

# Is Rock-Eval 6 thermal analysis a good indicator of soil organic carbon lability? – A method-comparison study in forest soils

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## Abstract

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Soil respiration tests and abundance of particulate organic matter (POM) are considered as classical indicators of the labile soil organic carbon (SOC) pool. However, there is still no widely accepted standard method to assess SOC lability and the pertinence of these two timeconsuming methods to characterize SOC turnover can be questioned. Alternate ways of determining the labile SOC fraction are thus much needed. Thermal analyses, in particular Rock-Eval 6 (RE6) analysis has shown promising results in the determination of SOC biogeochemical stability. Using a large set of samples (n = 99) of French forest soils representing contrasted pedoclimatic conditions, including deep samples (up to 0.8 m depth), we compared three different methods used for SOC lability assessment. We explored whether respired-C isolated by a 10-week laboratory soil respiration test, POM-C isolated by a physical SOC fractionation scheme (particle-size  $> 50 \mu m$  and d  $< 1.6 g \cdot cm^{-3}$ ) and several RE6 parameters were comparable and how they correlated. As expected, respired-C (mg CO<sub>2</sub>-C·g<sup>-1</sup> SOC) and POM-C (% of total SOC) fractions strongly decreased with depth. RE6 parameters showed that SOC from deeper soil layers was also thermally less labile, more oxidized and H-depleted. Indeed, SOC from deeper soil layers had lower proportion of thermally labile SOC, higher T<sub>50 HC PYR</sub> (temperature at which 50% of the pyrolysable hydrocarbons were effectively pyrolyzed) and T<sub>50 CO2 OX</sub> (temperature at which 50% of the CO<sub>2</sub> gas had evolved during the oxidation phase), larger oxygen index, and smaller hydrogen index. Surprisingly, the two classical indicators of the labile SOC pool (respired-C and POM-C) were only marginally correlated (p = 0.051) and showed layerspecific correlations. Similarly, respired-C was poorly correlated to RE6 parameters. Conversely, the POM-C fraction showed a strong negative correlation with  $T_{50 \text{ HC PYR}}$  ( $\rho =$ -0.73) and good correlations with other RE6 parameters.

Our study showed that RE6 parameters were good estimates of the POM-C fraction, which represents a labile SOC pool with a residence time of *ca.* a couple decades that is meaningful regarding SOC stock changes upon modifications in land management. RE6 thermal analysis could therefore be a fast and cost-effective alternative to more time-consuming methods used in SOC pool determination, and may be integrated into soil monitoring networks to provide high-throughput information on SOC dynamics. **Keywords**: soil organic carbon kinetic pools; Rock-Eval 6; particulate organic matter; soil basal respiration; deep soil organic carbon; French forest soils;

## 1. Introduction

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Soil organic matter (SOM) degradation has multiple consequences on major soil functions like nutrients cycling, soil emissions of greenhouse gases and affects soil carbon sequestration potential. In particular, the labile part of SOM (turnover < 20 years) is associated with biological (microbial) activity and nutrient cycling (Haynes, 2005) and is very relevant to these issues. In that context, information on the temporal trajectories of SOC storage at a fine spatial resolution, in the form of detailed mapping of SOC stock evolutions with time for different land management scenarios, are required. SOC dynamics models are the logical candidates to provide such information, but their predictive performance is not yet satisfying, and they would benefit from an improved initialization using fine-scale information on SOC kinetic pools (Luo et al., 2016). Soil monitoring networks have become more prominent in the last twenty years. However, currently they can only provide information relative to the recent temporal (decadal) evolution of total SOC stocks. To use the full potential of these networks and measure the effects of climate and land-use changes on SOC stocks will require indicators of the size of the different SOC pools. Respiration measurements and particulate organic matter (POM) quantification obtained by various methods of fractionation (particle size only / density only / density + particle-size) (von Lützow et al., 2007) have been used for decades and are now classical estimates of the labile SOC pool. Laboratory incubations are run under optimum temperature and moisture conditions and use the indigenous microflora. They thus represent a maximum potential rate of C mineralization and an index of C availability in the system, integrating the physical, chemical, and microbiological properties of the soil (Haynes, 2005). Incubations are fairly simple to set-up

but they require space and are time-consuming. Sieving and rewetting also tend to artificially increase the mineralizable pool (Haynes, 1986). Physical fractionation schemes are easy to implement and do not require expensive equipment although they can become costly when density fractionation is involved. Moreover they are very time-consuming, often requiring multiple and relatively long periods of agitation/settling and drying. The most important limitation is the ability of the fractionation scheme to isolate physical fractions that have homogeneous turnover and thus represent functional noncomposite SOC pools (von Lützow et al., 2007). While respired-C and POM-C fractions both represent a labile SOC pool, the former corresponds to a smaller SOC pool with a shorter mean residence time (usually < 1 year for temperate *in-situ* conditions) (Feng et al., 2016) while the latter corresponds to a larger SOC pool with a longer mean residence time (usually < 20 year for temperate *in-situ* conditions) (e.g., Trumbore et al., 1996; Balesdent, 1996). Because these two methods are both very timeconsuming, they cannot address the needs of soil monitoring, i.e., a methodology that is informative, high-yield and relatively cheap to implement, to allow for the analysis of numerous samples. Among thermal analyses used to characterize SOM, Rock-Eval 6 (RE6) analysis has shown promising results in the determination of SOM biogeochemical stability (e.g., Barré et al., 2016) and thus appears like a good candidate to fill this methodological gap. Originally developed for oil and gas exploration in sedimentary basins, the method was first applied to study soils with hydrocarbons contamination (Lafargue et al., 1998). RE was also shown to provide useful information on SOM originating from soil profiles worldwide (Disnar et al., 2003) and many studies on SOM characterization have been conducted, sometimes using RE analysis in conjunction with other methods like nuclear magnetic resonance (Albrecht et al., 2015), hydrocarbon analysis by gas chromatography (Di-Giovanni et al., 1998), infrared

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spectroscopy (Hetényi et al., 2006) or biochemical oxygen demand (Copard et al., 2006) to determine the origin and/or decomposition stage of the organic matter (Hetényi et al., 2005; Sebag et al., 2006). More recently, RE6 results have been compared with respiration test or SOM fractions at the plot (Gregorich et al., 2015) and the small landscape scale (Saenger et al., 2015) but in both cases the analyses were restricted to superficial soil layers. Gillespie et al. (2014) have also related thermal stability assessed by RE6 to respiration test and X-ray absorption near-edge structure spectroscopy for cryosolic soil profiles (up to 75 cm) in Northern Canada but only in four hummocks. Finally, RE6 thermal analysis has been used to look at SOM dynamics in a sample set with a large soil type variability and some deeper horizons (Sebag et al., 2016), but without comparison to other methodologies. The objective of this study was to properly "benchmark" RE6 thermal analysis with two classical yet time-consuming methods for labile SOC pool estimation: a soil respiration test isolating a respired-C fraction under controlled laboratory conditions and a physical SOC fractionation scheme isolating a POM-C fraction. We selected soil samples from the French forests monitoring network RENECOFOR at various depths. To our knowledge, this is the first study considering such a large set of samples (covering a wide pedoclimatic variability), including deep soil layers up to 0.8 m. Our sample set thus included soil samples that presumably contained very different proportions of the labile SOC pool. Because the difference in size of the C pool estimated by the respiration test and the POM fractionation (e.g., Haynes, 2005) and the previously observed correlations between stock of labile SOC estimated by RE6 parameters and the POM fraction (Saenger et al., 2015) on one hand and between cumulative C mineralized and a RE6 parameter (Gregorich et al., 2015) on the other hand, we were expecting that: 1/ the results provided by the two classical methods would differ quantitatively while the results from the three methods would be qualitatively

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comparable and correlated; 2/ we would be able to derive a significant quantitative relationship between RE6 parameters and the two classical indicators of the labile SOC pool.

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## 2. Material and methods

2.1. Sampling

We considered forest soils samples from 53 permanent forest sites of the French national network for the long term monitoring of forest ecosystems ("RENECOFOR"), established in 1992 (Ulrich, 1995) by the National Forest Service (ONF; http://www.onf.fr/renecofor) as a part of the European ICP-FORESTS (http://icp-forests.net/) level 2 network (Fig. 1a). They were representative of a good portion the national variability in terms of climate (with MAP and MAT ranging between 703–1894 mm and 4.8–12.3 °C respectively for the 1971–2000 period), soil type (entic Podzol; eutric Cambisol/Calcisol; dystric Cambisol) (IUSS Working Group, 2015) and forest vegetation (coniferous—silver fir; Norway spruce; European larch; Scots pine—and deciduous—beech; oaks *spp*—stands). At each site, samples representing two soil layers were obtained (0–10 cm and 40–80 cm; Fig. 1b). Samples of the top soil layer were composite, at each depth, of  $5 \times 5$  sampling points located over a 5000 m<sup>2</sup> plot, collected between 2007 and 2012 by digging a 50 cm wide soil profile (Ponette et al., 1997; Jonard et al., 2017). Samples of the deeper soil layer were composite from two soil pits located just outside the central plot and collected in 1994–1995 (Brêthes et al., 1997). The surface and deep samples thus originate from two different sampling campaigns. The deep samples were only collected once, during the first campaign, to limit perturbation on the monitoring plots. Basic soil parameters (pH and texture) were determined by Ponette et al. (1997) and are reported as supplementary information (Table SI-A1). Bulk soils were air-dried and stored in plastic buckets right after sampling. One liter of soil of each layer was retrieved for this study and sieved at 2 mm before analysis.

2.2. Elemental analysis

Bulk < 2 mm-sieved soil samples were ground (< 250  $\mu$ m; ultra-centrifugal mill ZM 200, Retsch Gmbh) and organic carbon and total nitrogen concentrations were determined by dry combustion with an elemental analyzer (CHN NA 1500, Carlo Elba). Samples with carbonates (total CaCO<sub>3</sub> = 3.5–835 g·kg<sup>-1</sup>) were first decarbonated following the same protocol as Harris et al. (2001). Briefly, 30 mg of ground samples were weighed in 5 mm × 9 mm silver boats followed by the addition of 50  $\mu$ L of distilled water. The boats were put in a glass bell jar, next to a beaker containing 100 mL of concentrated HCl (12 mol·L<sup>-1</sup>). The air in the jar was evacuated and samples let to sit in this HCl-saturated atmosphere to allow the acid to dissolve water and hydrolyze the carbonates for 8 h. Then, the decarbonated samples were dried at 60 °C in the silver boats for at least 48 h. Silver boats were further placed in 10 mm × 10 mm tin boats and analyzed for C and N.

POM fractions (see section 2.4.) were ground with a ball mill (mixer mill MM 200, Retsch Gmbh) or a mortar and pestle when the sample mass was less than 0.05 g. Carbon concentration was determined as for the bulk soil.

#### 2.3. Respiration test

For each sample, 20 g of 2 mm-sieved soil were transferred in a 120 mL glass-flask and rewetted at pF 2.5 (-0.033 MPa), which had been previously determined using a 5 Bar pressure plate extractor (#1600, Soilmoisture Equipment Corp.). The flasks were fitted with aluminum seals with PTFE-faced silicone septa to allow for headspace gas sampling and placed inside an incubator (AE240 BIO EXPERT, Froilabo SAS) kept at 20 °C for 10 weeks following a two-week period pre-incubation to allow the samples microbial activity to stabilize (data not included).

- Headspace gases were sampled at 1 to 2-week intervals during the 10-week incubation period
- and CO<sub>2</sub> concentrations were determined using an Agilent 490 micro-gas chromatograph
- equipped with the OpenLAB Chromatography Data System EZChrom software.
- When CO<sub>2</sub> concentrations had reached 2.5–3% or was expecting to do so before the next
- measurement, and/or when the cap had been pierced with the needle four times, flasks were
- opened and flushed with fresh and moist air to return CO<sub>2</sub> concentrations to ambient levels to
- avoid anoxia (while maintaining the moisture content), before returning them to the incubator.
- The CO<sub>2</sub> concentrations measured by the GC were converted in μ CO<sub>2</sub>-C·using equation 1:
- 186  $\mu g C-CO_2 = mmol air \times ppm CO_2 (\mu mol C/mol air) \times 10^{-3} (mol/mmol) \times 12 (\mu g C/\mu mol C)$
- 187 (equation 1)

- where "mmol air" corresponds to the millimoles of air present in the flask and was calculated
- with the ideal gas law (equation 2):
- 190  $n = PV / RT = (1 \times 100) / (82.05 \times 293)$  (equation 2)
- 191 As a result, we multiplied our concentrations of CO<sub>2</sub> expressed in percent by 499.16 to
- 192 convert them in μg C-CO<sub>2</sub>.
- 193 Finally, the 10-week mineralizable SOC (respired-C) was expressed in mg CO<sub>2</sub>-C·g<sup>-1</sup> SOC to
- account for differences in the C content of the various layers and sites.
- 196 2.4. Particle size and density SOC fractionation
- To isolate the particulate organic matter (POM) fraction, samples were first dried at 50 °C for
- 198 24 h before weighing 25 g and transferred them in polyethylene (PE) 250 mL flasks. We then
- added 180 mL of 0.5% sodium hexametaphosphate solution and ten 5 mm-diameter glass
- beads before shaking the samples overnight (50 rpm; 16 h) on an overhead shaker (Reax 2,
- Heidolph), in order to breakdown soil aggregates. Samples were thoroughly rinsed over a 50-
- 202 μm mesh with deionized water. The > 50 μm fraction was then transferred back to a dry PE

flask with a sodium polytungstate (SPT) solution of density =  $1.6 \pm 0.03$  g·cm<sup>-3</sup> (Golchin et al., 1994; Crow et al., 2007) and the solution was added up to *circa* 180 mL. The flasks were shaken overhead by hand 10 times and samples were left overnight to settle down after the cap of the flask was rinsed with the SPT solution. The floating material was collected with a spatula and placed over a 50- $\mu$ m mesh sieve. If necessary some SPT solution was added back to the flask and the previous step was repeated. This time, samples were placed in a centrifuge for 30 minutes to accelerate the separation (2750 rpm or 1250 g, Six et al., 1998). The floating material was again collected with the spatula or pipetted depending on the amount left. This step was repeated if the light fraction was abundant. If not, samples were left to settle down overnight before one last collection. The POM fraction on the sieve was thoroughly rinsed with deionized water throughout the whole process. The sieves and fractions were then placed in the oven at 50 °C for 24 h before being weighed. To account for differences in the C content of the different samples, we calculated the proportion of OC in the POM fraction (POM-C), expressed in g POM-C·g<sup>-1</sup> total SOC.

## 2.5. Thermal analysis: Rock-Eval 6

The thermal analysis of the samples was performed with a Rock-Eval 6 turbo device (Vinci Technologies, France). Details about the equipment have been previously published (Behar et al., 2001). We adapted the procedure developed for the analysis of soil organic matter by Disnar et al. (2003). Briefly, about 60 (20.7–62.1 depending on the sample's C content) mg of ground sample were exposed to two consecutive thermal treatments, first in a pyrolysis oven (200–650 °C; thermal ramping rate of 30 °C·min<sup>-1</sup>; under N<sub>2</sub> atmosphere) then in a combustion oven (300–850 °C; thermal ramping rate of 20 °C·min<sup>-1</sup>; under laboratory air atmosphere). At the beginning of the pyrolysis, there was an isothermal step (at 200 °C) during 180 seconds during which the free hydrocarbons (HC) were thermovaporized (S1

228 peak). The pyrolysis effluents (mostly HC) were detected and quantified with flame ionization 229 detection, while CO and CO<sub>2</sub> were quantified by infrared detection during both the pyrolysis 230 and oxidation stages (Fig. SI-A1). 231 Two standard RE6 parameters describing SOC bulk chemistry were determined: the hydrogen 232 and oxygen index values (HI and OI<sub>RE6</sub>). The HI index corresponds to the amount of 233 hydrocarbons formed during thermal pyrolysis of the sample (HC evolved between 200 and 234 650 °C minus the S1 peak) divided by the total SOC content of the sample and is expressed in mg HC·g<sup>-1</sup> SOC. It describes the relative enrichment/depletion of SOC in hydrogen-rich 235 236 moieties. The OIRE6 index describes the relative oxidation status of SOC. It was calculated 237 using the equation proposed by Lafargue et al. (1998):  $OI_{RE6} = 16 / 28 \times OI_{CO} + 32 / 44 \times OI_{CO2}$ 238 (equation 3) 239 Where OI<sub>CO2</sub> corresponds to the CO<sub>2</sub> yielded during thermal pyrolysis of the sample between 240 200 and 400°C divided by the total SOC of the sample and OI<sub>CO</sub> corresponds to the CO 241 yielded during thermal pyrolysis between 200 and 400–650°C (wherever a minimum of CO 242 production is observed; in the absence of a minimum, the default upper-limit temperature is set at 550 °C) divided by the total SOC of the sample. Thus OI<sub>RE6</sub> is expressed in mg O<sub>2</sub>·g<sup>-1</sup> 243 244 SOC. 245 We derived four additional RE6 parameters describing the thermal stability of SOC: (i) 246 T<sub>50 HC PYR</sub>, the temperature at which 50% of the HC resulting from the SOM pyrolysis had 247 evolved (ii) the T<sub>50 CO2 OX</sub>, the temperature at which 50% of the residual SOM was oxidized 248 to CO<sub>2</sub> during the oxidation phase. Because the signal was noisy at the beginning of the 249 pyrolysis, we started the integration for T<sub>50 HC PYR</sub> right after the S1 peak. For T<sub>50 CO2 OX</sub>, the 250 upper limit temperature for signal integration was set at 611 °C to obtain a total CO<sub>2</sub> signal 251 evolved from pure OM without interference of carbonates. Both these T<sub>50</sub> temperature

parameters and the HI index have been previously shown as good thermal indicators of SOM

253 biogeochemical stability (Gregorich et al., 2015; Barré et al., 2016). We also included two 254 thermal indices previously used in the literature: the (iii) R-index or (1 – R400), which 255 correspond to the integrated area of the HC thermogram above 400 °C over the total area of 256 the HC signal (Disnar et al., 2003; Sebag et al., 2016). The R-index estimates the proportion 257 of thermally stable SOC pool and varies between 0 and 1. We hypothesized that the 258 proportion (1 – R-index) would approximate a thermally labile/intermediate (turnover < 20 259 years) SOC pool. Finally, using equation 4, we calculated the (iv) I-index, which is an 260 indicator of the preservation of thermally labile immature SOM (Sebag et al., 2016): 261  $\log_{10}((A1 + A2) / A3)$ (equation 4) 262 where A1 + A2 corresponds to the integrated area of the HC thermogram below 400 °C and 263 A3 the integrated area of the HC thermogram between 400 °C and 460 °C. 264 Signal processing of the RE6 thermograms (signal integration and calculation of the 265 T<sub>50 HC PYR</sub>, T<sub>50 CO2 OX</sub>, R and I indices) was performed with the R environment software v.3.3 266 (R Core Team, 2016) using the hyperSpec (Beleites and Sergo, 2015) and pracma (Borchers, 267 2015) R packages.

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2.6. Calculations and statistical analyses

For RE6 analysis and the respiration test, samples with very low C content (< 0.2%) were not considered as the carbon flux they produced during the incubation or the thermal analysis was too low/too close to the limit of detection for reliable determination. This resulted in the selection of n = 46 for the soil layer 40–80 cm (total n = 99). The mean values of the variables derived from the SOC respiration test, fractionation and RE6 analysis for all layer depths were compared using standard non-parametric statistical methods such as Kruskal Wallis test one-way ANOVA by ranks and Wilcoxon signed-rank test. Relationships between the variables derived from the three methods were estimated using

Spearman rank correlation as the data did not meet the assumption of normality. Correlation tests were first performed on the whole dataset (n = 99) then within the 0–10 cm and the 40–80 cm layers, the three soil types and the two vegetation types individually. All comparisons were considered significant at an alpha value ( $\alpha$ ) of 0.05. A principal component of analysis (PCA) was performed to detect linear relations between parameters derived from the 3 methods. For that purpose, data were log-transformed, centered and scaled. Because the I-index was negative in some instances, we added the equivalent of the smallest I-index value + 0.2 to all the I-index values to run the PCA. To determine the number of principal components to select, we looked at the percentage of the total variance explained and used a scree plot and Kaiser's criterion. To analyze the relationship between RE-based and the two classical indicators of the labile SOC pool, we used a simple linear regression model and relied on the Cook's distance to identify potential outliers. All statistical analyses were performed using R 3.3 (R Core Team, 2016) using the factoextra (Kassambara and Mundt, 2016) and Hmisc (Harrell et al., 2016) packages.

#### 3. Results

3.1. Respiration test

The 10-week mineralizable SOC (respired-C) was expressed in mg CO<sub>2</sub>-C·g<sup>-1</sup> SOC to account for differences in the C content of the various layers and sites. Over the course of the 10-week incubation, the surface layer (0–10 cm) samples cumulatively respired on average 17  $\pm$  7.2 mg CO<sub>2</sub>-C·g<sup>-1</sup> SOC, while the deeper layer (40–80 cm) samples respired 13.4  $\pm$  6.9 mg CO<sub>2</sub>-C·g<sup>-1</sup> SOC (Table 1). There was a significant decrease in respired-C with depth (p = 0.003), indicating a smaller size of the labile C pool in the deeper layers of our forest soils. Within each soil layer, the large standard deviation (around 7.0 mg CO<sub>2</sub>-C·g<sup>-1</sup> SOC) illustrates an important inter-site variability.

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304	3.2. POM fractionation
305	The POM-C fraction (% of total C) decreased by half between layers 0–10 cm and 40–80 cm
306	with 22.6 $\pm$ 7.3% and 11.0 $\pm$ 6.1% respectively. This indicates a significantly ( $p$ < 0.001)
307	smaller labile C pool in the deeper (40–80 cm) soil layer. POM-C ranged between 12.1–
308	43.0% and 2.5–33.6% in the 0–10 cm and 40–80 cm layers, respectively, illustrating again an
309	important inter-site variability.
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311	3.3. RE6 thermal analysis
312	There was a significant effect of depth on all RE6 parameters. Particularly, the two $T_{50}$
313	parameters increased significantly ( $p < 0.001$ ) with depth: $421 \pm 9$ °C to $448 \pm 10$ °C and $399$
314	$\pm9$ °C to 431 $\pm18$ °C (Table 1), for $T_{50\_HC\_PYR}$ and $T_{50\_CO2\_OX}$ respectively, corresponding to
315	an increase in the thermal stability of total SOC (i.e. a relative decrease in the labile C pool
316	and increase of the stable C pool). OI <sub>RE6</sub> showed a similar increasing trend ( $p < 0.001$ ) with
317	depth (225 $\pm$ 37–439 $\pm$ 138 mg $O_2 \cdot g^{-1}$ total SOC; Table 1), reflecting a more oxidized SOC in
318	the deeper layers. Conversely, HI decreased significantly ( $p < 0.001$ ) with depth (276 $\pm$ 77–
319	$133 \pm 34$ mg HC·g <sup>-1</sup> total SOC; Table 1), suggesting a relative depletion of total SOC in H-
320	rich moieties with increased soil depth. The proportion of thermally stable SOC R-index, also
321	experienced a significant increase ( $p < 0.001$ ) with depth (59–69%; Table 1), while the I-
322	index decreased slightly (0.17–0.11; Table 1).
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324	3.4. Correlations between methods
325	3.4.1. For all samples
326	There were mainly significant and strong correlations between POM-C and the RE6

parameters (Table 2). Notably  $T_{50\_HC\_PYR}$ ,  $OI_{RE6}$  and R-index all had a strong negative

correlation with POM-C (Spearman  $\rho = -0.73$ , -0.76 and -0.69 respectively; Table 2; Fig. 3).  $T_{50 \text{ CO2 OX}}$  and HI moderately correlated with POM-C ( $\rho = -0.56$  and 0.67) and the I-index had a weak positive relationship with POM-C ( $\rho = 0.35$ ). I-index,  $T_{50 \text{ HC PYR}}$  and R-index were the only parameters that were significantly related to respired-C, with a weak correlation  $(\rho = 0.32, \rho = -0.26 \text{ and } -0.31 \text{ respectively}; \text{ Table 2})$ . The two classical methods of estimation of labile SOC (respired-C and POM-C) were weakly positively ( $\rho = 0.20$ ; Table 2; Fig. SI-B1 a) and indeed only marginally (p = 0.051) related. To describe the similarity or dissimilarity in the different indicators of SOC lability, we conducted a principal components analysis (PCA). As shown by the correlation test, T<sub>50</sub> HC PYR and R-index on the one hand and OI<sub>RE6</sub> and HI on the other hand were highly correlated ( $\rho = 0.93$  and -0.92 respectively; Table 2). We thus decided to conduct the PCA using only the 6 following explanatory variables = respired-C; POM-C; HI; T<sub>50</sub> CO<sub>2</sub> OX; T<sub>50</sub> HC PYR; I-index). The first two principal components (PC) explained approximately 73% of the total variance, with 53% explained by the first and 20% explained by the second PC, respectively (Fig. 2). PC1 clearly separated surface (0–10 cm) from deeper (40–80 cm) soil samples. Along PC1, POM-C and HI showed moderate negative loadings (-0.47 and -0.46 respectively; Table SI-B1) while T<sub>50</sub> HC PYR and T<sub>50</sub> CO2 OX had moderate positive loadings (0.53 and 0.45; Table SI-B1). Respired-C and the I-index showed strong positive loadings along PC2 (0.55 and 0.69; Table SI-B1), while they showed very weak negative loadings along PC1. Samples from layers 0–10 and 40–80 cm did not significantly differ along the

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3.4.2. For the 0-10 cm and 40-80 cm layer separately

These global correlations prompted us into looking at the influence of soil depth on the different parameters. The paired correlations between the 8 parameters differed in surface (0–

10 cm) and deep (40–80 cm) layers (Table 2). Specifically, the respired-C in the surface layers was weakly and negatively related to POM-C ( $\rho = -0.29$ ; Table 2). Conversely in the deep layers, respired-C and POM-C were moderately and positively correlated, as it would be expected ( $\rho = 0.47$ ; Table 2; Fig. SI-B1 a). In the surface layers, HI and OI<sub>RE6</sub> were also moderately (negatively and positively, respectively; Table 2) correlated to respired-C, while in the deep layers we observed again this negative and moderate correlation between T<sub>50 HC PYR</sub> and respired-C. For POM-C, we found the same negative correlations with  $T_{50~HC~PYR}$  and  $OI_{RE6}$  as in the "all samples" comparison but they were less strong ( $\rho = -0.35$ to -0.42; Table 2). In the surface layer, the C/N ratio, pH and clay content had all moderate and significant correlations with respired-C and T<sub>50 HC PYR</sub> (Table 2). These correlations were absent in the 40–80 cm layer. We also looked at the evolutions of the correlations as a function of vegetation and soil types, but there were no change as drastic as the ones we observed with depth (Table SI-C1). In both cases the changes affected only the correlations between respired-C and the other parameters. For instance, in coniferous plots, respired-C was weakly to moderately positively correlated to clay content ( $\rho = 0.27$ ) and pH (0.37) while those correlations were absent in deciduous plots (Table SI-C1). For the soil types, POM-C and respired-C were moderately and positively correlated in Podzols (0.42) and eutric Cambisols (0.46) but not in dystric Cambisols. Furthermore, in eutric Cambisols, respired-C was moderately and negatively correlated with  $T_{50 \text{ CO2 OX}}$  (-0.54), R-index (-0.50) and pH (-0.57; Table SI-C1).

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## 4. Discussion

4.1. Relationships between respiration test and POM fractionation

and marginally when considering all our samples.

Unexpectedly the two classical indicators of the labile SOC fraction correlated only weakly

POM-C is considered as a labile SOC fraction (Wander, 2004; Haynes, 2005; von Lützow et al., 2007), and we thus expected it would correlate significantly and strongly with the respired-C fraction isolated by the 10-week laboratory respiration test. Indeed, in his review, Haynes (2005) mentioned several studies reporting a positive and usually strong correlation between the respired-C and the POM-C fractions, appearing to support that hypothesis. However when carefully considering these papers (Janzen et al., 1992; Hassink, 1995; Campbell et al., 1999a; Campbell et al., 1999b; Wander and Bidart, 2000) and others (Liang et al., 2003; Hassan et al., 2016; Li et al., 2016), it emerged that the presented data were not normalized by the total SOC concentration of the samples. Without normalization it could be argued that the positive correlation between the POM-C and the respired-C fractions was in fact driven by variations in total SOC concentration and not SOC biogeochemical stability. It also prevented comparisons among studies, given the important difference in SOC concentration. The hypothesis of a positive correlation between the sizes of the labile SOC pool estimated by respiration test and POM fractionation schemes has actually not been properly tested on multiple sites, using SOC normalized data as it has been done in the present study. Indeed, the few studies that have reported moderate to strong positive correlations between the sizes of the labile SOC pool estimated by respiration test and POM fractionation were conducted on similar soils under different management (e.g., Alvarez and Alvarez, 2000) or correlations were made within sites (e.g., Janzen et al., 1992). When combining results from all sites, the correlation appeared to be weaker and it can therefore be hypothesized that in our study the weak and marginally significant correlation between POM-C and respired-C was partially due to the large inter-sites variability of soil properties for our sample set (Table SI-A.1). Finally, the labile SOC pools estimated by the two classical methods were so different in size (i.e. the labile SOC pool estimated as respired-C was about an order of magnitude smaller

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than the one estimated as POM-C; Table 1) that it is not surprising that the correlation did not hold specifically when introducing a lot of inter-sites variability. This constitutes another explanation to the lack of correlation between these two indicators of the size of the labile C pool. The two methods appeared to measure different SOC fractions (*i.e.* different sizes) (Table 2 and Fig. 2) that correspond to different SOC lability (*i.e.* mean residence time).

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4.2. Relationships between RE6 parameters and POM-C and respired-C Our RE6 results agreed with previous observations of thermal indicators of SOC lability. For instance, Sebag et al. (2016) reported a trend of decreasing HI and increasing OI<sub>RE6</sub> with soil depth. Trends of decreasing HI and increasing T<sub>50 CO2 OX</sub> were observed with increasing time since beginning of bare fallow experiments, which corresponded with a progressive decomposition of the labile SOC pool (Barré et al., 2016). Our correlations between the RE6 parameters and the POM-C fraction were close to those previously reported by Saenger et al. (2015). They indeed obtained a moderate positive correlation ( $R^2 = 0.50$ ) between the labile SOC pool stocks derived from a SOC fractionation scheme isolating POM-C, and the thermally labile SOC pool stocks derived from RE6 indices. We found a similar strong positive correlation between the proportion of labile SOC (1-R-index) and POM-C. The strong relationship between T<sub>50</sub> HC PYR and R-index could likely be explained as T<sub>50 HC PYR</sub> for our samples were very close to the 400 °C threshold used for the calculation of the R-index. As hypothesized we were able to derive a quantitative relationship between some of our RE6 parameters and POM-C (Fig. SI-B1 b-d). The best model was obtained for  $T_{50~HC~PYR}$  ( $R^2 = 0.52$ ; Fig. 3), while HI, R-index and  $OI_{RE6}$  were still moderately good predictors of POM-C ( $R^2 = 0.42-0.47$  (Fig. SI-B1 b-d). Nevertheless no strong relationship between respired-C and the other parameters could be established. Our correlations between the RE6 parameters and respired-C were smaller than

those previously reported by Gregorich et al. (2015). This could be explained by the fact that their study was, by design, very restricted in terms of its soil properties variability and also only considered surface soils (0–10 cm), in which the C/N ratios were around 10. Previous studies have also demonstrated that RE6 can be used to look at changes in the size of the SOC labile pools with time. For instance, RE6 was able to describe the decrease in the labile SOC pool in long-term bare fallows (Barré et al., 2016). Besides, RE6 captured differences in the size of the labile SOC pools in various land-uses and soil types over a small landscape (Saenger et al., 2015). Our results thus contradict the conclusions from Schiedung et al. (2017) who found no relationship between the thermally labile SOC (200–400 °C) and the C in the POM fractions. The latter (free and occluded POM—obtained by sonication) were indeed more stable at lower oxidation temperatures (300–350 °C) than the mineralassociated fraction. However, their analytical method was different from RE6 protocols: the thermal analysis they used was entirely realized under aerobic conditions (oxidation only), their temperature range was limited (only up to 400 °C) and they used a 50–100 °C temperature step every 15 minutes rather than a constant thermal ramping rate (standard in most thermal studies). For all these reasons, it is likely that their thermal indices differ greatly from our RE6-derived parameters. Moreover their study was based on topsoils (0–10 cm) of only three study sites. The good approximation of the POM-C fraction by RE6 we reported constitutes a very promising result. POM-C mean residence time (< 20 years in temperate conditions in the absence of an important charcoals contribution; e.g., Trumbore and Zheng, 1996; Balesdent, 1996; Balesdent et al., 1998; Baisden et al., 2002; Schrumpf and Kaiser, 2015) and its size (11 to 23% of total SOC in this study) are much larger than the one of the respired-C fraction, and is thus more meaningful regarding SOC stock evolutions upon changes of land management.

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This suggests that RE6 could be used to determine the size of the labile SOC pool with a decadal mean residence time.

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4.3. Effects of depth on correlations between the three methods estimating labile SOC Labile SOC content usually decreases with depth (e.g., Lorenz and Lal, 2005; Jenkinson et al., 2008). Such a trend was observed with the three methods used in the present study. Indeed, with depth, we observed a decrease in respired-C (respiration test), POM-C (POM fractionation) and HI alongside with an increase in T<sub>50 HC PYR</sub> and R-index that all signified the expected decrease in the size the labile SOC pool. Concurrently, OI<sub>RE6</sub> increased with depth, confirming the increase in SOC oxidative state with increasing decomposition (Hockaday et al., 2009; von Lützow and Kögel-Knabner, 2010; Hockaday et al., 2015). But more importantly, depth affected the correlations between the methods. The lack of correlation between two classical indicators of the labile SOC fraction previously mentioned appeared to originate from opposite trends in the surface and deep layers. In the 0–10 cm layer we observed a surprising negative (but weak) correlation between respired-C and POM-C while the expected positive and moderate correlation between the two indicators was found only in the deep layers. The differences in the sign of the correlations between respired-C and POM-C in the two considered layers (0–10 cm and 40–80 cm) may be related to pedological factors that can limit SOC mineralization in surface horizons. Indeed, the high C/N ratio found in the surface layer (Table SI-A1) is far from the expected C/N of the microorganisms and this lack of N may limit SOC respiration. Similarly, surface layers are on average more acidic (Table SI-A1) than deep layers which can also reduce SOC respiration. We could hypothesize that respired-C and POM-C correlate only when environmental conditions do not limit SOC mineralization explaining the absence of correlation in the acidic N-poor 0–10 cm layer. The significant correlations observed between respired-C and the C/N ratio, pH and the

clay content in the surface layer (Table 2) supports that hypothesis. This opposite behavior in the two layers also affected T<sub>50 HC PYR</sub>, which was not significantly correlated to respired-C in the surface layer while the two parameters were moderately and negatively correlated in the 40–80 cm layer (Table 2). These observations matched those from Peltre et al. (2013) who reported conflicting relationships between the parameter DSC-T50 (temperature at which half of the energy is released in differential scanning calorimetry) and mean soil respiration rates in two sets of high and low SOC content. Their DSC-T50 values were indeed negatively correlated with the respiration values for the low-C soils, whereas there was only a marginal positive correlation between the two parameters for the high-C soils. Their two groups were characterized by soil properties similar to our 0–10 and 40–80 cm layers: their low-C set consisted of samples with a higher pH and lower mean C/N ratio than those of the high-C. Similarly to our 0–10 cm samples, soils in their high-C set had a greater C concentration than those in the low-C set for similar clay contents (Table SI-A1). This would also explain why our results differ from those of Gregorich et al. (2015). In the deep layer, in which the C/N ratios are closer to those reported by Gregorich et al. (2015), we observed the same positive correlation they reported albeit less strong. Vegetation and soil types did not seem to have affected the correlations between the three methods we tested as much as depth did. However, these environmental factors are likely drivers of the size labile SOC pool as they have been shown to significantly influence RE6 parameters (e.g., Disnar et al., 2003; Sebag et al., 2006).

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4.4. Towards high-throughput information on SOC biogeochemical stability using RE6 analysis

Respiration tests and POM fractionation schemes are both time consuming, thus limiting the number of samples and/or replicates that are analyzed. With the RE6 set-up used in this study,

about 20 samples per day can be analyzed, and it requires only limited operator interventions (soil weighing and routine supervision of the RE6 analyzer). The lack of normalization in many studies using respiration tests and POM fractionation is an important issue and it should be recommended for further studies to include normalized data (% of TOC) when presenting their results. Moreover, despite the fact that POM-C and respired-C are considered as standard estimates of the labile SOC pool, the temperature and/or duration of incubations often varied from one study to the other. Similarly for the POM-C fraction, the density of the solution used for the flotation may drastically differ among studies. This makes data comparison almost impossible. In that regard, while the harmonization of RE6 programs would probably be much easier to implement than respiration tests or POM fractionation protocols as the number of users is still limited, protocol standardization is an important and pressing goal to achieve and this rather quickly as the method starts to gain interest. RE6 analysis is thus a rapid technique that captures differences in the labile SOC pool as well as other classical techniques. While the understanding of the underlying processes linking SOC thermal stability observed with RE6 and the laboratory or *in-situ* biogeochemical stability of SOC is not fully uncovered and further studies are needed, RE6 analysis appears like a very promising method to provide quick and inexpensive information on the labile SOC pool. Hence, it could constitute a standard method to complement C stock measurements in soil monitoring programs.

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- 690 pp. 67–102.

# Figure captions

- **Fig. 1**. (a) Location of the 53 study sites front the French national network for the long term
- 694 monitoring of forest ecosystems (RENECOFOR); (b) Number of samples by depths and
- analyses realized. Plot locations are also available via the Interactive Map Viewer.

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- 697 Fig. 2. Biplot of a principal components analysis (PCA) showing the loadings of the 6
- 698 parameters estimating the labile SOC (red arrows) and the 99 soil samples for the two layers
- (0-10 cm, n = 53; 40-80 cm, n = 46) along the first two principal component axes (PC1 and
- 700 PC2). The 95% ellipses for both soil layers were added for information; the circle in the
- center corresponds to the circle of correlations.

- Fig. 3. The proportion of OC in the POM fraction (POM-C) as a function of  $T_{50 \text{ HC PYR}}$  (the
- temperature at which 50% of the HC pyrolysis effluents have evolved) for all samples (n =
- 705 99; surface = 0-10 cm and deep = 40-80 cm).

# Figure captions

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Fig. 3. The proportion of OC in the POM fraction (POM-C) as a function of  $T_{50\_HC\_PYR}$  (the 718

temperature at which 50% of the HC pyrolysis effluents have evolved) for all samples (n =

720 99; surface = 0-10 cm and deep = 40-80 cm).

Table 1. Mean (and minimum; maximum; standard deviation) of the RE6 (HI,  $OI_{RE6}$ ,  $T_{50\_HC\_PYR}$ ,  $T_{50\_CO2\_OX}$ , R-index, I-index), respiration test (10-week mineralizable C, respired-C) and POM fractionation (POM-C) parameters, as well as the bulk SOC content for each soil layer (0–10 and 40–80 cm) of the 53 RENECOFOR plots.

	n		ng HC FOC)	O	E6 (mg 2 / g OC)	_	HC_PYR	_	co2_ox °C)		ndex SOC)	I-iı	ndex	(mg C	ired-C CO <sub>2</sub> -C / OC)		-C (% OC)		C (%) k soil)
			(161;		(161;		(400;		(382;		(50;		(0.00;		(4.4;		(12.1;		(1.2;
0–10 cm	53	276	443;	225	288;	421	439;	399	422;	59	68;	0.17	0.32;	17.0	33.7;	22.6	43.0;	5.1	15.1;
			77)		37)		9)		9)		4)		0.07)		7.2)		7.3)		2.7)
			(75;		(236;		(421;		(390;		(59;		(-0.18;		(3.6;		(2.5;		(0.2;
40–80 cm	46	133	202;	439	875;	448	480;	431	470;	69	79;	0.11	0.39;	13.4	32.2;	11.0	33.6;	0.9	3.9;
			34)		138)		10)		18)		5)		0.14)		6.9)		6.1)		0.8)

Table 2. Spearman correlation coefficients between 10-week mineralizable SOC (respired-C), the proportion of OC in the POM fraction (POM-C), the RE6 parameters and the C/N ratio, pH and clay content of the bulk soil, for both the 0–10 cm (n = 53) and 40–80 cm (n = 46) layers and each layer individually. Significance is indicated as follows: \*\*\*: p < 0.001; \*\*: p < 0.01; \*: p < 0.05. The very high correlations are marked in bold.

All (n = 99)	respired-C	POM-C	T <sub>50_HC_PYR</sub>	T <sub>50_CO2_OX</sub>	HI	$OI_{RE6}$	I-index	R-index	C/N	pН
POM-C	0.20									
$T_{50\_HC\_PYR}$	-0.26**	-0.73***								
$T_{\rm 50\_CO2\_OX}$	-0.16	-0.56***	0.76***							
HI	0.06	0.67***	-0.78***	-0.66***						
$OI_{RE6}$	-0.02	-0.76***	0.78***	0.63***	-0.92***					
I-index	0.32**	0.35***	-0.48***	-0.17	0.10	0.06				
R-index	-0.31**	-0.69***	0.93***	0.64***	-0.64***	0.67***	-0.74***			
C/N	-0.13	0.63***	-0.55***	-0.52***	0.67***	-0.78***	0.16	-0.50***		
pН	0.23*	-0.55***	0.49***	0.44***	-0.57***	0.66***	-0.27**	0.49***	-0.64***	
clay content	0.20	-0.19	0.04	-0.06	-0.17	0.31**	-0.22**	0.13	-0.49***	0.43***
0–10 cm	respired-C	POM-C	T <sub>50_HC_PYR</sub>	T <sub>50_CO2_OX</sub>	HI	$OI_{RE6}$	I-index	R-index	C/N	pН
POM-C	-0.29*									
$T_{50\_HC\_PYR}$	0.12	-0.44***								
$T_{\rm 50\_CO2\_OX}$	0.13	-0.19	0.45***							
HI	-0.43**	0.32*	-0.30*	0.07						
$OI_{RE6}$	0.52***	-0.41**	0.40**	-0.06	-0.91***					
I-index	0.11	0.37**	-0.87***	-0.43**	0.04	-0.17				
R-index	0.07	-0.44**	0.99***	0.44***	-0.24	-0.35*	-0.92***			
C/N	-0.51***	0.56***	-0.55***	-0.19	0.67***	-0.80***	0.37**	-0.52***		
pН	0.62***	-0.35**	0.46***	0.36**	-0.61***	0.70***	-0.25	0.42**	-0.70***	
clay content	0.43**	-0.29*	0.44**	-0.03	-0.71***	0.75***	-0.31*	0.43**	-0.70***	0.60***

40–80 cm	respired-C	POM-C	T <sub>50_HC_PYR</sub>	T <sub>50_CO2_OX</sub>	HI	$OI_{RE6}$	I-index	R-index	C/N	pН
POM-C	0.47***									
$T_{50\_HC\_PYR}$	-0.41**	-0.35*								
$T_{50\_CO2\_OX}$	-0.01	-0.01	0.26							
HI	-0.03	0.06	-0.24	-0.24						
$\mathrm{OI}_{\mathrm{RE6}}$	0.13	-0.42**	0.10	0.22	-0.47***					
I-index	0.41**	0.13	-0.19	0.43**	-0.49***	0.40**				
R-index	-0.52***	-0.28	0.64***	-0.19	0.20	-0.20	-0.85***			
C/N	-0.17	0.30*	0.10	-0.18	0.25	-0.65***	0.30*	0.23		
pН	0.03	-0.34*	-0.03	-0.18	0.11	0.25	-0.07	0.06	-0.27	
clay content	-0.08	-0.42**	-0.01	0.11	-0.07	0.62***	-0.15	0.17	-0.59***	0.35*

Figure 1 Click here to download high resolution image

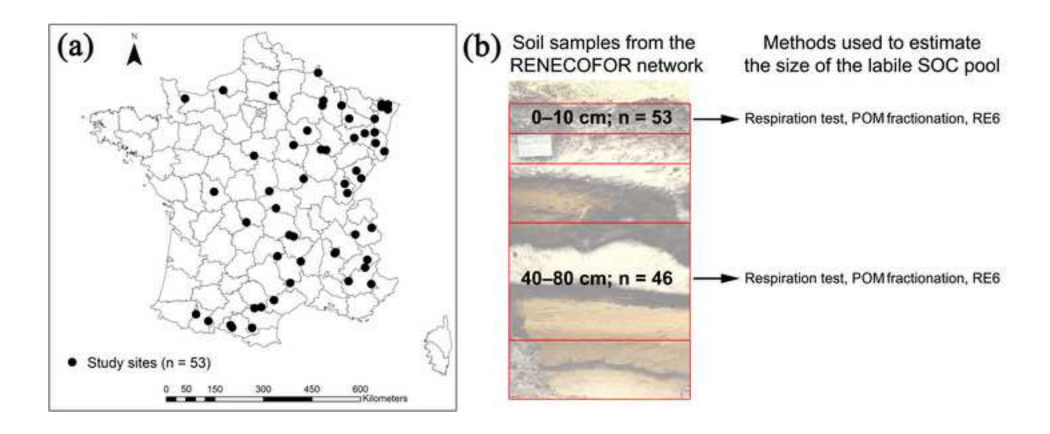


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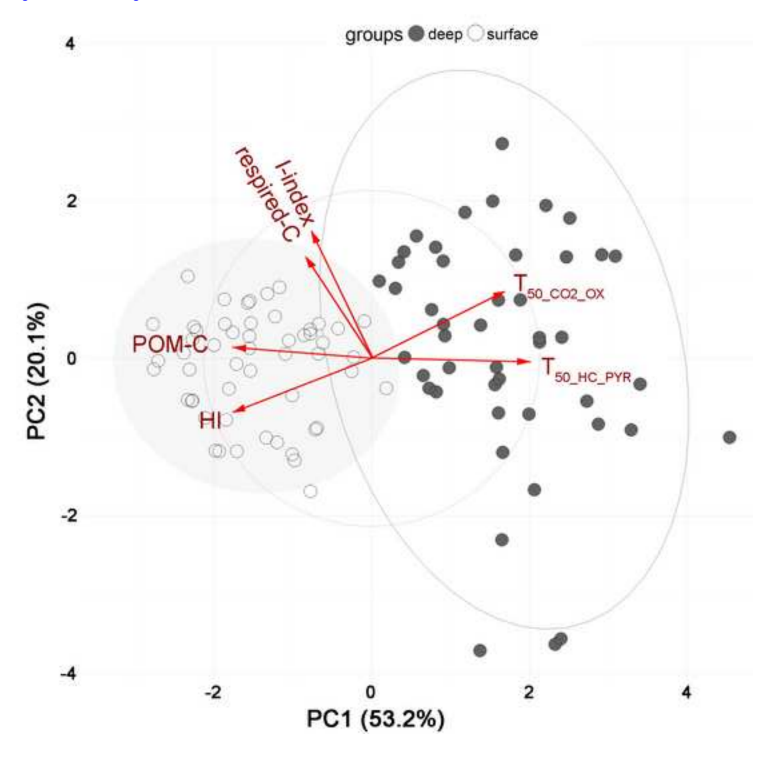
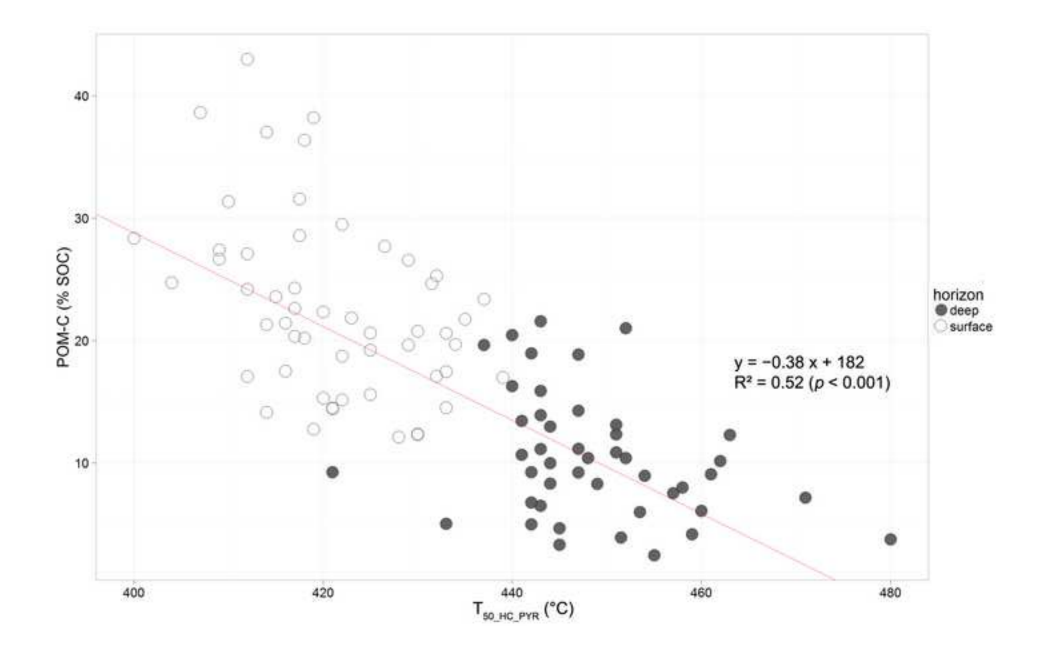


Figure 3
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Is Rock-Eval 6 thermal analysis a good indicator of soil organic carbon lability? – A method-comparison study in forest soils

Laure Soucémarianadin<sup>1,\*</sup>, Lauric Cécillon<sup>2</sup>, Claire Chenu<sup>3</sup>, François Baudin<sup>4</sup>, Manuel Nicolas<sup>5</sup>, Cyril Girardin<sup>3</sup> and Pierre Barré<sup>1</sup>

# **Supporting Information**

**Table SI-A.1**. Mean (+ standard deviation) particle-size distribution, pH and C/N ratio of the studied samples.

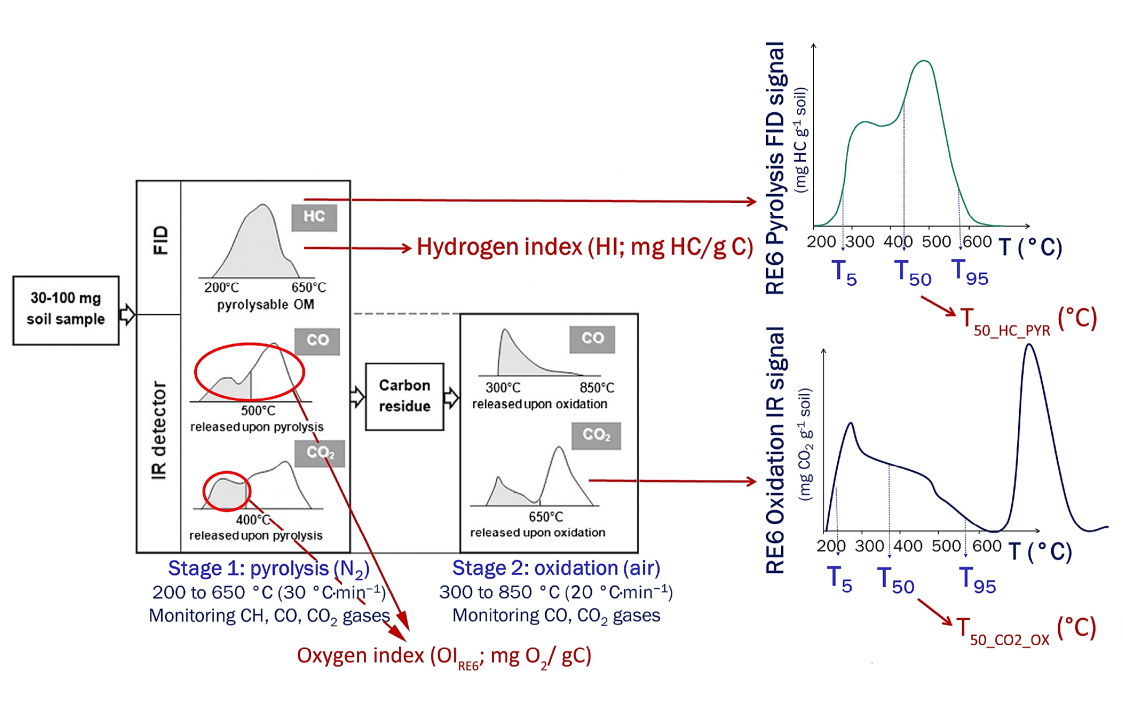
**Table SI-B1**. Percentage of variance explained and loadings of the first three principal components (PC) after Box-Cox transformation to correct for skewness for the PCA of all (0–10 cm and 40–80 cm) samples (n = 99). Values in bold indicate the variables with loading greater than the mean of the absolute loading in each PC.

**Table SI-C1**. Spearman correlation coefficients between 10-week mineralizable SOC (respired-C), the proportion of OC in the POM fraction (POM-C), the RE6 parameters and the C/N ratio of the bulk soil, for the three soil types and the two vegetation types. Significance is indicated as follows: \*\*\*: p < 0.001; \*\*: p < 0.01; \*: p < 0.05. The very high (> 0.9) correlations are marked in bold.

**Fig. SI-A1**. Description of the Rock-Eval 6 thermal analysis (adapted from Saenger et al., 2013) and calculation of four RE6-derived parameters (Hydrogen index; Oxygen index; T<sub>50 HC PYR</sub>, the temperature at which 50% of the HC resulting from the SOM pyrolysis had

evolved; T<sub>50\_CO2\_OX</sub>, the temperature at which 50% of the residual SOM was oxidized to CO<sub>2</sub> during the oxidation phase).

**Fig. SI-B1**. The proportion of OC in the POM fraction (POM-C) as a function of (a) respired-C (the proportion of total SOC mineralizable during a 10-week laboratory incubation); (b)  $OI_{RE6}$  (the oxygen index); (c) HI (the hydrogen index); (d) R-index (the proportion of thermally stable SOC pool) for all samples (n = 99; surface = 0–10 cm and deep = 40–80 cm).



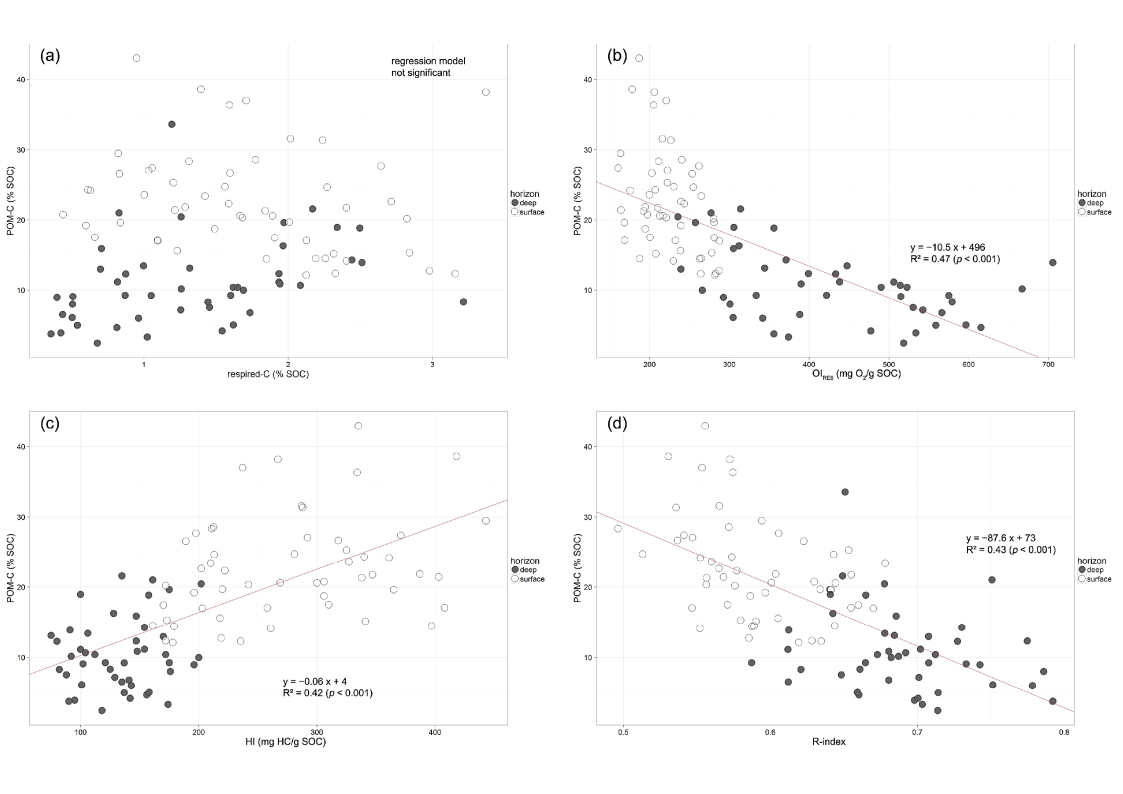


Table SI-A.1. Mean (+ standard deviation) particle-size distribution, pH and C/N ratio of the studied samples in each layer of the 53 plots.

depth (cm)	n	clay (%)	silt (%)	sand (%)	$pH_{\text{water}}$	C/N bulk soil	
0–10	53	22.5 (13.6)	35.5 (18.0)	42.0 (28.8)	4.9 (1.0)	16.9 (4.5)	
40-80	46	21.0 (15.4)	32.8 (16.2)	46.2 (26.7)	5.9 (1.5)	11.8 (3.8)	

Table SI-B1. Percentage of variance explained and loadings of the first three principal components (PC) after Box-Cox transformation to correct for skewness for the PCA of all samples (n = 99). Values in bold indicate the variables with loading greater than the mean of the absolute loading in each PC.

PC	PC1	PC2	PC3
% variance explained	53.2	20.1	13.1
respired-C	-0.22	0.55	0.77
POM-C	-0.47	0.06	0.00
$T_{50\_HC\_PYR}$	0.53	-0.02	0.11
$T_{50\_CO2\_OX}$	0.45	0.36	-0.18
HI	-0.46	-0.29	-0.15
I-index	-0.20	0.69	-0.58

Table SI-C1. Spearman correlation coefficients between 10-week mineralizable SOC (respired-C), the proportion of OC in the POM fraction (POM-C), the RE6 parameters and the C/N ratio of the bulk soil, for the three soil types and the two vegetation types. Significance is indicated as follows: \*\*\*: p < 0.001; \*\*: p < 0.01; \*\*: p < 0.05. The very high (> 0.9) correlations are marked in bold.

SOIL TYPE											
		respired-C	POM-C	$T_{50\_HC\_PYR}$	$T_{50\_CO2\_OX}$	HI	I-index	$OI_{RE6}$	R-index	C/N	pН
	POM-C	0.15									
	$T_{50\_HC\_PYR}$	-0.36*	-0.68***								
ol	$T_{\rm 50\_CO2\_OX}$	-0.18	-0.67***	0.77***							
bis	HI	0.17	0.62***	-0.81***	-0.65***						
Jan	I-index	0.47**	-0.01	-0.20	0.05	-0.17					
dystric Cambisol	$OI_{RE6}$	-0.15	-0.73***	0.82***	0.73***	-0.89***	0.15				
/str	R-index	-0.50**	-0.52**	0.90***	0.61***	-0.62***	-0.58***	0.63***			
Ð,	C/N	0.11	0.68***	-0.67***	-0.66***	0.58***	0.14	-0.75***	-0.64***		
	рH	0.14	-0.60***	0.58***	0.51**	-0.61***	0.08	0.60***	0.42*	-0.38*	
	clay content	-0.30	-0.16	0.15	-0.02	-0.03	-0.45**	0.23	0.32	-0.27	-0.02
	POM-C	0.46*									
	$T_{50\_HC\_PYR}$	-0.45*	-0.68***								
eutric Cambisol	$T_{50\_CO2\_OX}$	-0.54**	-0.53**	0.77***							
	HI	0.31	0.72***	-0.52**	-0.48**						
am	I-index	0.39*	0.37*	-0.58***	-0.36	0.21					
၁	$OI_{RE6}$	-0.42*	-0.88***	0.68***	0.55**	-0.83***	-0.39*				
utri	R-index	-0.50**	-0.68***	0.94***	0.72***	-0.55**	-0.78***	0.70***			
O	C/N	0.41*	0.84***	-0.60***	-0.49**	0.71***	0.17	-0.82***	-0.54**		
	pН	-0.57***	-0.56**	0.69***	0.71***	-0.54**	-0.20	0.62***	0.62***	-0.56**	
	clay content	-0.03	-0.13	-0.08	-0.08	-0.03	-0.14	-0.05	0.01	0.01	-0.25
	POM-C	0.42*									
	$T_{50\_HC\_PYR}$	-0.35*	-0.86***								
		-0.24	-0.52**	0.69***							
zol	HI	0.18	0.71***	-0.75***	-0.57***						
od	I-index	0.31	0.55***	-0.69***	-0.30	0.27					
entic Podzol	$OI_{RE6}$	-0.15	-0.71***	0.75***	0.48**	-0.96***	-0.26				
ent		-0.32	-0.83***	0.97***	0.62***	-0.68***	-0.81***	0.68***			
	C/N	0.08	0.54**	-0.55***	-0.44*	0.74***	0.13	-0.70***	-0.46**		
	pН	-0.29	-0.69***	0.83***	0.72***	-0.71***	-0.48**	0.72***	0.79***	-0.55***	
	clay content		-0.10	-0.06	-0.29	-0.15	0.05	0.23	-0.06	-0.39*	0.02

VEGETATION TYPE

		respired-C	POM-C	$T_{50\_HC\_PYR}$	$T_{50\_CO2\_OX}$	HI	I-index	$OI_{RE6}$	R-index	C/N	pН
	POM-C	0.20									
	$T_{50\_HC\_PYR}$	-0.25	-0.67***								
	$T_{50\_CO2\_OX}$	-0.24	-0.52***	0.72***							
sn	HI	0.05	0.70***	-0.82***	-0.67***						
èro	I-index	0.29*	0.38**	-0.52***	-0.16	0.21					
coniferous	$OI_{RE6}$	0.00	-0.75***	0.76***	0.58***	-0.92***	-0.27				
3	R-index	-0.25	-0.68***	0.95***	0.63***	-0.75***	-0.74***	0.74***			
	C/N	-0.13	0.60***	-0.46***	-0.33*	0.73***	0.19	-0.80***	-0.47***		
	pН	0.37**	-0.54***	0.42**	0.37**	-0.64***	-0.18	0.70***	0.45***	-0.66***	
	clay content	0.27*	-0.20	-0.11	-0.32*	-0.15	-0.09	0.33*	-0.01	-0.51***	0.50***
	POM-C	0.21									
	$T_{50\_HC\_PYR}$	-0.29	-0.80***								
	$T_{50\_CO2\_OX}$	-0.07	-0.56***	0.70***							
ns	HI	0.10	0.69***	-0.83***	-0.73***						
deciduous	I-index	0.40**	0.17	-0.24	0.15	-0.04					
eci	$OI_{RE6}$	-0.05	-0.77***	0.84***	0.69***	-0.92***	0.03				
ð	R-index	-0.44**	-0.73***	0.92***	0.52***	-0.66***	-0.57***	0.67***			
	C/N	-0.16	0.67***	-0.66***	-0.66***	0.81***	-0.20	-0.86***	-0.46**		
	pН	0.07	-0.55***	0.52***	0.42**	-0.53***	-0.19	0.60***	0.52***	-0.61***	
	clay content	0.09	-0.12	0.16	0.15	-0.26	-0.25	0.27	0.24	-0.34*	0.35*