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Organo-mineral associations largely contribute to the stabilization of century-old pyrogenic organic matter in cropland soils



Victor Burgeon^{a,*}, Julien Fouché^{a,b}, Jens Leifeld^c, Claire Chenu^d, Jean-Thomas Cornélis^a

^a TERRA Teaching and Research Centre, Gembloux Agro-Bio Tech, University of Liège, Gembloux 5030, Belgium

^b LISAH, Univ Montpellier, INRAE, IRD, Institut Agro, Montpellier, France

^c Climate and Agriculture Group, Agroscope, Reckenholzstrasse 191, 8046 Zurich, Switzerland

^d UMR Ecosys, Université Paris-Saclay, INRAE, AgroParisTech, 78850 Thiverval-Grignon, France

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ABSTRACT

Understanding the processes underlying carbon (C) stability in soils is of utmost importance in the context of climate change. In this setting, biochar is often studied for its persistence in soils and reported to have positive impacts on soil fertility. Whilst recent research has mostly focused on the short-term effects of biochar amendments to soil, a better understanding of its long-term persistence in soils is needed.

Our study focuses on agricultural soils enriched in charcoal residues, produced in preindustrial kiln sites ca. 220 years ago, as a proxy for aged biochar. Our aim is to better understand the processes governing the long-term persistence of pyrogenic organic matter (PyOM) in cultivated soils. To achieve this, we focus on i) the effect of PyOM on soil aggregation, ii) its distribution amongst soil fractions and iii) its chemical and thermal properties. For this purpose, we combined a soil particle size-density fractionation with elemental and thermal analyses on topsoil samples collected in charcoal enriched (CHAR) and adjacent reference (REF) soils in a conventionally cropped field in Wallonia (Belgium).

The presence of charcoal in soils resulted in a 91 \pm 34% higher C content in CHAR soils than REF soils, of which 84 to 94% was PyOM and 6 to 16% was additional non-PyOM. In CHAR soils, macroaggregation was promoted at the expense of microaggregation (CHAR_{MACRO} = 48.9 \pm 12.8; CHAR_{MICRO} = 25.5 \pm 8.3 g 100 g⁻¹ soil) whereas these were similar in REF soils (REF_{MACRO} = 41.3 \pm 9.5; REF_{MICRO} = 36.0 \pm 8.7 g 100 g⁻¹ soil). Elemental and thermal analyses revealed that PyOM did not only occur as free light fractions but was also occluded in aggregates or sorbed onto mineral phases (56.4 \pm 22.9% of total PyOM) suggesting PyOM may be further stabilized through organo-mineral associations. Century-old PyOM showed increased H:C and O:C atomic ratios compared to recently pyrolyzed OM, particularly in occluded as opposed to free light fractions, vouching for the functionalization of its surfaces. Furthermore, regardless of high C contents in CHAR soils, similar C:N ratios between studied soils in mineral dominated fractions suggested increased amounts of N-rich non-PyOM in charcoal rich soils. Our results demonstrate that, over centuries in cultivated soils, PyOM plays an active role in aggregation patterns. We conclude that century old PyOM is broken down from coarse to fine particles, and through functionalization of its surfaces contributes to organo-mineral associations. Beyond its intrinsic chemical recalcitrance, the long-term persistence of biochar is enhanced by occlusion and sorption processes.

1. Introduction

Globally, human land use and agriculture in particular have resulted in a substantial loss of soil organic C (SOC) (Sanderman et al., 2017). As soil organic matter (OM) plays a crucial role in both the global C cycle and in ensuring food security (Le Quéré et al., 2009; Chenu et al., 2019) implementing practices that allow for SOC restoration and sequestration in cultivated soils offers a great potential to contribute to the mitigation of anthropogenic greenhouse gas emissions, whilst improving soil agronomic and ecological functions (Lal, 2004; Minasny et al., 2017; Chenu et al., 2019).

Amongst the various strategies allowing for SOC sequestration (i.e., a

* Corresponding author.

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E-mail addresses: victor.burgeon@uliege.be (V. Burgeon), julien.fouche@supagro.fr (J. Fouché), jens.leifeld@agroscope.admin.ch (J. Leifeld), claire.chenu@inra. fr (C. Chenu), jtcornelis@uliege.be (J.-T. Cornélis).

net removal of atmospheric CO₂), the study of C forms displaying long residence times in soils such as biochar has received much attention. Biochar is a form of pyrogenic organic matter (PyOM) whose purpose is to be amended to soils for its conjoint potential in terms of long-term C storage (centennial or longer) (Forbes et al., 2006) as well as soil fertility improvement (Lehmann et al., 2006). Meta-analyses of short-term (annual to decadal) biochar impacts in agro-ecosystems report an overall increased plant productivity as explained by the increase in soil water holding capacity, plant nutrient availability and liming effects (Jeffery et al., 2011; Biederman and Harpole, 2013). Such changes are particularly marked in highly weathered, coarse textured, tropical soils (Crane-Droesch et al., 2013). Biochar properties and their resulting agronomic benefits depend on its organic precursor and the pyrolysis conditions (e.g., heating rates and maximum temperatures) (Zimmerman et al., 2011; B,P. Singh et al., 2012; Baveye, 2014; Fang et al., 2015). Beyond reported positive impacts on soil fertility, biochar is studied for its potential as a C sequestration tool and related implications on the global C cycle (Schmidt and Noack, 2000). The stability of biochar and PyOM in general is attributed to their aromatic moieties which confers them recalcitrance against biotic or abiotic oxidation (Hammes et al., 2008; Kuzyakov et al., 2014). The distribution and organization of these aromatic moieties in charcoal fragments are characterized by the aromaticity and degree of aromatic condensation of biochar (Wiedemeier et al., 2015) which will in turn determine its persistence in ecosystems (Hammes et al., 2008; Kuzyakov et al., 2014). Despite this persistence in soils, biochar properties evolve in soils over time through breakdown and functionalization (i.e., ageing) (Cheng et al., 2006; Nguyen et al., 2008; Cheng and Lehmann, 2009; Hardy et al., 2017b). Recent studies have shown that PyOM may decompose faster than initially predicted (Hamer et al., 2004; Hammes et al., 2008) and that the persistence of PyOM in soils may have been overestimated particularly in arable settings (Lehndorff et al., 2014; Lutfalla et al., 2017; N. Singh, 2012). The persistence of PyOM, as for other OM forms, in soils is therefore strongly dependant on ecosystem properties rather than solely inherited chemical recalcitrance (Schmidt et al., 2011; Lehmann and Kleber, 2015; Lehmann et al., 2020). Therefore, spatial inaccessibility for decomposer organisms (i.e., occlusion in aggregates) and interactions with mineral surfaces (i.e., adsorption) or metals (von Lützow et al., 2006; Kögel-Knabner et al., 2008) need to be carefully investigated as stabilization processes for PyOM in soils.

While short-term biochar-induced changes in agro-ecosystems have been extensively characterized (Jeffery et al., 2011; Biederman and Harpole, 2013; Crane-Droesch et al., 2013), insights into its long-term effects are needed to verify its potential in climate-smart agriculture (Baveye, 2020). Worldwide, various field models have been addressed to better understand the long-term fate of biochar in soils. Amongst these are the fertile Amazonian Dark Earth (Glaser and Birk, 2012), soils containing wildfire derived PyOM (Schmidt and Noack, 2000; Bird et al., 2015), historical industrial sites (Cheng et al., 2008b), crop residues burning or relic charcoal hearths sites (Borchard et al., 2014; Lehndorff et al., 2014; Hernandez-Soriano et al., 2016b; Kerré et al., 2016; Hardy et al., 2017a; Hirsch et al., 2017; Schneider et al., 2019). All these studies address aged PyOM however varying production environments may result in a final product differing from that obtained through oxygen regulated pyrolysis as is done today to produce biochar (Bird et al., 2015).

In southern Belgium, pre-industrial charcoal production kiln sites are easily spotted in agricultural fields as "black patches" (Hardy and Dufey, 2015). The charcoal still found in these sites stands as left-overs from the production kiln that has been incorporated in agro-ecosystems since their conversion from woodlands *circa* 150–200 years ago (Hardy et al., 2017a). Due to similar oxygen-limited conditions in kilns as those obtained in modern biochar production units, this setting is ideal for the study of century-old biochar in temperate conventional cropping systems (Hardy et al., 2017a). Recent studies investigating similar kiln sites in the region showed that black patches stored 60 to 80% more C than adjacent reference soils in cropped systems, of which from 25% to 40% was non-pyrogenic organic matter (non-PyOM) (Hernandez-Soriano et al., 2016a; Kerré et al., 2016; Hardy et al., 2017a). The total N content as well as the C:N and C:P ratios are higher in kiln soils than reference soils (Hardy et al., 2017a). In addition, kiln soils display greater cation exchange capacity (CEC), which was associated with larger content of exchangeable Ca^{2+} and Mg^{2+} (Hardy et al., 2017a). Long-term ageing and continuous cultivation have resulted in the increase in H:C and O:C ratios of charcoal particles in comparison to freshly pyrolyzed OM (Hardy et al., 2017b). These modifications are associated with the oxidation of charcoal particles and the creation of functional groups (mostly carboxyl, phenol or carbonyl) at the edge of aromatic rings (Lehmann et al., 2005; Cheng et al., 2008a) which in turn promotes the precipitation of 2:1 phyllosilicates and CaCO₃ on the surfaces of PyOM (Hardy et al., 2017b).

In the present study we aim to investigate the various associations between century-old biochar and soil minerals in order to assess their contribution to PyOM persistence in soils. For this purpose, we separate SOC into pools displaying similar stabilization mechanisms (i.e., chemical recalcitrance, spatial inaccessibility, interactions with mineral surfaces) (von Lützow et al., 2006; von Lützow et al., 2007). Numerous physical (size, density) and chemical (oxidation, extraction) fractionation protocols have been developed to isolate SOC pools of comparable stabilization mechanisms and turnover rates (von Lützow et al., 2007; Poeplau et al., 2018). In soils, the most labile SOC pool, displaying turnover rates inferior to 10 years, are represented by free microbial biomass and the coarse light fraction (LF - density < 1.6-2 g cm⁻³) whereas the SOC pools with the highest turnover rates comprise the oxidation-resistant SOC pool and the heavy fraction after macroaggregate dispersion (HF - density $> 2.8 \text{ g cm}^{-3}$) (von Lützow et al., 2007; Virto et al., 2010; Poeplau et al., 2018). In cultivated soils, macroaggregates display rapid turnover as revealed by their composition of a mix of young and older SOC originating from both labile organic matter, (i. e., plant roots, mucilage, exudates, fungal hyphae, microbial biomass) (Rillig, 2004; Rasse et al., 2005) and more persistent physically protected and adsorbed organic compounds (Moni et al., 2010; Poeplau et al., 2018). Their dispersion is therefore required to isolate SOC groups with similar turnover rates (Virto et al., 2010; Poeplau et al., 2018). The few fractionation studies performed on soils enriched in aged biochar showed that the century-old biochar particles were mostly present as free particulate organic matter (i.e., non-associated with mineral particles) (Glaser et al., 2000; Herath et al., 2014; Kerré et al., 2016; Grunwald et al., 2017). However PyOM has also been reported to increase aggregate formation (Awad et al., 2013) and stability (Pituello et al., 2018) as well as to increase the abundance of non-PyOM in the aggregate-protected silt and clay fraction (Liang et al., 2010; Hernandez-Soriano et al., 2016a; Kerré et al., 2017) thus showing interactions with mineral phases.

Although the effects of PyOM in soils have been widely addressed, there is a lack of consensus concerning the processes involved in its longterm persistence. Nonetheless it is known that ageing in soils causes an oxidation of the charcoals surfaces and results in the formation of phenolic and carboxylic functional groups along with a charging of the structures surfaces (Cheng et al., 2008a; Nguyen et al., 2008; Cheng and Lehmann, 2009). As a result, physical protection through aggregation (Brodowski et al., 2006) as well as an increased association with the mineral phases (Nguyen et al., 2008; Rodionov et al., 2017; Hardy et al., 2017b) could play an important role in PyOM stabilization (Lehmann and Kleber, 2015) beyond its chemical recalcitrance to oxidation.

In this study, we investigated the long-term persistence of PyOM in soils. To achieve this, soil samples from five preindustrial charcoal kiln sites were compared to five adjacent reference soils in a conventionally farmed Luvisol (Isnes, Belgium). These charcoal kilns are visible as black spots on bare soils and procure an ideal setting for the study of aged biochar in conventionally cropped systems. This site hence serves as a proxy to study the long-term effect of biochar in soils if it was to become a common soil amendment in temperate soils of the region.

Our objective is to understand how and where PyOM persists in the soil matrix, as either free standing particulate OM (POM) or protected through aggregate occlusion or mineral sorption. In particular, this study aimed i) to quantify the effects of long-term PyOM accumulation on soil aggregation through a size and density fractionation protocol, ii) to investigate the distribution of PyOM amongst the fractions and iii) to determine the physico-chemical and thermal characteristics of organic matter in soil fractions. Numerous analytical techniques exist to identify, characterize and quantify PyOM; physical density separations, chemical oxidation, thermal analyses, optical or molecular marking techniques (Schmidt and Noack, 2000; Brodowski et al., 2005; Hammes et al., 2007; Leifeld, 2007; Plante et al., 2009; Bird et al., 2015; Kerré et al., 2016). Among these, thermal analyses and differential scanning calorimetry (DSC) in particular, has been reported to be an efficient way to characterize number of samples rapidly within the whole OM continuum, and was hence used to characterize PyOM forms (Leifeld, 2007; Plante et al., 2009).

We postulate that charcoal is found in all, light and heavy, soil fractions as a consequence of particle physical breakdown and functionalization of its surfaces. Furthermore, we postulate that the elemental composition of OM controls its thermal stability within the different fractions. Finally, we postulate that pyrogenic C forms may be further stabilized as a result of organo-mineral associations which increases its potential as a C sequestration tool.

2. Materials and methods

2.1. Study site

In cropped agroecosystems of the province of Namur (Belgium), century-old charcoal kiln sites are easily spotted visually in the field or by aerial imagery as darker ellipses on the soil surface (Fig. 1). The site selected for this study is located in Isnes ($50^{\circ}31'N 4^{\circ}44'E$) and was



Fig. 1. Aerial view of the study site. Kiln sites are visible as black patches when the soil is bare. The image comes from the "Service Public de Wallonie: Orthophotos 2009/2010", arrows indicate the approximate location of the five soil sampling sites per studied soil.

converted from forest to cropland at least 170 years ago as revealed by land occupation maps of Vandermaelen (1846–1854). Charcoal was produced in kiln sites during the pre-industrial era to meet energy demands thus providing an estimated age for the charcoal of interest of ca. 250 years (Hardy et al., 2017a). Hardy et al. (2017b) thoroughly characterized kiln sites of the region and reported that charcoal mixtures are generally dominated by deciduous hardwood such as oak (Quercus sp.), hornbeam (C. betulus), beech (F. sylvatica), hazel (Corylus avellana L.) and birch (Betula sp.).

The soil of the studied site has a homogeneous silt loam texture throughout and is slightly sloped (Table 1). It is characterized as a Haplic Luvisol (WRB, 2014) developed from decarbonated quaternary loess. The average annual precipitation is of 840 mm of rain and the mean annual temperature is 9.8 °C. The field follows a four-year crop rotation established approximately in 1985 and alternates between chicory, winter wheat, beet root, potatoes and cover crops (mustard and phacelia).

2.2. Soil sampling and characterization

In this study, five field replicates of two soils, charcoal enriched (CHAR) and reference soils (REF) at least 40 m away from one another, were compared to study the effect of charcoal. Every soil sample collected for each of the five replicates of CHAR and REF sites was bulked from five cores sampled in the topsoil layer (0–30 cm) within a radius of 1 m. These samples were dried at 40 $^{\circ}$ C for 48 h before further analysis.

A subsample of each soil was ground and sieved to 2 mm for baseline physico-chemical characterization. Briefly, pH was determined in distilled water and KCl (1 N) (ratio soil: solution 1:5, ISO 10390). The soil particle-size distribution was determined by gravity sedimentation (NF X 31–107), C and N content were quantified by near infra-red spectrometry after flash dry combustion (respectively ISO 10694; ISO 13878). We measured the cation exchange capacity (CEC) by percolation of 1 M ammonium acetate (pH 7) on soil columns according to the Metson procedure (Metson, 1956). The bioavailable major elements, Ca, Mg, P and K were extracted using ammonium acetate-EDTA 1 M (pH = 4.65; (Lakanen and Erviö, 1971)) and quantified by atomic absorption spectroscopy (Ca, Mg, K) or spectrophotometry (P).

2.3. Fractionation protocol

Soil samples from both studied soils followed a four step fractionation protocol (Fig. 2), which was adapted from the method described by Chenu and Plante, (2006), Gulde et al. (2008) and Virto et al. (2010). In the first step, wet sieving fractionated the soil into four fractions according to size (>2000 μ m, coarse sand size: 2000–250 μ m, fine sand size: 250–50 μ m, silt and clay size (S&C): < 50 μ m). The particulate

Table 1		
Main bulk soil	physico-chemical	parameters

	CHAR	REF
Sand (%)	7.7 ± 0.3	$\textbf{7.5}\pm\textbf{0.4}$
Silt (%)	$\textbf{77.3} \pm \textbf{1.1}$	$\textbf{76.9} \pm \textbf{0.8}$
Clay (%)	15.0 ± 1.0	15.5 ± 0.7
C (%)	$2.6\pm0.5^{*}$	1.4 ± 0.0
Norg (%)	$2.3\pm0.2^{*}$	1.4 ± 0.1
C/N (-)	11.3 ± 1.5	9.9 ± 0.6
CEC (cmol _c kg ⁻¹)	14.0 ± 1.8	12.0 ± 0.6
pH (H ₂ O)	7.7 ± 0.2	$\textbf{7.8} \pm \textbf{0.2}$
pH (KCl)	7.0 ± 0.3	7.1 ± 0.2
P (mg 100 g^{-1})	7.3 ± 0.9	8.0 ± 0.5
K (mg 100 g^{-1})	21.4 ± 2.9	18.2 ± 1.1
Mg (mg 100 g^{-1})	9.9 ± 1.0	8.6 ± 0.7
Ca (mg 100 g ⁻¹)	$314.4 \pm 39.1*$	251.6 ± 31.7

Means \pm SD (n = 5). When followed by * results are significantly different (p.val in text) according to a student paired T-test.



Fig. 2. Flow-chart of the four step fractionation protocol. S&C silt and clay; LF light fraction; HF heavy fraction; Mp macroaggregate protected; mp microaggregate protected; occ occluded; non-occ non-occluded. Dashed squares highlight the difference between the three size classes.

material bigger than 2 mm was discarded. The second step used density to differentiate light fractions (free LF: d < 1.85 g cm⁻³) from heavy fractions (HF: d > 1.85 g cm⁻³): macroaggregate, microaggregate and free silt and clay size heavy fractions (free S&C HF). Step 3 focused on breaking down the macroaggregates (2000–250 µm) to isolate its components into three finer size class fractions (coarse sand, macroaggregate protected fine sand, macroaggregate protected-S&C). Finally, step 4, through a sequential size and density fractionation and dispersion of microaggregates, isolated non-occluded and occluded S&C LF from S&C-size HF within each size class sizes from step 1.

We used the wording *free* for LF and HF after wet sieving, and *occluded* for LF obtained after ultrasonic dispersion of aggregates. This fractionation protocol was applied on both studied soils, CHAR or REF, and for all five replicates.

(i) Step 1 - Size separation by wet sieving

We applied the wet sieving size separation protocol presented by Six et al. (1998). A soil sample of 80 g, previously air-dried at 40 °C for 48 h was submerged in deionized water for 5 min prior to deposition in a water bath on a 2 mm sieve. The sieve was then moved up and down and tilted in a rotational movement for 50 rotations in two minutes at room temperature. The sieve was then removed from the water bath and its inner and outer walls were rinsed with deionized water. The reject (>2 mm) was collected by back-washing in a glass tray and air dried at 60 °C. This wet sieving operation was repeated firstly using a 250 µm sieve on the soil that went through the 2 mm sieve, then repeated a second time on a 50 µm sieve to separate the two smallest fractions. All sieve rejects were back-washed in glass trays after rinsing the sieve and dried at 60 °C. This way four fractions per sample were differentiated as follows: particulate (>2000 µm) > coarse sand size (2000–250 µm) > fine sand size (250–50 µm) > Silt & Clay size materials (<50 µm) (Fig. 2 - step1).

(ii) Step 2 - density fractionation

In this step we divided a 10 g subsample of the three size fractions smaller than 2 mm (coarse sand size > fine sand size > S&C-size materials) obtained from step 1 into a light fraction (LF) and a heavy fraction (HF) as described by Six et al. (1998). For this density separation, subsamples were submerged in 30 ml of sodium polytungstate (SPT, $Na_6(H_2W_{12}O_{40})$, - 1.85 g cm⁻³) in centrifuge tubes. These were then closed and gently stroked 10 times to remove bubbles and without breaking aggregates. The centrifuge tubes were then reopened and with an extra 20 ml of SPT its inner walls were rinsed to gather all the sample in the solution. The samples were placed in a vacuum (-140 kPa) for 10 min, left to rest for 20 min and centrifuged at 4700 g for 10 min. The supernatant was transferred on a filtering flask where a gentle vacuum was applied (glass fiber filter Mechery-Nagel Germany, $\Phi = 0.45 \ \mu m$). After having repeated the centrifuge and the filtering thrice, the LF on the filter was thoroughly rinsed to remove the remaining SPT under a light vacuum. The LF was then recovered in a glass tray by gently back washing the filter with deionized water and then oven dried (60 °C).

To recover the HF, 50 ml of deionized water was added to the pellet in a centrifuge tube and the tube gently stroked until the pellet detached. An additional 20 ml of deionized water was used to clean the tubes inner walls before centrifuging for 10 min at 4700 g. This rinsing was repeated thrice to rid the HF of SPT before recovering the sample in a glass tray and oven drying at 60 °C for 24 h. This step produces a LF and a HF per size class obtained after step 1. These LF are referred to as free LF (macroaggregate, microaggregate or S&C-size) and the HF as macroaggregates, microaggregates or free S&C-size (Fig. 2 - step 2).

(iii) Step 3: Macroaggregate breakdown

This step focused on the HF collected after density fractionation (step 2) of the macroaggregate soil fraction separated from coarse sand size

(2000–250 µm) material. The purpose of this step was to separate this HF into: coarse sands, macroaggregate protected fine sands and macroaggregate protected S&C-size material. To achieve this, the sample was placed in a 250 µm sieve in contact with 50 glass beads in a water bath previously installed on a shaker plate. Deionized water was set to run continuously on the sample while maintaining a 2 cm water front on top. The sample running with the water through the first sieve was collected by means of a funnel and deposited onto a 50 µm sieve. This breakdown procedure was set to last for 5 min on the shaker plate. After this step, deionized water was used to thoroughly rinse the funnel into the 50 μm sieve. The 250 μm sieve was back-washed and the sample material oven dried (60 °C for 24 h) in a glass tray. Then the wet sieving procedure was repeated for the fraction collected in the 50 µm sieve whereby the sieve was tilted and lifted up and down 50 times within two minutes. Again after rinsing the sieve, the reject was back-washed and collected in a glass tray. Both fractions (250–50 μ m and < 50 μ m) were collected in glass trays and oven dried (24 h at 60 °C).

(iv) Step 4: Isolating occluded LF and HF

The first purpose of this step was to isolate the non-occluded (nonocc) S&C-size fraction (<50 µm) contained in macroaggregates from the S&C-size HF (i.e., S&C aggregates or free minerals). For this step an adapted density fractionation was undertaken. The sample was submerged in 30 ml SPT (1.85 g cm^{-3}) and gently shaken into suspension. Using an extra 20 ml of SPT the inner walls of the centrifuge tube were rinsed before being centrifuged at 4700 g for 60 min (all material bigger than 0.7 µm sediments). Using a micropipette, the topmost 15 ml were sampled and deposited on a glass fiber filter (0.4 µm) without vacuum application (composes the non-occ LF). After each filtration the filter was rinsed thoroughly to rid it of the SPT and then back-washed in a glass tray to collect the LF in order to avoid filter clogging, this step was repeated thrice. After the third centrifuge, 40 ml of the supernatant were collected and poured in the filter, rinsed thoroughly with distilled water through the filter. Finally, this filter was back-washed to collect the LF in a glass tray and oven dried (24h at 60°C).

Next, using SPT, the pellet of the centrifuge tube was shaken into suspension and then dispersed by ultrasonication at 440 J ml⁻¹ for 2 min (Soniprep 150 MSE) (Amelung and Zech, 1999) in order to break down remaining S&C-sized aggregates thus freeing the occluded LF. This step was undertaken for macroaggregate protected S&C-size material, microaggregates HF and free S&C-sized material. Following this dispersion, the LF was separated from the HF through a density fractionation (1.85 g cm⁻³) to obtain three new fraction: occluded LF in macroaggregates (occ Mp-LF), occluded LF in microaggregates (occ mp-LF) and occluded LF (occ LF).

Finally, to conclude step 4, a size separation through wet sieving on the HF obtained following the ultrasonication of microaggregate HF only, is undertaken as described previously to separate fine sands from S&C size HF.

The density of SPT was controlled before each density fractionation step. Previous studies reported solubilization of organic compounds with SPT use leading to a loss of C and a decrease in the C recovery after density fractionation (Chenu and Plante, 2006).

The elemental composition (C, H, N) was determined via dry combustion for all fractions (LF and HF). The O concentration was determined on LF only through sample pyrolysis at 1000 °C. The C, H, N, O concentrations were quantified using a Euro EA elemental analyzer (Hekatech, Germany). Prior to the analysis, HF samples were acidified with HCl to remove residual inorganic C.

2.4. Differential scanning calorimetry (DSC)

We used thermal analysis to compare soil fractions isolated by the fractionation protocol to highlight the presence of PyOM in fractions as well as its position in the OM continuum by comparing CHAR and REF fractions. For this thermal analysis, three out of the five field replicates were selected for each studied soil. For these repetitions, all fractions isolated by the fractionation protocol were analyzed. In addition, three products were characterized as reference materials; hand-picked pyrolyzed OM (hand-picked PyOM, n = 3), hand-picked non pyrolyzed coarse OM (hand-picked non-PyOM, n = 3) and freshly pyrolyzed beech (F. Sylvatica – fresh PyOM, n = 3). Hand-picked non-PyOM are crop residues hand-picked from REF soils and hand-picked PyOM are charcoal fragments hand-picked from CHAR soils. The freshly pyrolyzed beech was made in an industrial pyrolyzer (Greenpoch SA, Belgium). Finally, three bulk soils were also used as comparison materials (n = 3).

Samples were analyzed using a differential scanning calorimeter (Netzsch STA 449, Selb, Germany) at the Agroscope facilities in Zürich, Switzerland). Before sample analysis, the sensor was calibrated for temperature and sensitivity (i.e. reaction enthalpy) with six different standards (indium, tin, bismuth, zinc, aluminium and silver).

Prior to scanning, samples were ground to powder. For HF, between 15 and 25 mg of samples were weighed and placed in open Al₂O₃ pans for analysis. For LF 2 mg of samples were mixed to approximately 18 mg Al₂O₃ powder. Precise weights were recorded to enable a correction by the dilution factor. Samples were analyzed under a constant flow of 50 ml min⁻¹ synthetic air from 50 °C to 700 °C at a heating rate of 20 °C min⁻¹. During data treatment a baseline correction was applied on thermograms to account for increased heat capacity of samples. Resulting thermograms were drawn between 200 and 600 °C to highlight the temperature range of interest as data below 200 °C or above 600 °C were never used.

To characterize samples, we used values of peak height as the heat flow at a given temperature (W g⁻¹), peak temperatures as the temperature at which main exotherms occur (°C), the area under the curve as the total heat of reaction of a sample (J g⁻¹) delimited by the baseline, and the thermogram 50% burn-off temperatures (°C) as the temperature where 50% of the total heat flow has occurred. Finally ratios between areas or peak heights considered as thermally labile or stable were used (Leifeld, 2007). In accordance with previous studies, a tipping temperature of ~ 380–390 °C was used to differentiate the thermally labile or stable C forms (Dell'Abate et al., 2000; Lopez-Capel et al., 2005).

Amongst various indicators available for the characterization of C forms using DSC, the sum of peak heights of the exotherms ($R^2 = 0.78 - p$ -value < 0.0001) and the total heat release ($R^2 = 0.91 - p$ -value < 0.0001) displayed strong linear correlation with the C content of fractions.

2.5. PyOM and non-PyOM quantification in charcoal rich soils

In CHAR soils, the PyOM was quantified as the additional heat flow occurring in the thermally stable region (>390 °C) as opposed to the heat flow in adjacent REF soils. This tipping temperature was established based on our results and in line with previous findings (Dell'Abate et al., 2000; Lopez-Capel et al., 2005). This additional heat flow over the sum of punctual heat flows (peak heights) of CHAR samples multiplied by the samples C content corresponds to the PyOM content of CHAR soils (eq (1)).

$$PyOM = \frac{\sum CHAR_{st,peaks} - \sum REF_{st,peaks}}{\sum CHAR_{peaks}} * C_{CHAR}$$
(1)

where **PyOM** the PyOM content (g kg⁻¹), st. peaks the heat flow of exotherms occurring above 390 °C and C_{CHAR} the C content of CHAR samples.

Changes in non-PyOM contents, δ (non-PyOM), were calculated as the ratio of differences in CHAR to REF labile peaks height (<390 °C) to the difference in sum of peaks between studied soils, the whole multiplied by the difference of C content between CHAR and REF samples (eq (2).).

$$\delta(non-PyOM) = \frac{CHAR_{lab.peak} - REF_{lab.peak}}{\sum CHAR_{peaks} - \sum REF_{peaks}} * (C_{CHAR} - C_{REF})$$
(2)

where δ (*non-PyOM*) the change in non-PyOM content (g kg⁻¹), *lab. peak* the heat flow of exotherms occurring below 390 °C, *C*_{CHAR} and *C*_{REF} the C content of CHAR and REF samples (g kg⁻¹).

2.6. Data analysis and signal treatment

All data analysis, DSC signal treatment and graph plotting were performed using R studio v. 3.4.2 (R Core Team, 2017). Results of all parameters are presented as the means \pm standard deviation (SD). Comparison between fractions of similar density and size class were undertaken by analysis of variances (ANOVA) or by paired t-tests for soil baseline characteristics of bulk soils. Mean classification was pursued through a least significant differences test (LSD-test) and indicated by different letters following mean \pm SD unless indicated otherwise in the caption. For the application of this test the normality of the distribution and homoscedasticity of variances was presumed given the low number of repetitions (n < 10).

To ensure the validity of our soil fractionation results, the mass recovery and C content (mass by concentration) were calculated at each fractionation step. These were obtained as the sum of fraction masses and C content divided by that of sample used as input for the fractionation step.

3. Results

3.1. Soil characteristics

Beyond the presence of century-old charcoal kiln sites, the study site was selected for its homogeneity (REF soil) in terms of texture (sand = 7.5 \pm 0.4%, silt = 76.9 \pm 0.8%, clay = 15.5 \pm 0.7%), C concentration (1.4 \pm 0.0%), CEC (12.0 \pm 0.6 cmol_c kg⁻¹) and pH (H₂O – 7.8 \pm 0.2). On average, C concentrations in CHAR soils ([1.9 – 3.3]%) were 91 \pm 34% higher than in REF soils ([1.3 – 1.4]%) (p-val. < 0.01). Similarly, organic N concentrations were 66 \pm 11% higher (p-val. < 0.001) in CHAR ([2.1 – 2.6] g kg⁻¹) than in REF soils ([1.2 – 1.5]%). C, N concentrations as well as C:N ratios were more variable among replicates in the CHAR soils than in the REF soils. The two studied soils were of identical texture (silty loam) and had similar pH values. No significant differences in CEC were reported. Bioavailable Ca²⁺ concentrations were on average 26 \pm 15% higher in CHAR soils (p-val. < 0.05) (Table 1).

3.2. Weight and carbon distribution amongst fractions

In CHAR soils, macroaggregates contributed significantly more (p-val. < 0.01) to the total soil weight than microaggregates or free S&C fractions (CHAR_{MACRO} = 48.9 \pm 12.8, CHAR_{MICRO} = 25.5 \pm 8.3, CHAR_{Free} S&C = 25.6 \pm 4.6 g 100 g⁻¹ soil). In reference soils, macroaggregate and microaggregate fractions were equal and accounted significantly more to the total soil weight (p-val. < 0.01) than the free S&C fraction (REF_{MACRO} = 41.3 \pm 9.5, REF_{MICRO} = 36.0 \pm 8.7, REF_{Free} S&C = 22.7 \pm 2.6 g 100 g⁻¹ soil - Fig. 3).

The weight recovery following the wet sieving procedure was of 99.4 \pm 0.3% and 99.6 \pm 0.1% for CHAR and REF respectively. For step 4 of our fractionation protocol, 93.6 \pm 4.8% and 89.0 \pm 7.5% of initial mass were recovered (CHAR and REF respectively). For this step the C content recovery was of 87.9 \pm 52.1% (CHAR) and 100.6 \pm 20.1% (REF).

Macroaggregate-size free LF was the most abundant free LF in CHAR soils (0.7 \pm 0.2%) and was also significantly more abundant (p-val. < 0.05) in CHAR soils than in REF soils (Table 2). Micro and S&C-size free LF displayed similar contribution to the soil mass between the two studied soils (Table 2). Furthermore, free LF displayed systematically higher C contents (i.e., fraction C concentration multiplied by fraction



Fig. 3. Size class distribution. Weight distribution amongst fractions of different sizes following wet sieving of the studied soils. Points represent the 5 field replicates. Symbols and box shadings highlight different fraction sizes. Weight recoveries (mean \pm SD) are: CHAR = 99.4 \pm 0.3%, REF = 99.6 \pm 0.1%.

mass) in CHAR soils than in REF soils (Table 2) due mainly to higher C concentrations (i.e., mass of C per mass of the fraction). Macroaggregate size free LF of charcoal rich soils contributed the most to the C content compared to all other free LF studied soils for both treatments.

While no differences between studied soils were observed, occluded LF contributed more to the total soil weight than free LF (Table 2). In both studied soils, microaggregate-size fractions accounted the most to the total soil C content for both occluded LF and free LF. Microaggregate-size occluded LF stored twice the amount of C than microaggregate-size free LF (Table 2).

The macroaggregate protected HF was significantly more abundant than microaggregate protected and free HF in charcoal rich soils. In reference soils, macroaggregate and microaggregate protected HF, which displayed the same contribution to the total soil mass, were more abundant than free HF (Table 2). The C concentrations of HF ranged between 0.7 ± 0.2 to $0.8 \pm 0.2\%$ in reference soils and from 0.9 ± 0.1 to $1.0 \pm 0.2\%$ in CHAR soils. In charcoal rich soils, macroaggregate protected HF (3.5 ± 1.2 g kg⁻¹ soil) contributed more to the total soil C content than microaggregate protected HF and free HF (2.2 ± 0.6 , 2.2 ± 0.3 g kg⁻¹ soil). In REF soils, macroaggregate and microaggregate protected HF accounted similarly to the total soil C content, which was a greater contribution than free HF (Table 2).

Summing all fractions regardless of size into either LF or HF showed that although HF accounted for most of the soil mass (CHAR = 89.0 \pm 2.2, REF = 86.5 \pm 1.0%), it merely accounted for 39.0 \pm 25.0% of the recovered soil C in CHAR soils (CHAR_HF = 7.9 \pm 1.4 g C kg⁻¹) and 48.0 \pm 23.0% in reference soils (REF_{HF} = 6.8 \pm 1.2 g C kg⁻¹). In charcoal rich soils, LF (free and occluded combined) comprised 12.2 \pm 3.1 g C kg⁻¹ soil of which 6.8 \pm 3.0 g C kg⁻¹ soil occurred in occluded LF rather than free LF (5.4 \pm 0.7 g C kg⁻¹ soil). In reference soils, LF combined comprised 7.5 \pm 1.5 g C kg⁻¹ soil, mainly as occluded LF (REF_{occ.LF} = 4.7 \pm 1.3 g C kg⁻¹ soil) rather than as free LF (REF_{freeLF} = 2.8 \pm 0.7 g C kg⁻¹ soil). C concentrations of free LF increased with aggregate size and ranged between 25.6 \pm *na* and 41.9 \pm 7.4% for charcoal rich soils and between 10.0 \pm 2.9 and 36.7 \pm 10.1% in reference. Microaggregate occluded LF (RAR = 31.1 \pm 5.4, REF = 30.3 \pm 4.2%) than macroaggregate occluded LF (CHAR = 26.1 \pm 2.4, REF = 21.3 \pm 2.4% - Table 2).

(µm) (g free LF CHAR 2000–250 6. 250–50 4.		J	INIUL	,	N	CIN	PYOIN
free LF CHAR 2000–250 6. 250–50 4.	(g kg ⁻¹ soil)	(%)	(%)	(g kg ⁻¹ soil)	(mg kg ⁻¹ soil)	(-)	(g kg ⁻¹ soil)
250–50 4.	$6.7\pm2.0~{\rm A}$	$41.9 \pm 7.4~\mathbf{A}$	$15.3\pm2.8~\mathrm{A}$	$2.7\pm0.4~\mathrm{A}$	$98.4 \pm 17.7~\mathbf{A}$	$27.5\pm3.0~\mathrm{A}$	1.1 ± 0.7
	$4.2\pm1.5~\mathrm{B}$	$36.5\pm4.5~\mathbf{A}$	$14.6\pm1.4~\mathrm{AB}$	$1.7\pm0.5~\mathrm{B}$	$67.2\pm18.4~\mathrm{B}$	$25.0 \pm 1.5~\mathbf{A}$	0.3 ± 0.1
<50 3.	$3.8\pm3.3~\mathrm{B}$	$25.6 \pm na$	$11.8\pm2.8~\mathrm{B}$	$1.0\pm na$	$49.5\pm0.0~\mathrm{BCD}$	$21.5\pm na$	$0.1\pm na$
REF 2000–250 3.	$3.9\pm0.7~\mathrm{B}$	$36.7\pm10.1~{ m A}$	$14.8\pm1.2~\mathrm{AB}$	$1.5\pm0.6~\mathrm{BC}$	$58.2\pm10.3~\mathrm{BC}$	$25.0\pm7.6~\mathrm{A}$	
250-50 2.	$2.6\pm0.8~{ m B}$	$34.9\pm3.9~\mathrm{AB}$	$15.5\pm2.9~\mathrm{A}$	$1.0\pm0.3~\mathrm{CD}$	$42.6\pm16.0~\mathrm{CD}$	$22.8 \pm 2.5~\mathbf{A}$	
<50 3.	$3.6\pm1.8~\mathrm{B}$	$10.0\pm2.9~{ m C}$	$6.5\pm2.2~{ m C}$	$0.3\pm0.2~\mathrm{D}$	$22.3\pm12.0~\mathrm{D}$	$15.7\pm1.8~\mathrm{B}$	
occ LF CHAR 2000–250 7.	$7.8\pm6.4~\mathrm{FG}$	$26.1\pm2.4~\mathrm{FG}$	$17.9\pm1.6~\mathrm{GH}$	$2.1\pm1.8~\mathrm{FG}$	$142.4\pm117.6~\mathrm{FG}$	$14.6\pm0.8~\mathrm{FG}$	$1.2 \pm na$
250–50 11	$11.1\pm 6.7~{ m F}$	$31.1\pm5.4~\mathrm{F}$	$20.2\pm4.7~\mathrm{FG}$	$3.8\pm2.3~\mathrm{F}$	$247.1\pm149.6~{\rm FG}$	$15.7\pm1.5~\mathrm{F}$	0.2 ± 0.0
<50 4.	$4.2\pm0.9~{\rm FG}$	$24.4\pm4.2~\mathbf{G}$	$15.3\pm1.6~\mathrm{H}$	$1.0\pm0.4~{ m G}$	$63.3\pm18.3~\mathrm{G}$	$15.8\pm1.6~\mathrm{F}$	0.5 ± 0.2
REF 2000–250 5.	$5.9\pm2.4~\mathrm{FG}$	$17.7\pm3.4~\mathrm{H}$	$13.7\pm3.1~{ m H}$	$1.0\pm0.5~{ m G}$	$82.0\pm41.4~{\rm G}$	$13.1\pm1.4~\mathrm{GH}$	
250–50 9.	$9.1\pm1.9~{\rm FG}$	$30.3\pm4.2~\mathrm{F}$	$22.2\pm4.4~\mathrm{F}$	$2.7\pm0.8~{ m FG}$	$200.2\pm60.3\mathrm{FG}$	$13.9\pm1.4~\mathrm{GH}$	
<50 4.	$4.1\pm3.5~{\rm G}$	$21.3\pm2.4~\mathrm{GH}$	$17.1\pm1.9~\mathrm{GH}$	$0.9\pm1.0~{\rm G}$	$70.1\pm72.9~{ m G}$	$12.4\pm0.7~{ m H}$	
HF CHAR 2000–250 41	$414.2 \pm 127.6 \mathrm{J}$	$0.9\pm0.2\mathrm{JK}$	$0.9\pm0.3~{ m J}$	$3.5\pm1.2~\mathrm{J}$	$403.6\pm 228.8~{\rm J}$	$9.8\pm2.4~\mathrm{K}$	0.3 ± 0.2
250-50 22	$221.6 \pm 70.4~\mathbf{L}$	$1.0\pm0.2\mathrm{J}$	$0.8\pm0.2~\mathrm{JKL}$	$2.2\pm0.6~\mathrm{KL}$	$168.6\pm33.4~\mathrm{KL}$	$13.1\pm3.2~\mathrm{JK}$	0.2 ± 0.1
<50 25	$\textbf{252.1} \pm \textbf{56.1} ~ \textbf{JK}$	$0.9\pm0.1~ m JK$	$0.7\pm0.2~\mathrm{JKL}$	$2.2\pm0.3~\mathrm{KL}$	$183.5\pm100.7~\mathrm{KL}$	$13.9\pm3.3~\mathrm{J}$	0.4 ± 0.1
REF 2000–250 3:	$337.5\pm80.5~\mathrm{JK}$	$0.8\pm0.2~{ m K}$	$0.9\pm0.4~\mathrm{JK}$	$2.7\pm1.0~\mathrm{JK}$	$306.6 \pm 168.9 ~ \mathrm{JK}$	$10.0\pm2.8~{\rm K}$	
250-50 31	$310.7\pm79.6~\mathrm{JKL}$	$0.8\pm0.1~\mathrm{JK}$	$0.6\pm0.1~{ m KL}$	$2.6\pm0.7~\mathrm{JK}$	$178.3\pm43.5\mathrm{KL}$	$14.5\pm2.2~{\rm J}$	
<50 21	$217.1\pm25.0~\mathrm{L}$	$0.7\pm0.2~{ m K}$	$0.5\pm0.2~\mathrm{L}$	$1.5\pm0.3~\mathrm{L}$	$109.3\pm25.8\mathrm{L}$	$14.0\pm1.2~\mathrm{J}$	
hand-picked PyOM		37.8 ± 7.9	5.4 ± 1.7			76.6 ± 27.4	
fresh PyOM		$\textbf{79.6}\pm\textbf{0.8}$	8.8 ± 0.6			90.6 ± 6.9	

3.3. Differential scanning calorimetry

Three reference materials and two bulk soils were analyzed to allow for the discrimination of various forms of OM (Fig. 4). Three distinct exotherms characterized hand-picked PyOM (top - Fig. 4) with a major peak at 443 \pm 4 °C. Fresh PyOM was characterized by a single exotherm (485 \pm 2 °C) and a higher punctual heat flow. Hand-picked non-PyOM (crop residues) were characterized by a strong first exotherm occurring at 350 \pm 2 °C and a second smaller at 430 \pm 1 °C.

The total heat flow of CHAR bulk soils was on average $84 \pm 59\%$ greater than REF bulk soils (Table 4). CHAR bulk soils were characterized by four exotherms occurring at temperatures similar to those occurring in hand-picked PyOM and hand-picked non-PyOM combined (top - Fig. 4). Similarly, exotherms of reference bulk soils occurred at similar temperatures to those which characterized hand-picked non-PyOM.

Fig. 5 illustrates the thermograms of the three main types of fractions addressed in this study, the free LF, occluded LF and HF (respectively row A, B, C - Fig. 5) as a function of the aggregate size class in which they occur (col. 1: macroaggregate, col. 2: microaggregate, col. 3: free S&C-size - Fig. 5).

Thermograms of the free LF (Fig. 5 - A) showed more pronounced exotherms occurring between 400 and 500 °C for CHAR than REF fractions whereas thermograms of REF fractions tended to be more spread out throughout the thermal continuum than CHAR fractions. Nonetheless, 50% burn-off temperatures were not significantly smaller in REF than CHAR fractions (p-val. > 0.05). Finally, a decrease in total heat flow as a function of fraction size was reported whereby heat flows of S&C-size free LF were significantly smaller (CHAR = 232 \pm 60 J g⁻¹, REF = 172 \pm 30 J g⁻¹) than microaggregate-size free LF (CHAR = 665 \pm 254 J g⁻¹, REF = 753 \pm 160 J g⁻¹), and macroaggregate-size free LF (CHAR = 821 \pm 185 J g⁻¹, REF = 645 \pm 262 J g⁻¹) regardless of the studied soil (p-val. < 0.01). However, total heat flows are equal between fractions when normalized by their C content.

The occluded LF (Fig. 5 - B) were characterized by two distinct exotherms visible as individual peaks or marked shoulders occurring between 325-342 °C and 430–480 °C for both studied soils (Table 4). In CHAR soils, the second exotherm (430–480 °C) had a higher relative importance as shown by the increase of thermally stable:total peaks ratios by 42.6% in macroaggregate occluded-LF, 44.3% in microaggregate occluded-LF and 50.0% in S&C-size occluded-LF compared to REF soils (Table 4). No differences in 50% burn-off temperatures were noted between studied soils within a size class yet macroaggregate protected occluded LF have significantly smaller 50% burn-off temperatures than microaggregate protected or S&C occluded LF (p-val. < 0.05) (Table 4).

HF thermograms were characterized by a main exotherm occurring for both studied soils at \sim 323 °C. CHAR fractions were differentiated from REF HF by a marked shoulder in the DSC curves occurring at \sim 405 °C. Thermograms of HF (Fig. 5 - C) were more systematic between repetitions as highlighted by the average heat flow. The average heat flow occurring in HF was of 187.1 \pm 42.3 J g^{-1} for CHAR soils and 147.5 \pm 31.5 J g⁻¹ for REF soils where the standard deviation represented respectively 22.5 and 21.3% of the total heat flow. In occluded LF the average total heat flow was of 356.1 \pm 159.0 J g^{-1} for CHAR and 242.8 \pm 147.1 J g⁻¹ for REF soils with the standard deviation representing 44.6 and 60.6% for CHAR and REF soils. In free LF, it represented 55.0% for CHAR samples and 55.5% in reference samples (CHAR_{free LF} = 495.2 \pm 272.6 J g⁻¹, REF_{free LF} = 523.8 \pm 291.228 J g⁻¹). For HF, 50% burn-off temperatures were always significantly higher for charcoal rich than reference soils (p-val. < 0.01) in all size classes and notably smaller than in LFs (Table 4).

Finally, thermograms differed between hand-picked reference materials (Fig. 4) and fractions isolated by the fractionation protocol (Fig. 5). Indeed the first (\sim 386 ± 3 °C) and last (\sim 520–528 °C) exotherms of hand-picked PyOM (Fig. 4) were not found in soil fraction

Table



Fig. 4. DSC reference curves. DSC thermograms of reference material for the characterization of C forms. Top panel: compares three types of OM forms, hand-picked (abbreviated h.p) non-PyOM, hand-picked century-old PyOM and fresh PyOM. Bottom panel: compares CHAR and REF bulk soils.

thermograms (Fig. 5).

3.4. Elemental analyses

Aged hand-picked PyOM displayed much higher H:C (0.70 \pm 0.10) and O:C (0.40 \pm 0.10) atomic ratios than fresh PyOM (H:C = 0.02 \pm 0.00; O:C = 0.10 \pm 0.00) (Table 3). Differences in H:C and O:C ratios were greater between soil fractions (free or occluded) than between CHAR and REF soils (Fig. 6). Indeed, free LF had lower ratios than occluded LF for both studied soils (Table 3). However, S&C-size free LF had higher H:C and O:C ratios in REF soils than in CHAR soils. For occluded LF, differences in H:C ratios between studied soils were not significant, and only O:C ratios of macroaggregate-size occluded LF were greater in REF fractions than in CHAR fractions.

H:C atomic ratios increased from free LF, to occluded LF and HF for both studied soils (Table 3). For microaggregate protected and free HF, H:C ratios were higher in REF (microaggregate protected HF: 3.22 \pm 0.34, free HF: 4.24 \pm 0.39) than CHAR soils (microaggregate protected HF: 2.71 \pm 0.15; free HF: 3.44 \pm 0.43).

Inversely, C:N atomic ratios decreased from free LF ([15.7 \pm 1.8 – 27.5 \pm 3.0]), to occluded LF ([12.4 \pm 0.7 – 15.8 \pm 1.6]) and HF ([9.8 \pm 2.4–14.5 \pm 2.2]). In free LF, C:N ratios decreased as a function of size while their values were similar in occluded LF of the three size classes. Furthermore, C:N ratios of HF were lower in macroaggregate protected HF than in microaggregate protected and free HF materials (p-val. <

0.05), for both studied soils (Table 3).

3.5. Combining analytical practices

Amongst the different fractions of both studied soils, H:C atomic ratios decreased (Table 3) alongside an increase in stable:total heat flow ratios (Table 4). HF (i.e., S&C minerals) were characterized by most of the heat flow occurring at low temperatures as opposed to LF. While H:C ratios were similar between studied soils, HF of CHAR soils displayed higher stable:total heat flows than HF of REF soils (CHAR = 24.0 ± 3.0 , REF = $16.1 \pm 2.5\%$). On average in CHAR soils, free LF showed lower H: C and higher stable:total ratios (H:C = 0.69 ± 0.09 ; stable:total = $76.6 \pm 11.1\%$) than for occluded LF (H:C = 1.14 ± 0.16 ; stable:total = $68.2 \pm 7.8\%$) whereas in REF soils, average H:C ratios for occluded fractions were lower than for free LF (occ LF = 1.27 ± 0.15 , free LF = 1.08 ± 0.44) while stable:total ratios were similar (occ LF = 63.2 ± 4.8 , free LF = $64.1 \pm 6.2\%$ - Table 4)

3.6. PyOM and non-PyOM quantification in charcoal rich soils

Charcoal rich bulk soils contained between 5.8 and 11.9 g kg⁻¹ soil of PyOM. This amounts for between 29.9 and 41.3% of the total SOC in bulk CHAR soils. In free LF, the PyOM accounted between 13.7 and 41.2% of the total SOC in these fractions. Most of the PyOM was found as macroaggregate-size free LF and accounted for up to 1.1 ± 0.7 g kg⁻¹ soil (Table 2). Similar amounts of PyOM were found in occluded LF and accounted between 5.4 and 55.9% of total SOC contained in occluded fractions. HF contained between 0.2 ± 0.1 and 0.4 ± 0.1 g kg⁻¹ soil of PyOM. This represented between 7.3 and 16.2% of the SOC contained in HF and was highest for free HF (0.4 ± 0.1 g kg⁻¹ soil).

A higher heat of reaction occurring at thermally labile temperatures in CHAR soils than REF soils shows that differences in C content did not solely occur as thermally stable OM but also as labile PyOM or non-PyOM. Bulk charcoal rich soils accounted 0.4 to 2.3 g C kg⁻¹ soil of additional thermally labile OM compared to reference soils. This represented between 5.8 and 16.2% of the additional C content of CHAR soils. The combined heat flows of occluded LF were on average 38.7% higher in CHAR than REF fractions (CHAR_{occ} = 9.8 ± 4.3 W g⁻¹, REF_{occ} = 7.1 ± 1.8 W g⁻¹) although non-significantly different due to the variability between CHAR repetitions. However, an increase in thermally labile peak height was not systematic between samples and hence not statistically significant for occluded LF (p-val. > 0.05) nor HF (p-val. > 0.05).

4. Discussion

4.1. Physico-chemical characteristics of charcoal rich soils

Our results show close to a twofold increase in C contents as a result of century-old PyOM accumulation in cultivated soils as previously reported for similar sites (Hernandez-Soriano et al., 2016b; Kerré et al., 2016; Hardy et al., 2017a). As both studied soils were sampled in the same field, they have received similar amendments and field works in identical climatic conditions. We can therefore be confident that such an increase in C content is induced by the presence of PyOM and corresponds to increases in both PyOM and non-PyOM contents in kiln soils. Accumulation of non-PyOM in kiln soils have been previously shown to originate from a decreased mineralization rates of recent non-PyOM (Hernandez-Soriano et al., 2016a; Kerré et al., 2016) and from the adsorption of dissolved organic compounds onto charcoal particles (Hernandez-Soriano et al., 2016b; Kerré et al., 2017). A relative greater aboveground biomass production on kiln soils (Heidarian Dehkordi et al., 2020) may also be responsible for a positive feedback loop through an increase in non-PyOM inputs to the charcoal rich soils.

Previous studies on similar kiln sites in Wallonia reported significant increases in CEC related to the presence of charcoal (Kerré et al., 2016;



Fig. 5. DSC signatures of soil fractions. Graphs are presented as: **free LF (light fraction)** (A - top row) either macroaggregate-size ($2000-250 \mu m$), microaggregate-size ($250-50 \mu m$) or silt and clay size ($50> \mu m$). Middle (B) and bottom (C) rows show **occluded LF** or **HF (heavy fraction)** of silt and clay size, either protected in macroaggregates or microaggregates or as free material. In each graph the three replicas were represented, each one was plotted by pair with the adjacent samples of the opposite studied soil (CHAR/REF). Curves were vertically shifted to facilitate the comparison between replicas. Vertical lines show the mean (n = 3) 50% burn-off temperature per fraction (REF = dashed line, CHAR = solid line) and the horizontal line at the top of each graph represents the standard deviation (REF = light grey, CHAR = dark grey).

Hardy et al., 2017a). Our results show slight yet non-statistically significant increases. Furthermore, higher CEC has been estimated for century-old PyOM fragments (PyOM = 414 cmol_c kg⁻¹) compared to non-PyOM (non-PyOM = 213 cmol_c kg⁻¹) explained by an increase in the amount of functional groups with charcoal ageing and cultivation combined with the high specific surface area of PyOM (Hardy et al., 2017a). Additionally, CHAR soils displayed greater bioavailable Ca²⁺ contents than adjacent REF soils as previously reported (Hardy et al., 2017a, 2017b). Increased Ca²⁺ contents in CHAR soils result from the

high affinity between Ca²⁺ and carboxylate groups found on aged PyOM thus potentially serving as a mediator for PyOM interaction with soil constituents (Kalinichev and Kirkpatrick, 2007).

4.2. Elemental C:N, H:C and O:C ratios support charcoal-induced organomineral associations

The presence of PyOM in charcoal enriched soils resulted in a greater macroaggregate formation at the expense of free microaggregates in

Table 3

H:C and O:C atomic ratios of soil fractions.

fraction	soil	size (µm)	H:C (-)	0:C (-)
free LF	CHAR REF	2000–250 250–50 < 50 2000–250 250–50	$\begin{array}{c} 0.69 \pm 0.11 \ \mathrm{C} \\ 0.72 \pm 0.06 \ \mathrm{C} \\ 0.67 \pm na \\ 0.95 \pm 0.20 \ \mathrm{B} \\ 0.75 \pm 0.13 \ \mathrm{C} \end{array}$	$\begin{array}{c} 0.30 \pm 0.10 \text{ A} \\ 0.31 \pm 0.00 \text{ A} \\ 0.41 \pm na \\ 0.34 \pm 0.00 \text{ A} \\ 0.41 \pm 0.00 \text{ A} \end{array}$
		<50	$1.55\pm0.11~\text{A}$	$1.30\pm0.50\;B$
occ LF	CHAR	2000–250 250–50 < 50	1.24 ± 0.12 F 1.19 ± 0.20 FG 1.01 ± 0.12 G	$\begin{array}{c} 0.39 \pm 0.04 \; \text{G} \\ 0.44 \pm 0.03 \; \text{G} \\ 0.38 \pm 0.01 \; \text{G} \end{array}$
	REF	2000–250 250–50 < 50	$\begin{array}{c} 1.32 \pm 0.20 \; \text{F} \\ 1.31 \pm 0.12 \; \text{F} \\ 1.18 \pm 0.14 \; \text{FG} \end{array}$	$\begin{array}{c} 0.85 \pm 0.00 \; F \\ 0.37 \pm 0.00 \; G \\ 0.39 \pm 0.05 \; G \end{array}$
HF	CHAR	2000–250 250–50 < 50	$\begin{array}{l} 3.14 \pm 0.37 \text{ KL} \\ 2.71 \pm 0.15 \text{ L} \\ 3.44 \pm 0.43 \text{ KL} \end{array}$	
	REF	2000–250 250–50 < 50	$\begin{array}{l} 3.58 \pm 0.50 \text{ K} \\ 3.22 \pm 0.39 \text{ K} \\ 4.24 \pm 0.39 \text{ J} \end{array}$	
hand-picked PyOM fresh PyOM			$\begin{array}{c} 0.70 \pm 0.10 \\ 0.02 \pm 0.00 \end{array}$	$\begin{array}{c} 0.40\pm0.10\\ 0.10\pm0.00\end{array}$

Either free LF (light fraction), occluded LF and HF (heavy fraction) as a function of treatment and of the size classes in which they were contained. Different letters indicate significant differences (p \leq 0.05) between studied soils as a function of fraction. "na" indicated that not enough fraction material was recovered for a complete set of analysis. No O:C ratios were measured for mineral dominated fractions.

comparison to reference soils. Macroaggregate turnover is known to be relatively rapid particularly in cropped ecosystems where tillage damages soil structure (Six et al., 2000; Chenu et al., 2019). In CHAR soils, where macroaggregates are more abundant than in REF soils, it is likely that PyOM acts as an additional binding agent between soils constituents and microaggregates (Six et al., 2000). Furthermore, a larger macroaggregate fraction in CHAR soils may also result from higher aggregate stability (Pituello et al., 2018) caused by increased inter-particular cohesion (Sun and Lu, 2014) or faster aggregate formation as a result of the specific surface area of PyOM. Century-old PyOM therefore improves soil aggregation which is a key property for soil structure, essential in maintaining soil fertility and which may also further increase the stability of OM through increased physical disconnection with

Table 4

DSC curve characteristics of soil fractions.

decomposers (Schmidt et al., 2011; Lehmann and Kleber, 2015).

Our results showed significant differences in O:C and H:C ratios between free and occluded LF for both studied soils thereby showing that the position of PyOM and non-PyOM in the soil matrix and their chemical composition are strongly linked. Lower O:C and H:C atomic ratios suggest a greater abundance of large aromatic structures in free LF than in occluded LF whereas the latter may be enriched in either PyOM with more functional entities such as carboxyl groups or additional non-PyOM (Brodowski et al., 2005; Cheng et al., 2008a).

Long-term residence of PyOM in soils results in higher O:C and H:C atomic ratios (Cheng et al., 2008a; Nguyen et al., 2008; Hardy et al., 2017b) compared to fresh PyOM (Hardy et al. 2017b). Ageing of PyOM is known to occur mainly through oxidation of the particles and results in the development of phenol, carboxyl and carbonyl functional groups on the PyOM aromatic structures (Cheng et al., 2008a). Although free LF are highly weathered compared to fresh PyOM, their lower H:C and O:C ratios in comparison to occluded LF suggests free LF may be less oxidized than occluded LF. In turn, this higher functionalization of PyOM in occluded LF compared to free LF supports its increased reactivity towards surrounding soil constituents. As all the PyOM in CHAR soils is of similar age range and as most of the oxidation occurs rapidly following the addition of PyOM to the soil, it is expected that particles would have faced similar oxidative agents (Nguyen et al., 2008). Therefore, differences in O:C and H:C ratios between free and occluded LF may originate from products with varying initial properties. Indeed, slight changes in pyrolysis conditions throughout the kiln during charcoal production may result in varying degrees of aromaticity or aromatic condensation (B.P. Singh et al., 2012; Wiedemeier et al., 2015) resulting in different degrees of hydroxylation and carboxylation (Lehmann et al., 2005) and in turn leading to charcoal fragments of varying persistence. However, it may be that when first produced, the PyOM was physically and chemically homogeneous in composition and that, with time and farming practices, the PyOM fragments were broken down to different size classes which resulted in different degrees of oxidation. In addition, higher H:C and O:C atomic ratios of occluded LF than free LF in charcoal enriched soils may also originate from higher amounts of non-PyOM such as microbial related biomass (Lehmann et al., 2011) or stabilized fresh non-PyOM residues (Hernandez-Soriano et al., 2016a). In HF, differences in H:C and O:C ratios result either from mineral phases or changes in OM. However, as texture and mineralogy are similar throughout the study site, differences in H:C and O:C ratios between

				exother	exotherm 1 exotherm 2		erm 2		
fraction	soil	size	50% Burnoff T	peak T	Heat flow	peak T	Heat flow	Labile: Total Heat flow	Stable: Total Heat flow
		(µm)	(°C)	(°C)	(W g ⁻¹)	(°C)	$(W g^{-1})$	(%)	(%)
free LF	CHAR	2000-250	$436.3\pm35.9~\text{AB}$			$442.3\pm38.6~\text{A}$	$22.9\pm3.7~\mathrm{A}$	$28.5\pm15.2\text{ BC}$	$72.0\pm14.9~\text{AB}$
		250-50	$443.3\pm33.5\text{ AB}$			$448.7\pm24.0~\text{A}$	$16.7\pm5.6\;\text{AB}$	$24.2\pm13.3~\mathrm{C}$	$76.2\pm13.0~\text{A}$
		< 50	$462.3\pm3.1~\mathrm{A}$			$471.0\pm2.0~\text{A}$	$6.8\pm2.6\;\text{C}$	$22.6\pm1.0~\mathrm{C}$	$77.5\pm1.0~\text{A}$
	REF	2000-250	$413.5\pm21.9~\text{B}$			$442.3\pm38.6~\text{A}$	$11.0\pm7.3~\text{BC}$	$41.2\pm7.5~B$	$59.3\pm7.4~\text{B}$
		250-50	$439.8\pm9.9~\text{AB}$			$454.0\pm6.2~\text{A}$	$10.5\pm3.5\text{ BC}$	$32.4\pm2.4~\text{BC}$	$67.9\pm2.4~\text{AB}$
		< 50	$446.8\pm23.2\text{ AB}$			$483.0\pm4.4~\text{A}$	$3.2\pm0.6\;\text{C}$	$34.0\pm6.2\text{ BC}$	$66.2\pm6.2~\text{AB}$
occ LF	CHAR	2000-250	$414.3\pm16.6~\text{G}$	$335.0\pm1.4~\mathrm{F}$	$3.8\pm3.7~\text{F}$	$449.7\pm15.0~\text{G}$	$6.7\pm6.3~\text{F}$	$40.8\pm6.7~\text{FG}$	$59.5\pm6.8~\text{GH}$
		250-50	$458.3\pm2.0\ \text{F}$	$328.7 \pm 1.5 \ \mathrm{H}$	$3.0\pm0.4\ F$	$469.7\pm4.6\ F$	$8.5\pm3.1\;\text{F}$	$26.6\pm2.8~\mathrm{H}$	$73.6\pm2.8~\text{F}$
		< 50	$449.5\pm18.8\ \text{F}$	$342.0\pm0.0\;\text{F}$	$3.0\pm2.1~\text{F}$	$472.3\pm5.7~\text{F}$	$7.9\pm5.0\;\text{F}$	$28.7\pm7.0~\mathrm{H}$	$71.5\pm6.9\ F$
	REF	2000-250	$418.0\pm20.2~\text{G}$	$335.0 \pm 1.4 \; \text{F}$	$2.5\pm0.4\ F$	$471.7 \pm 1.2 \; \text{F}$	$2.9\pm0.8\;\text{F}$	$42.8\pm5.4~\text{F}$	$57.5\pm5.4~\mathrm{H}$
		250-50	$447.3\pm8.8\ \text{F}$	$332.0\pm4.0\;G$	$3.5\pm1.7\;\text{F}$	$464.3\pm8.3\ F$	$6.8\pm3.2\ \text{F}$	$31.7 \pm 1.5 \text{ GH}$	$68.5\pm1.5\;\text{FG}$
		< 50	$441.7\pm18.9\ FG$	$342.0\pm0.0\;F$	$1.1\pm0.5\;\text{F}$	$475.7\pm4.0\;F$	$2.1\pm1.3~\text{F}$	$38.8 \pm 6.2 \; \mathrm{FG}$	$61.4\pm6.2~\text{GH}$
HF	CHAR	2000-250	$338.2\pm7.8~\text{JK}$	$324.0 \pm 1.7 \text{ K}$	$0.4\pm0.1\;J$	405.0 ± 0.0	$0.2\pm0.0~\text{JKL}$	$76.9\pm4.6~\text{KL}$	$23.5\pm4.6~\text{JK}$
		250-50	$341.8\pm3.8~J$	$329.7 \pm 2.1 \text{ J}$	$0.4\pm0.1~J$	405.0 ± 0.0	$0.2\pm0.0~\text{JK}$	$77.0 \pm 4.2 \text{ KL}$	$23.4\pm4.2~\mathrm{JK}$
		< 50	$339.2\pm4.2~\mathrm{J}$	$322.0\pm4.4~\text{K}$	$0.4\pm0.0\;J$	405.0 ± 0.0	$0.2\pm0.0~J$	$75.3\pm2.2~\mathrm{L}$	$25.1\pm2.2~\mathrm{J}$
	REF	2000-250	$330.8\pm3.4~\text{KL}$	$322.3\pm1.5~\text{K}$	$0.4\pm0.1~J$	405.0 ± 0.0	$0.2\pm0.0~\text{KL}$	$81.7\pm2.7~JK$	18.6 \pm .,7 KL
		250-50	327.0 \pm . 2.6 KL	319.0 \pm . 3.6 K	$0.4\pm0.0\;J$	405.0 ± 0.0	$0.1\pm0.0\;L$	$85.3\pm1.2~\mathrm{J}$	$15.1\pm1.2~\text{L}$
		< 50	$325.2\pm2.,6\text{ KL}$	$319.3\pm3.8~\text{K}$	$0.4\pm0.1~J$	405.0 ± 0.0	$0.1\pm0.0\;L$	$85.6\pm1.6~J$	$14.7\pm1.6~L$

50% burn-off T, peak T and heat flows of visible exotherms as well as ratios between labile:total and stable:total heat flow (mean \pm SD) are represented followed by results of the analysis of variance. For a given fraction, etiher free LF (light fraction) occ LF or HF (heavy fraction) identical letters indicate equal means.



Fig. 6. Van Krevelen diagram of fractions. Results show H:C and O:C atomic ratios of LF (light fraction) isolated by the fractionation protocol. Squares illustrate free-LF (f-LF, encircled by solid line) and circles illustrate occluded LF (occ-LF, encircled by dashed line). The point color depends on studied soils, white = REF, grey = CHAR. Crosses show ratios for fresh PyOM or hand-picked (h.p.) century-old PyOM.

studied soils are mainly attributed to the presence of PyOM. Lower H:C ratios in HF of charcoal enriched soils than in reference soils demonstrate the presence of PyOM (C-rich) associated with minerals through S&C-size microaggregation or adsorption.

Similar C:N ratios between studied soils, despite generally higher C content in CHAR, indicate that larger amounts of N occur in charcoal enriched soils than in adjacent REF soils. This may result from significantly higher nitrate contents retained by the biochar (Hagemann et al., 2017) or N-rich organic compounds derived from root and microbial activities (Brodowski et al., 2005; Meier et al., 2017). Indeed, microbial-derived proteins and root exudates are known to strongly bind to minerals with possibilities for their polar chains to serve as binding sites (Kleber et al., 2007; Vogel et al., 2014; Poirier et al., 2018) and to interact with PyOM (Brodowski et al., 2005). In HF this is supported by C:N ratios similar to those expected for microbial-derived OM sorbed onto mineral surfaces ranging [7–14] (Aufdenkampe et al., 2001; Bell et al., 2014).

4.3. Thermal analysis confirms organo-mineral associations

The presence of charcoal in the bulk soil of kiln sites is distinguished from reference soils by three distinct exotherms (Fig. 4). These exotherms occur at higher temperatures than in reference thermograms and correspond to peak temperatures characterizing hand-picked charcoal particles (Hardy et al., 2017a). At first, fresh PyOM is characterized by a single sharp exotherm. With time, as the PyOM particle oxidizes from its surface inwards (Brodowski et al., 2005; Cheng et al., 2008a) the aromatic structures are broken down and functional groups appear (Plante et al., 2009). The occurrence of these functional groups decreases the thermal stability of PyOM as shown by the occurrence of a new exotherm at ${\sim}386$ °C (Fig. 4), which corresponds to the combustion of Orich compounds (Hardy et al., 2017b). Hardy et al. (2017b) suggested that Ca^{2+} acted as a thermal catalyst thereby resulting in a decrease of thermal stability of functional groups visible in thermograms as a decrease in peak temperatures towards more thermally labile OM. Interestingly, hand-picked PyOM is characterized by the occurrence of a

third peak (~528 °C) with higher thermal stability (Fig. 4) than fresh PyOM as previously reported by Hardy et al. (2017b). In this study authors showed an increase in this peak height with time of cultivation thereby suggesting that amounts of highly thermally stable C increase relatively to other, less stable forms which are preferentially altered or lost from the system. The third type of OM used as a reference curve is hand-picked non-PyOM which is characterized by the occurrence of both thermally labile (336 ± 2 °C) and thermally stable (442 ± 2 °C) OM (Fig. 4). These peak temperatures correspond to those of cellulose and lignin like molecules (Lopez-Capel et al., 2005). The occurrence of these exotherms in both studied soils highlight that PyOM in soils can be determined as the additional heat flow occurring at thermally stable temperatures in CHAR as opposed to REF soils.

In both studied soils, LF (i.e. occ. or free) are characterized by higher thermal stability than HF. Free LF in CHAR soils are characterized by a punctual heat flow comparable to that of pristine fresh PyOM which suggests little functionalization of PyOM particles in these fractions. Such a recalcitrance to oxidation for CHAR free LF is explained by a greater aromaticity or degree of condensation of the PyOM (Wiedemeier et al., 2015). Thermograms of REF free LF on the other hand show more diversity in the type of OM composing the fraction as shown by a multitude of peaks. In agricultural soils, after biomass inputs, it may be expected that free LF would be quickly deprived of small and energy rich forms of OM due to rapid mineralization of thermally labile organic compounds more beneficial to decomposers (Leifeld, 2008; Rovira et al., 2008; Barré et al., 2016). As a result, following the preferential decomposition of energy rich OM, free LF in CHAR soils are expected to be dominated by larger, more persistent molecules with higher binding energy and hence to be thermally more stable (Leinweber et al., 1992; Lopez-Capel et al., 2005; De la Rosa et al., 2008). This is supported by higher 50% burn-off temperatures in free LF compared to other fractions (Leifeld, 2008) (Fig. 5).

Occluded LF differ from free LF through the appearance of a second class of exotherms occurring around ~330 °C (Fig. 5). The occurrence of this thermally labile peak suggests an accumulation of energy-rich OM resulting from physical inaccessibility of this OM to decomposers and organo-mineral associations (Lehmann and Kleber, 2015). In occluded LF of REF, soils significant heat flows occur at high thermal stability. These exotherms coincide with peak temperatures attributed to PyOM (Fig. 5) which complicates a definite distinction between PyOM and non-PyOM (Leifeld, 2007). Such exotherms highlight the importance of studying CHAR soils in comparison to thermograms of REF soils to determine the PyOM pool as the additional heat flow occurring in CHAR soils. In CHAR occluded LF the presence of PyOM is mainly underlined by a slightly higher 50% burn-off temperatures (Fig. 5) and higher stable:total heat flow ratios (Table 4).

In HF the presence of an elbow occurring at \sim 410 °C in CHAR and not in REF soils strongly suggests the presence of PyOM tightly associated with minerals. Indeed, although these fractions are dominated by thermally labile OM (Fig. 5 - C – peaks \sim 330 °C), the sustained presence of PyOM in fractions dominated by microaggregation and adsorption suggests that century-old charcoal displays strong interactions with soil minerals (Chenu and Plante, 2006; Virto et al., 2010). The decrease in peak temperatures of PyOM originates from the creation of phenol, carbonyl and carboxyl functionalities at the surface of century-old charcoal particles (Hardy et al., 2017a). Our results support this assumption as this peak value in HF corresponding to PyOM (~410 °C) occurs between the two first peaks of hand-picked PyOM (p1 = 386 ± 2 $^{\circ}\text{C}, \text{ p2}$ = 443 \pm 4 $^{\circ}\text{C}\text{)}.$ Furthermore, because functional groups are needed for sorption to occur (Kleber et al., 2015) it is expected that PyOM particles tightly bound to mineral phases have been strongly oxidized through either biotic or abiotic processes (Cheng et al., 2006). As microaggregation and adsorption are the main stabilization mechanisms for OM in soils, it is expected that beyond its chemical recalcitrance, PyOM is further stabilized by organo-mineral associations (von Lützow et al., 2006; Dungait et al., 2012; Lehmann and Kleber, 2015) yet

that this additional physical protection is not discernable by thermal analyses (Plante et al., 2011).

4.4. A major part of PyOM is associated with mineral phases

DSC results show PyOM amounts for an important part of the total SOC [29.9-41.3%] in bulk charcoal rich soils. Due to the variability in chemical composition of PyOM no single method can correctly quantify it (Lorenz and Lal, 2014; Kerré et al., 2016). Our methodology consisted in estimating the amount of PyOM by comparing peak height ratios between CHAR and adjacent REF soil with respect to differences in C content and is therefore not a direct quantification. Furthermore, our quantification approach is developed on the assumption that no PyOM is found in REF soils. However, it may be expected that small amounts of PyOM could be found in reference soils suggesting PyOM amounts are slightly underestimated in charcoal rich soils. Using standard additions of PyOM to mineral soils free of PyOM, Hardy et al. (2017a) reported that charcoal-C content in kiln sites amounts for 79.5% of the difference in C content between CHAR soils and REF soils. Using this relationship on our charcoal rich soils we obtain an estimate of PyOM content ranging [25.3 – 39.2%], slightly lower than our estimate [29.9–41.3%]. Using DSC on isolated soil fractions enabled an understanding of the distribution of PvOM amongst soil fractions. Results show that after two centuries of residence in soils, PyOM is located in all soil fractions and that, although a large amount occurred as free LF (43.6 \pm 22.9% of total PyOM), the majority interacts with mineral phases (56.4 \pm 22.9% of total PyOM) through occlusion (29.6 \pm 28.8% of total PyOM) or sorption (26.78 \pm 5.9% of total PyOM) to mineral phases (Table 2).

Despite decades of incorporation in the soil matrix and sustained farming practices, results show that a large amount of PyOM is still found as free LF in CHAR soils. This result illustrates the importance of PyOM composition for its stability in soils and the role of charring against oxidative agents through its polycondensed aromatic structure (Wiedemeier et al., 2015) and against decomposers by making it a hardly decomposable form of OM (Rovira et al., 2008; Wang et al., 2016). Nonetheless, in the presence of specific decomposers, PyOM may decompose faster than often reported (Hamer et al., 2004). Recent studies show that the persistence of PyOM in soils may be overestimated (Lehndorff et al., 2014; Lutfalla et al., 2017; Hammes et al., 2008; N. Singh, et al., 2012). Nonetheless, given the high temperatures and low oxygen contents found in kiln sites, it may be expected that the stability of the charcoal produced in such environment would be higher than wildfire residues (Bird et al., 2015). In the studied CHAR soils, the presence of PyOM amongst occluded LF and OM in the HF shows that its stability is not exclusively explained by its chemical structure but further enhanced by a decreased accessibility for microorganisms (Brodowski et al., 2006; Lützow et al., 2006). Charcoal-mineral associations likely increase with PyOM ageing through increasing proportions of functional groups on the charcoal surfaces as well as the occurrence of biotic or abiotic binding agents (Brodowski et al., 2006; Kögel-Knabner et al., 2008; Kopittke et al., 2020). Finally, our results show increased amounts of non-PyOM in charcoal rich soils which suggests PyOM may influence the stability of non-PyOM in soils by serving as a hotspot for its sorption onto already existing organo-mineral clusters (Kopittke et al., 2020) or by creating microaggregates in which it becomes physically inaccessible to decomposers.

5. Conclusion

Our work combined for the first time thermal and elemental analyses on soil fractions extracted from a unique setting where PyOM from kiln sites was incorporated in conventionally cropped systems for over two centuries.

The C content of CHAR soils was significantly higher than that of adjacent REF soils thus confirming the potential of PyOM as a tool for C storage over centuries. Furthermore, our results show that such

increases in C content were not solely attributed to increases in PyOM content, but also to additional non-PyOM accumulation in CHAR soils. The presence of PyOM in all soil fractions showed that after centuries of incorporation in strongly anthropogenized environments pyrogenic matter still stands as free LF yet also strongly associates with mineral phases. This association occurs through occlusion into aggregates and sorption of PyOM onto mineral surfaces. Compared to fresh PyOM, century-old PyOM isolated from charcoal rich soils showed higher H:C and O:C atomic ratios particularly in occluded LF vouching for a functionalization of the PyOM. As such its reactivity with other soil constituents increases as suggested by increased Ca²⁺ concentrations in CHAR soils. Furthermore, similar C:N ratios in CHAR and REF fractions suggest increased amounts of N-rich organic products in charcoal rich soils which probably serve as binding agents between PyOM and mineral phases. As a result of more binding sites, we observed increased amounts of water stable macroaggregates in CHAR soils. The related improved soil structure in charcoal rich soils is fundamental in maintaining ecosystem services such as diminished erosion and enhanced plant primary production.

Our results highlight that beyond its chemical composition, PyOM is further stabilized by organo-mineral associations and can not only serve as a C sequestration tool but also as a mean to improve the soil quality of conventionally cropped agroecosystems in temperate Luvisols.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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