) are characterized by broad excitation/emission spectra, with representative peaks in the wavelength range of 300-370/400-500 nm. Zhongqui et al. characterized 13 IHSS standard humic substances (aquatic and soil derived humic and fulvic acids) with PARAFAC, and two components resembled C1 and C2, while the spectral characteristics of IHSS-SRFA in Mobed et al. showed the same peaks at 320/405 and 350/470. C3 and C4 featured maxima at 285/335 and 275/310 nm respectively. Similar peaks in NOM were associated to protein-like structures. The intensities of the PARAFAC components are reported in Table S9, and their relative percent contributions to the total signal of each sample are presented in Figure S6. The BC WSOCs were mainly composed by C1 and C2 and lacked in the C3 and C4 structures, that featured the OL. Overall, the total signal intensity of BC WSOCs sharply decreased from BC450 and likewise that of C1 and C2, even though were still detectable at 650°C (Table S9).
Contrarily, the total EEM intensity of the OL was not dependent on the pyrolysis temperature and showed values one order of magnitude higher than that of the BC (Table S9). The percent contribution of C1 sharply increased with the pyrolysis temperature in the BC WSOCs while C2 decreased accordingly. Similar trends were observed by Uchimiya et al.\textsuperscript{19} in which pyrogenic DOC of lignocellulosic and animal based BC were investigated: two peaks (310/420 and 350/470 nm) were attributed to polyphenolic pyrolysis products and aromatic humic-like compounds that decomposes above 350°C, with the first one increasing and the second one decreasing with the pyrolysis temperature. C3 presented a maximum in OL350 and decreased at higher temperatures, while C4 smoothly increased. A high contribution of C3-C4-like components was observed in non-completely pyrolyzed BC from sawmill waste feedstocks\textsuperscript{18} and the pyrolysis of lignocellulosic biomass with low nitrogen and sulfur content is known to produce phenolic species. Given the low nitrogen content in the BC (1%)\textsuperscript{11}, and their similarity with the EEMs of the lignin markers (Figure S5), C3 and C4 could be associated to phenolic-like species. However, it cannot be excluded that protein-like structures could contribute to C3, as several nitrogen-containing compounds were detected in the OL. Moreover, the standard PAHs solution showed a similar peak at 280/335 nm, that could be attributed the naphthalene or fluorene\textsuperscript{36}. Nevertheless, PAHs exhibited distinctively narrower peaks compared to the broader ones observed for C1 and C2. Noteworthily, C3 and C4 had lower emission wavelengths than C1 and C2. A red shift in the excitation/emission maximum can be associated to an increased aromaticity and higher molecular weight,\textsuperscript{17,18} therefore C3 and C4 presented a lower degree of aromaticity compared to C1 and C2. Similarly, Uchimiya et al.\textsuperscript{17} observed a component comparable to C2 (380/460 nm) that was associated to recalcitrant polyaromatic fraction substituted with carboxyl and phenolic functionalities, especially in low temperature BC (350-500°C). In summary, BC WSOCs were composed of fulvic-like structures
and depleted in the phenolic-like less aromatic functionalities C3 and C4. Interestingly C1 and C2 were also primary components of the OL suggesting that biomass pyrolysis could intrinsically produce aromatic NOM-like moieties.

**Effects on the germination of cress seeds**

Germination tests are simple bioassays to screen the potential phytotoxicity of BC in soil due to the presence of harmful compounds (metals, PAHs, carboxylic acids, phenols). Tests on cress seeds aimed to evaluate the performance of BC containing variable amounts of WSOCs in consequence to the pyrolysis temperature. The ratio BC/DW of 40 g/l was chosen as indicative of a relatively high load of BC in soil application (40 t/ha). Lower loads were considered less active, while higher values less realistic, and were not examined in this study. The germination rate with BC was not significantly different from the controls without BC ($p > 0.05$) and the average value of all the treatments was $97 \pm 2.5 \%$. In agreement with previous studies on corn stalk BC with high VOC content, WSOCs did not present inhibiting effect. Surprisingly, the seedlings emerged after the germination showed significantly longer shoots in all the BC treatments versus the controls ($p < 0.001$), and the values for the BC with greater WSOCs content (BC 350-500) were higher ($p < 0.05$) than those of the more carbonized ones (BC 550-650) (Figure 5). BC WSOCs could be involved in the enhancement of plant growth. Plant stimulants such as karrikins were recently detected in green waste BC and in the corresponding pyrolysis water, particularly KAR1 (3-methyl-2Hfuro[2,3-c]pyran-2-one), that induced longer shoot lengths of tomato and lettuce seedlings in germination tests. Interestingly, ESI(-)FT-ICR-MS mass spectrum of all the OL presented a peak at $m/z$ 149.02442 [M-H]$^-$ with molecular formula [C$_8$H$_5$O$_3$]$^-$, that could be associated to KAR1. However, the peak was not revealed in corn stalk BC WSOCs, probably because of the trace concentration reported in BC compared to pyrolysis water. Moreover, BC
WSOCs featured aromatic fulvic-like structures and humic substances were found to promote the growth of shoot and roots of plants at different stages.\textsuperscript{41} With regards to the semi-volatile fraction, a solution of VFA mimicking the concentrations detected in BC350 (3 mg/g\textsubscript{BC}) was tested without and with BC650 (this BC was almost deprived of VFA Table S4). Total VFA concentration of 125 mg/l (equivalent to 40 g/l load of biochar) exhibited the same effects of the control in the case of the solution with only VFA, and the same effect of BC650, when added to the corresponding biochar (Figure 5). These findings indicated that VFAs were not involved in the root and shoot growth. Contrary to the improved shoot growth, significantly shorter root lengths were observed in the BC versus the controls ($p < 0.001$), especially BC550-650 ($p < 0.001$) (Figure 5). Reduced root growth was observed by Buss et al.,\textsuperscript{38} who associated the inhibition to high pH and available K of biochar produced in the temperature range of 550-650°C. However, the same study reported also the inhibition of shoot growth.\textsuperscript{38} Lengthening of root in plants is connected to the exploration of soil for water and nutrients during drought periods.\textsuperscript{42} In Petri dish bioassays, where the water content is not a limiting factor and the nutrients are provided by the seeds, BC could have possibly promoted the investment of carbon in the shoots rather than the roots. This would explain the improved shoot and reduced radical length for all BC. A similar effect was observed also on maize seedlings with maize BC produced at 450°C.\textsuperscript{43} However, the inhibition of root growth due to inorganic components or pH, cannot be excluded, and other factors could contribute to the improved shoot lengths, like the provision of additional nutrients by the BC.

\textit{Implications for biochar environmental applications}

In this study, it was shown that after three days in water at r.t, even highly carbonized BC released WSOCs. Interestingly, in a leaching study lasted for 17 days, most of the BC WSOCs were released within the first 3 days of the experiment.\textsuperscript{18} Thus, under environmental conditions, a wide
array of compounds can contribute to the pool of natural organic matter in soil. Overall, SPME-GC-MS, ESI(-)FT-ICR-MS and fluorescence-PARAFAC indicated that the release of WSOCs from BC was strongly reduced above 450°C, in agreement with the trend observed for VOCs, that began to decrease above 400°C.\textsuperscript{11} The investigation of OL WSOCs revealed original clues about the formation and release of those from BC. Organic acids were the main semi-volatile components released in water, suggesting that the more abundant OL components of lignin were strongly adsorbed onto the BC matrix or efficiently volatilized during pyrolysis. Given the porous structure of biochar, pores could be accessible by water solutions and the retention of phenolic compounds could possibly occur due to a hydrophobic effect, or by π-π interactions,\textsuperscript{44} that become more pronounced as the matrix gets more carbonized\textsuperscript{45}. Previous studies categorized biochar water-extractable organic compounds into classes used to describe NOM and evidenced that low molecular weight acids were important species even in BC produced at higher temperatures, while humic acids and low molecular weight neutral species were the principal components of the lower temperature BC (<450°C)\textsuperscript{14,46}. The higher molecular weight and aromatic structures of BC WSOCs were comparable to the species recurring in NOM. In D’Andrilli et al.\textsuperscript{33} the standard IHSS-SRFA was dominated by C\textsubscript{c}H\textsubscript{c}O\textsubscript{x} species by ESI(-)FT-ICR-MS and likewise BC and OL WSOCs of this study, that displayed similar structures in the lignin region of the Van Krevelen diagrams, but unique formulas in that of carbohydrates and lipids. Besides, the principal mass spacing patterns in the mass spectra (ESI-FT-ICR-MS) were alkyl chain elongation (14.01565 Da) and substitution of CH\textsubscript{4} versus O (0.0364 Da)\textsuperscript{47} in both SRFA and WSOCs. Fluorescence EEMs further confirmed the fulvic-like nature of the BC WSOCs, that was composed of labile (C1) less aromatic, and more recalcitrant polyaromatic (C2) structures substituted with carboxyl and hydroxyl groups, in agreement with previous hypotheses.\textsuperscript{10,17,48} WSOCs were extracted from five
BC produced by pilot plant continuous pyrolysis units. The profiles of semi-volatile and aromatic WSOCs are reported in Figure S7 and S8, respectively. BC1, 2 and 3 did not present detectable peaks associated to semi-volatile species. These BC were stored outdoors for several days after production and the more volatile components, if present, could have been lost. MSP550 and SWP550 BC, sealed upon production to prevent contamination, presented trace intensities of acetic acid, in line with the carboxylic acids released in water by corn stalk BC produced at the same temperature (BC550). However, the patterns of semi-volatile WSOCs were markedly reduced compared to BC550 (Figure S7), indicating that some vapor re-condensation could have occurred in the bench scale reactor. Interestingly, the EEMs of the WSOCs extracted from all the commercial BC revealed fluorophores resembling the aromatic fulvic-like structures C1 and C2 (Figure S8). This finding suggests that even at the pilot plant scale these aromatic units are formed probably by the interaction between the pyrolysis vapors and BC and survived into the pores. In conclusion, WSOCs influence the suitability of BC for environmental applications. Previous studies proposed that high amounts of PAHs\textsuperscript{49}, phenolic and carboxylic acids\textsuperscript{4,8} in BC WSOCs caused harmful effects on cress seeds. In this study, fulvic-like WSOCs and concentrations of VFA< 3mg/g induced statistically significant positive effects on the seedlings of cress, corroborating the hypothesis that the complex biological effects of BC WSOCs are the results of an interaction between contrasting factors.\textsuperscript{3,40} However, this study demonstrated that BC to soil application can be sustainable when BC contamination (organic and inorganic) is limited.

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Aromatic fulvic-like

Aliphatic fatty acid-like

350°C → 650°C
**FIGURES**

**Figure 1**: Total ion chromatograms of BC350 and OL350 WSOCs after DI-SPME-GC-MS analysis. Principal compounds are evidenced, while the complete lists of the volatile and semi-volatile WSOCs are reported in Tables S1 and S2.
Figure 2: Van Krevelen diagrams of WSOCs by ESI(-)FT-ICR-MS, in BC produced at 350, 450, 650 °C and the corresponding OL
Figure 3: Plots of DBE vs Carbon number of BC 350, 450, 650 WSOCs and the corresponding OL
**Figure 4:** Spectral characteristics of the four components (C1-4) PARAFAC modeling of BC and OL WSOCs
Figure 5: Box and whisker plots of shoot and root lengths (cm) of the cress seedlings for each treatment with (BC), without BC (Control) and VFA solution. 25-75 percent quartiles are drawn using a box. The median is shown with a horizontal line inside the box. Horizontal lines outside the box represent the variability outside the lower and upper quartiles. Values >1.5 times the box height are reported as circles.
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SUPPORTING INFORMATION

Figures and Tables with detailed quantitative, semi-quantitative and qualitative results of the analysis of BC and OL WSOCs by DI-SPME-GC-MS, ESI-FT-ICR-MS and Fluorescence-PARAFAC.