

Evaluation of optical techniques for characterising soil organic matter quality in agricultural soils

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1 **EVALUATION OF OPTICAL TECHNIQUES FOR CHARACTERISING SOIL**
2 **ORGANIC MATTER QUALITY IN AGRICULTURAL SOILS**

3

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24 Soil organic matter (SOM) is one of the main global carbon pools. It is a measure of soil
25 quality as its presence increases carbon sequestration and improves physical and
26 chemical soil properties. The determination and characterisation of humic substances
27 gives essential information of the maturity and stresses of soils as well as of their health.
28 However, the determination of the exact nature and molecular structure of these
29 substances has been proven difficult. Several complex techniques exist to characterise
30 SOM and mineralisation and humification processes. One of the more widely accepted
31 for its accuracy is Nuclear Magnetic Resonance (NMR) spectroscopy. Despite its
32 efficacy, NMR needs significant economic resources, equipment, material and time.
33 Proxy measures like the fluorescence index (FI), cold and hot-water extractable carbon
34 (CWC and HWC) and SUVA₋₂₅₄ have the potential to characterise SOM and, in
35 combination, provide qualitative and quantitative data of SOM and its processes.
36 Spanish and British agricultural cambisols were used to measure SOM quality and
37 determine whether similarities were found between optical techniques and ¹H-NMR
38 results in these two regions with contrasting climatic conditions. High correlations
39 ($p < 0.001$) were found between the specific aromatic fraction measured with ¹H-NMR
40 and SUVA₋₂₅₄ ($R_s = 0.95$) and HWC ($R_s = 0.90$), which could be described using a linear
41 model. A high correlation between FI and the aromatics fraction measured with ¹H-

Abbreviations: (SOM) Soil organic matter; (NMR) Nuclear magnetic resonance; (FI) Fluorescence index; (CWC) Cold-water extractable carbon; (HWC) Hot-water extractable carbon; (LOI) Loss of ignition; (DOM) Dissolved organic matter; (DOC) Dissolved organic carbon; (EEM) Excitation-emission matrix; (GS-UK) Soil with a grass cover, United Kingdom; (CC-UK) Cereal crops from United Kingdom; (CC-ES) Cereal crops from Spain; (OG-ES) Olive grove from Spain; (HIX) Humification index; (BIX) Biological/autochthonous index; (RU) Raman units; (SUVA-254) Specific absorbance at 254 nm

42 NMR ($R_s=-0.976$) was also observed. In view of our results, optical measures have a
43 potential, in combination, to predict the aromatic fraction of SOM without the need of
44 expensive and time consuming techniques.

45 **KEYWORDS:** SOM quality; hot-water extractable carbon; cold-water extractable
46 carbon; fluorescence index; EEM; SUVA₋₂₅₄; ¹H-NMR; aromatic fraction.

47 **1. Introduction:**

48 *1.1. The importance of organic matter*

49 Soil organic matter (SOM) is composed of organic residues that are originated from
50 plant and animal remains and microbial products at different stages of decomposition or
51 humification (Hur *et al.*, 2013). Additionally, it is one of the main global carbon pools,
52 storing three times more carbon than living organisms or the atmosphere (Fischlin *et al.*,
53 2007; Brevik, 2012). Aside from carbon sequestration, SOM is also a measure of soil
54 quality because of the beneficial function it has on a variety of soil processes. For
55 instance, it reduces erosion and, therefore, increases crop production by increasing the
56 elasticity and resistance to deformation and compactability as well as porosity and water
57 retention (Sellami *et al.*, 2008; Paradelo and Barral, 2013). Increased water retention
58 decreases potential runoffs by improving water infiltration in to soils and provides a
59 store of water for plant uptake, buffering against moisture and rainfall fluctuations (Lal,
60 2004). This is of importance considering that the lack of water retention leads to a
61 change in the hydrological patterns of agricultural areas and promotes the quantity and
62 severity of floods and water-led erosion. Also, SOM leads to an increased vegetative
63 cover, which ultimately reduces soil erosion (Cerdà, 1998, 2000; Novara *et al.*, 2011;
64 Zhao *et al.*, 2013). Carbon mineralisation is crucial in SOM dynamics and along with
65 carbon input, determines how much carbon accumulates in soil and releases nutrients
66 that are essential for plant growth. Factors that affect mineralisation are the size of

67 labile carbon, environmental conditions and the local microbial community (Zhao *et al.*,
68 2008; Li *et al.*, 2013). SOM and soil assemblage; SOM decomposition and transport by
69 organisms contribute to soil stabilisation and the improvement of soil structure (Brevik
70 *et al.*, 2015). Moreover, SOM quantity has been directly related to the preservation of
71 soil aggregates, which in turn reduces soil erodibility (Novara *et al.*, 2011). Also, the
72 direct processing of SOM along with its decomposition contribute to the improvement
73 of soil chemical properties and stability (Brevik *et al.*, 2015). Therefore, optimal
74 quantities of SOM improve structure, water retention, and nutrient holding capacity of
75 soils, which has an effect in multiple aspects of the soil system. These are essential
76 considering the wider context of Earth System, as SOM conservation techniques have
77 been proven to improve the fertility of degraded soils of a wide variety of ecosystems
78 that are the main resource of large communities of developing areas of our planet, as
79 well as have an influence on biogeochemical cycles and climate change mitigation
80 (Batjes, 2014; Saha *et al.*, 2014; Srinivasarao *et al.*, 2014).

81 *1.2. SOM carbon fractions and their importance*

82 SOM has been conceptualised as containing three pools, with different
83 residence/turnover times (Trumbore, 2000). These pools are the active SOM (living
84 biomass of microorganisms and partially decomposed residues; associated with 1 year
85 turnover); the slow SOM (resistant plant material; associated to a turnover from years to
86 centuries); and passive SOM (humic substances and inert organic matter), which has
87 been traditionally associated with longer residence time (thousands of years) and more
88 stability. Some authors consider that the inert organic matter should not be considered
89 as part of the passive pool, but as a fourth pool (Trumbore, 1997; Ohno, 2002; Agren
90 and Bosatta, 2002; Sparks, 2003; Bell and Lawrence, 2009; Dungait *et al.*, 2012).
91 Although the traditional view has been that decomposition led to complex molecules

92 that were very stable as a result of their structure; it has recently been known that
93 environmental conditions, organo-mineral associations and other processes influence
94 more in SOM stability than structure, which only plays a secondary role. As a result of
95 this new view, recent research has found that humic substances, which have always
96 been considered high molecular mass polymers, could be simpler than originally
97 thought (Kleber and Johnson, 2010; Schmidt *et al.*, 2011). Still, their structure is on
98 discussion and the separation of SOM into fractions with different turnovers remains a
99 major challenge (Kleber, 2010; Schmidt *et al.*, 2011; Schrumpf and Kaiser, 2015).
100 Non-humic substances are composed by microbial biomass, decomposable plant
101 material (active SOM); and resistant plant material, mainly waxes, lignified tissues and
102 polyphenols (slow SOM) (Dungait *et al.*, 2012). Microbial biomass has been used for
103 comparing natural and degraded ecosystems and as an early indicator of soil processes,
104 fertility and health (García-Gil *et al.*, 2000; Brevik, 2009; Chen *et al.*, 2013).

105 *1.3. Current SOM quality measurements. Challenges*

106 Measures such as % Organic Matter measured by Loss of Ignition (LOI) are useful,
107 popular and inexpensive methods to determine bulk SOM (Luke *et al.*, 2009; Salehi *et*
108 *al.*, 2011). However, some studies have concluded that bulk SOM measurements cannot
109 be used, on their own, as a representative indicator of carbon in soil due to their
110 limitations (Koarashi *et al.*, 2005; Salehi *et al.*, 2011).

111 Humic substances have also been measured to determine soil quality, as their presence
112 has been associated with a higher quality of soils as stated in section 1.2. Their study is
113 relevant in agricultural soils, as they increase crop yield and root dry weight, although
114 this response is not fully understood (Rose *et al.*, 2014). Therefore, the determination
115 and characterisation of humic substances gives essential information of the maturity and
116 stresses of soils as well as of their health.

117 Traditionally, alkali and acid abstraction methods have been used, to later interpret the
118 chemistry of the extracted functional groups (Olk and Gregorich, 2006). Afterwards,
119 these were combined with other complex techniques that enabled scientists to obtain
120 new information on the structure and dynamic associations of humic substances (Sutton
121 and Sposito, 2005; Schmidt *et al.*, 2011). Despite these advances, SOM dynamics and
122 cycling still have many questions to answer, with models differing in SOM fluxes
123 results for the future, due to their sensitivity to SOM turnover time assumptions
124 (Schmidt *et al.*, 2011). There are a number of powerful but complex and expensive
125 techniques that have been used for the study of soil fluxes (Helal *et al.*, 2011). The
126 economic resources needed, along with the time required to prepare the samples and
127 conduct the analyses, make its use with a large number of samples difficult and delays
128 experiments, while more work is still needed to accurately determine and define the
129 molecular structures and linkages between the SOM components (Weishar *et al.* 2003;
130 Helal *et al.*, 2011).

131 Nuclear Magnetic Resonance (NMR) spectroscopy is a non-destructive technique that is
132 valuable for the characterisation of SOM and humification processes, providing
133 information on static and dynamic properties of molecules. This is due to its high
134 performance to assess intermolecular interactions. The relationship between SOM,
135 contaminants and metals can also be studied with NMR (Cardoza *et al.*, 2004). Of the
136 various variants that exist, ¹H-NMR spectroscopy was used in this study. This technique
137 analyses humic and fulvic acids dissolved in neutral or alkaline solutions to characterise
138 the components of the substance, and gives a semi-quantitative notion of aromatic,
139 aliphatic and carboxylic groups (Hemminga and Buurman, 1997). One of the main
140 drawbacks of this technique is the quantity of economic resources that are necessary for
141 its regular application in research laboratories/centres. This is due to the expensive

142 deuterated solvents and NMR tubes, as well as the expensive equipment and significant
143 sample preparation that are required (Weishar *et al.*, 2003; Cardoza *et al.*, 2004; MIT,
144 2008). Also, the technique is time consuming not only when measuring, but when
145 interpreting 2-D or 3-D data resulting from it (Cardoza *et al.*, 2004). Simpler methods
146 for the characterisation of SOM are required.

147 *1.4. Proxy measures. Opportunities to improve the ability to characterise SOM*
148 *quality*

149 Water extractable carbon is the most active component in the carbon cycle. Its quantity
150 and biological nature is affected by the extraction temperature (Bu *et al.*, 2010). Hot-
151 water extractable carbon (HWC) contains simple compounds such as microorganisms,
152 soluble carbohydrates and other compounds that account for the labile fraction of SOM
153 (Ghani *et al.*, 2003). HWC responds to land use changes in the short term and has been
154 used to detect the effects of different land management practices and for determining
155 the effects of soil amendments such as biochar or agricultural residues (Leifeld and
156 Kögel-Knabner, 2005; Uchida *et al.*, 2012; Alburquerque *et al.*, 2014; Fernández-
157 Romero *et al.*, 2014). For these reasons, it has been proven useful to obtain information
158 about soil quality (Ghani *et al.*, 2003; Xue *et al.*, 2013).

159 Fluorescence has become popular because of its potential to characterise SOM and
160 study humic substances, as it is non-destructive, simple, non-separative and accurate. As
161 a result, it has been used for determining the compositional and structural properties of
162 SOM (Chen *et al.*, 2003; Senesi and D'Orazio, 2005; Sun *et al.*, 2007; Kwiatkowska *et*
163 *al.*, 2008; Henderson *et al.*, 2009; Hur and Kim, 2009; Tang *et al.*, 2011). The intensity
164 and position of the peaks detected in the spectra are unique to each substance structural
165 and functional characteristics. For instance, higher fluorescence intensities are related
166 to a higher humification (Martins *et al.*, 2011).

167 The Fluorescence Index (FI) was developed to assess different properties of dissolved
168 organic matter (DOM). It was defined by McKnight *et al.* (2001) as the ratio of
169 emission intensities at 450-500 nm excited at 370 nm. The 450 nm point was chosen for
170 specific characteristics of the experiment. Later, Cory *et al.* (2010) modified the ratio to
171 470-520 nm to reflect corrections specific to the instruments used. This index has been
172 correlated to the aromaticity of DOM (Korak *et al.*, 2014).

173 Fluorescence spectroscopy can be used in combination with UV-Visible spectroscopy to
174 characterise humic substances, as absorbance measures transitions from the ground state
175 to the excited state, as opposed to fluorescence spectroscopy (Skoog *et al.*, 2007). Its
176 spectra are usually uniform and provide with qualitative data when a specific
177 wavelength is selected (Hassouna, *et al.*, 2012). Also the specific absorbance at 254 nm
178 (SUVA₋₂₅₄) has been recognised as a method to determine SOM aromaticity (Fuentes *et*
179 *al.*, 2006; Chow 2006). This parameter is very useful for assessing the nature of the
180 general composition of dissolved organic carbon (DOC), due to its high correlation with
181 it (Weishar *et al.*, 2003).

182 Considering what has been described in sections 1.3 and 1.4 above, both the
183 fluorescence and NMR techniques can be used in combination to determine the humic
184 substances properties and the degree of aromaticity; while HWC could contribute
185 further to the understanding of soil quality, given its usefulness to detect the
186 biodegradation of soil biochemical properties (Ghani *et al.*, 2003; Saab and Martin-
187 Neto, 2007; González-Pérez *et al.*, 2007).

188 As an illustrative and additional way to characterise and represent some of the analyses
189 conducted and the results obtained, Excitation-Emission Matrix (EEM) spectra have
190 been plotted. These provide information on the relative intensity of fluorescence at
191 different excitation and emission wavelengths regions in a fast manner that is also easy

192 to interpret (Coble, 1996). Several peaks have been identified that are used to describe
193 EEM fluorescence spectra. Peak A and C refer to humic peaks. Peak A is referred to as
194 UVC-excited and is located at an excitation wavelength (λ_{Ex}) between 240-260 nm and
195 an emission wavelength (λ_{Em}) between 400-460 nm. Peak C, also referred to as UVA-
196 excited; is located at a λ_{Ex} between 320-360 nm and a λ_{Em} between 420-460 nm. There
197 are also peaks that indicate biological activity material (peaks B and T, which are
198 defined as tyrosine-like and tryptophan-like peaks respectively). B has λ_{Ex} of 270-280
199 nm and λ_{Em} of 300-315 nm whereas T has λ_{Ex} between 270-280 nm and λ_{Em} of 345-
200 360 nm (Birdwell and Engel, 2010).

201 *1.5. Aim/objective of this study*

202 The aim of this study was to evaluate the use of fluorescence spectroscopy to measure
203 SOM quality (specifically the grade of humification). The specific objectives were: (1)
204 characterise water extractable SOM quality using liquid state $^1\text{H-NMR}$; (2) characterise
205 the quality of water extractable organic matter using fluorescence spectroscopy and UV-
206 VIS; (3) compare measures of quantity and quality of water extractable organic matter
207 with the specific organic matter fractions measured by $^1\text{H-NMR}$ like aromaticity. If
208 robust relationships and similarities between optical measures and $^1\text{H-NMR}$ are found,
209 there may be potential for fluorescence spectroscopy as a fast and more cost-effective
210 method of organic matter characterisation.

211 **2. Materials and methods**

212 *2.1. Field sites description*

213 Cambisols were sampled in two regions with contrasting climatic conditions;
214 Andalusia (South Spain) and Berkshire (South East England) (Table 1). Berkshire has a
215 temperate climate, characterised generally by relatively mild winters and summers and
216 rainfall throughout the year. The annual mean temperature ranges from 6.7 °C and

217 14.5 °C (30 years, annual mean temperature: 10.5 °C) and the average annual rainfall is
218 635.4 mm (UK Met Office, 2014).

219 Andalusia has a Mediterranean climate, which characteristics are hot and dry summers
220 contrasted with cool and wet winters. In the province of Cordoba (Hinojosa del Duque
221 and Pozoblanco), the annual mean temperature is 17.6 °C and the average annual
222 rainfall is 536 mm (Aemet, 2014). As for the province of Jaen (Torredelcampo), its
223 annual mean temperature is 16.2 °C and the average annual rainfall is 646.3 mm
224 (REDIAM, 2007).

225 In Berkshire, samples were collected from University of Reading Farms at Sonning,
226 Arborfield and Shinfield. The site sampled at Sonning (GS-UK-1) has been covered
227 with grass for over 15 years. Arborfield (CC-UK-1) was permanent pasture until
228 autumn 2012, and has been subsequently drilled with wheat or winter barley. Shinfield
229 (CC-UK-2) has been in an arable rotation for over 20 years and sown with either winter
230 wheat, maize or spring barley.

231 In Andalusia, soil samples were collected between the provinces of Jaen
232 (Torredelcampo) and Cordoba (Hinojosa del Duque, Pozoblanco) that were managed
233 with conventional tillage of cereals and olive crops. CC-ES-3 was managed with a
234 wheat-barley-fallow cycle, whereas the other sites (CC-ES-1 and CC-ES-2) were
235 covered by wheat crops throughout the year. OG-ES-1, OG-ES-2, and OG-ES-3 were
236 covered by olive grove.

237 2.2. *Sample collection and preparation*

238 Soil samples were collected from each horizon. Total soil depths are included in Table
239 2. Only the first horizon from each soil (specified in section 2.3) was used for the
240 analyses. Samples were air dried and sieved with a 2 mm sieve.

241

242 2.3. *Analytical methods*

243 Cold and hot water extractable carbon were determined following Ghani *et al.* (2003).
244 This consists in a cold and a hot extraction of the supernatant of the samples (room
245 temperature and 80 °C respectively). To do this, 30 ml of ultrapure water was added to
246 3 g of soil. Then, there was a 30-minute extraction in a shaker at 20°C. After this, the
247 sample was centrifuged during 20 minutes at 3500 rpm. Once this was done, the
248 supernatant was extracted (cold extraction) and analysed (DOC). The resulting pellet
249 was used for the rest of the steps. 30 ml of ultrapure water were added to the pellet,
250 which was then shaken on a Vortex to mix the ultrapure water with the pellet. The
251 sample was then left in a bath at 80 °C during 16 hours. The sample was centrifuged at
252 3500 rpm during 20 minutes and was filtered with a 0.45 µm cellulose nitrate filter (hot
253 extraction). The filtered supernatant was analysed for DOC, fluorescence, absorbance
254 and NMR.

255 The techniques that were compared in this study were applied to the first horizon of
256 each soil sample, as this was considered sufficient for the purposes of comparing these
257 techniques. This horizon was at different depths depending on the soil sample (at 12 cm
258 in GS-UK-1; 10.5 cm in CC-UK-1; 5.7 cm in CC-UK-2; 15 cm in CC-ES-1; 20 cm in
259 CC-ES-2; 30 cm in CC-ES-3; 10 cm in OG-ES-1; 10 cm in OG-ES-2 and 18 cm in OG-
260 ES-3).

261 Fluorescence of all the hot water extracts was measured in a Varian Eclipse
262 Fluorescence spectrophotometer (Agilent Technologies, Santa Clara, CA, USA) at an
263 emission wavelength from 300 to 600 nm at 5-nm increments and an excitation of 240-
264 450 nm at 5-nm increments. All samples were run in 1 cm quartz cuvettes and in
265 triplicate.

266 FI was calculated as the ratio of intensities at 450 over 500 nm with an excitation of
267 370 nm, as described by Cory *et al.* (2010). McKnight *et al.* (2001) introduced this
268 index approach for the characterization of the fulvic and fraction of DOM.

269 The Humification Index (HIX) was calculated with the following formula:

$$270 \quad (\sum I_{435 \rightarrow 480}) / (\sum I_{300 \rightarrow 345}) + (\sum I_{435 \rightarrow 480})$$

271 where I is the fluorescence intensity at each wavelength (modified of Zsolnay *et al.*,
272 1999 by Ohno, 2002).

273 The Biological/Autochthonous Index (BIX) has been calculated as the ratio of
274 intensities at 380 nm over 430 nm with an excitation of 310 nm, as described by Huguet
275 *et al.* (2009). This index assesses the relative contribution of autochthonous DOM in
276 water and soil samples.

277 A subsample of the HWC extract was frozen and subsequently freeze-dried to remove
278 all the water present. Deuterium oxide was added as a solvent to avoid disruptions in the
279 spectrum, as other solvents have a proton signal that causes disruptions, as
280 demonstrated by Cardoza *et al.* (2004), prior to a second freezing and a second freeze-
281 drying. This second stage of freezing-freeze-drying was used to avoid that H₂O peaks
282 interfered in the spectra. After that, deuterium oxide (100%, density of 1.107 g/ml at
283 25 °C) was added again as a solvent to conduct the NMR tests. NMR was measured in
284 1.5-1.7 ml of sample with a Bruker Avance III 700 MHz NMR spectrometer (Bruker
285 Corporation, Billerica, MA, USA). The deuterium oxide peak was used as a calibration
286 reference and placed in 4.75 ppm.

287 Absorbance of all the hot water extracts was measured with a Varian Cary 300 UV-
288 Visible spectrometer (Agilent Technologies, Santa Clara, CA, USA), from 200-800 nm
289 with 1 nm intervals using a 1 cm cuvette.

290 SUVA₋₂₅₄ has been used for this study. This is the absorbance at 254 nm divided by the
291 DOC concentration (of the hot water extracts). Values are expressed in $l \cdot mg^{-1} \cdot m^{-1}$.
292 DOC of the cold and hot water extracts was calculated with a Shimadzu TOC 5000 total
293 organic carbon analyser (Shimadzu Corporation, Kyoto, Japan). Standards were
294 calculated using Stock solutions of 1,000 ppm.

295 % Organic matter measured by LOI was calculated using a modified version of Hierl *et al.*
296 *al.* (2001). A soil sample of 10 g was heated at 105 °C for 24 hours. Then, the sample
297 was weighted again (w_1) and heated in a Muffle furnace at 550 °C for 16 hours to ignite
298 the organic matter. The sample was weighted after this (w_2) and the % of organic matter
299 was obtained by weight difference between w_1 and w_2 .

300 Soil pH was determined in 1:2.5 soil to water ratio. Texture was determined by laser
301 granulometry, using a Coulter LS 230 (Beckman/Coulter Inc. Brea, CA, USA). This
302 technique uses polarised light at three different wavelengths (450 nm, 600 nm and
303 900 nm) to analyse the particle size distribution specifically in the 0.04 μm to 0.4 μm
304 range.

305 EEM fluorescence spectra were obtained by collecting a series of emission scans of λ_{Ex}
306 240-450 nm at 5 nm intervals and λ_{Em} 300-600 nm, also at 5 nm intervals. EEM spectra
307 were plotted using RStudio v0.98.1091 (Rstudio Inc., Boston, MA, USA).

308 Intensities are reported in Raman units (RU). Raman scattering was mitigated
309 subtracting blanks that had been collected on ultrapure water from each spectrum.

310 2.4. *Statistical methods*

311 Data were tested for normality to verify the model assumptions. As the data failed the
312 normality test, non-parametric tests were used. The effect of the hot water extraction in
313 the extractability of carbon (compared to the cold water extraction) was analysed using
314 a Paired Sample Wilcoxon Signed Rank Test. The correlation between the different

315 analytical methods was tested using the Spearman Rank and assessing the significance
316 of the resulted Spearman Correlation Coefficient (Minitab 16 for Windows. Minitab
317 Inc., State College, PA, USA). Linear regressions of the Spearman Correlations were
318 also plotted (SigmaPlot 12.0 for Windows. Systat Software Inc., San Jose, CA, USA).
319 Differences of $p < 0.05$ were considered statistically significant.

320 **3. Results and discussion**

321 *3.1. Soil characteristics*

322 The majority of CC-UK soils had sandy texture, with a relatively high proportion of silt
323 (Table 2). The only exception was CC-UK-2, although the sand proportion was quite
324 close to that of silt. This texture was similar to that of CC-ES-1 and CC-ES-2, although
325 CC-ES-3 had a higher proportion of silt and higher proportion of clay than of sand. All
326 the OG soils presented a texture of a majority of silt (56.7-59.5%), followed by clay. It
327 is worth considering that silt is the most erodible fraction (Table 2).

328 pH in the UK soils was generally acid, although GS-UK-1 and CC-UK-1 had values
329 between 3.58-5.66 and CC-UK-2 had values close to 7. ES soils had generally higher
330 values, ranging from 5.53-6.98 (Table 2). Some authors have related pH and
331 measurement of aromatics (Weishar *et al.*, 2003). GS-UK-1 has the lowest pH (3.58)
332 and is the one with the highest aromatics content and highest HWC.

333 *3.2. Total extractable carbon by cold and hot extractions*

334 CWC and HWC measurement results are in Table 3. HWC extracted significantly more
335 carbon ($P < 0.01$) than CWC, thus proved to be a more exhaustive extraction method.
336 The increases in the values ranged from 161-605%. This was equivalent to higher
337 values by a factor between 3 and 7 respectively. The most significant increase was
338 obtained in GS-UK-1. Gregorich *et al.* (2003) obtained a similar trend in maize-cropped
339 soils of Ottawa, Canada, and found HWC exceeding CWC by a factor of two.

340 Moreover, Landgraf *et al.* (2006) found that HWC had higher carbon concentrations
341 than CWC by a factor that varied from 4 to 6 in surface horizons of forest soils in SE
342 Germany.
343 A correlation analysis was run to assess whether the C extractability differed between
344 the different soils and locations. The correlation resulted to be high ($R_s=0.91$, $p<0.01$).
345 Wang and Wang (2007) found a similar correlation ($r=0.93$, $p<0.01$) in forest oxisols
346 of Southern China. Klose and Makeschin (2003) also found that HWC and CWC
347 increased with the same proportion in forest soils of NE Germany. On the other hand,
348 Ghani *et al.* (2003) found a positive but poor correlation.
349 HWC and CWC data were compared with SOM (measured by LOI) to assess which of
350 the two extraction methods would imply a higher correlation with the organic matter
351 from the samples (Table 4). HWC implied a higher correlation with SOM measured
352 with LOI ($R_s= 0.70$, $p<0.05$) than CWC ($R_s= 0.55$). Despite the fact that the correlation
353 between CWC and SOM was lower than that of HWC; other authors have found even
354 poorer correlations. For instance, Van Migroet *et al.* (2005) found a r^2 of 0.2 in forest
355 soils in Utah, USA.

356 3.3. *Quality of cold-water extractable carbon using UV-Vis and Fluorescence*

357 Cold-water extractable carbon (CWC) data was compared with UV-Vis and
358 fluorescence spectroscopy, using $SUVA_{-254}$ and FI respectively. The Spearman Rank
359 Correlation was calculated to evaluate the quality and reproducibility of CWC using
360 optical techniques (Table 4).

361 There were high correlation patterns between CWC and $SUVA_{-254}$ ($R_s=0.82$, $p<0.05$)
362 although not between CWC and FI ($R_s=-0.29$), where the slight correlation that could
363 be appreciated was inverse. Chow (2006) obtained a worse correlation between CWC
364 and $SUVA_{-254}$ ($R^2=0.38$), whereas that of Van Migroet *et al.* (2005) was even lower

365 ($R^2=0.01$). As Weishar *et al.* (2003) point out; some authors have found conflicting
366 conclusions when using SUVA₋₂₅₄ to determine the aromaticity of DOC.

367 3.4. *Quality of hot-water extractable carbon using UV-Vis and Fluorescence*

368 HWC data was compared with SUVA₋₂₅₄ and FI to assess whether it had higher
369 correlation than CWC. As it can be seen in Table 4, there is a high correlation ($R_s=0.88$,
370 $p<0.001$) between HWC and SUVA₋₂₅₄ whereas that of HWC with FI is not as
371 significant ($R_s=-0.53$). In both cases, HWC was more significantly correlated to the
372 results obtained with fluorescence and SUVA₋₂₅₄ than CWC.

373 3.5. *EEM spectra general characteristics*

374 A set of reference fluorescence spectra has been represented in Figure 1. As it can be
375 seen in Figure 1.a and 1.b, the two soils with the highest % aromatics measured with
376 ¹H-NMR as well as high SUVA₋₂₅₄ and HWC values (GS-UK-1 and CC-UK-1) showed
377 strong peaks C (10-12 RUs), which indicate UVA-excited humic peaks as described by
378 Coble (1996). They also showed weak peaks A.

379 On the other hand, Figure 1.c. represents the soil with the highest FI, but lowest %
380 aromatic measured with ¹H-NMR as well as comparatively low values of SUVA₋₂₅₄
381 (CC-ES-1). The EEM spectrum of this soil is very different from the ones represented in
382 Figures 1.a and 1.b, with a strong peak A or UVC-excited (8 RUs), and weak peaks C
383 and T. The presence of the latter indicates biological activity.

384 3.6. *Relationship between optical measures and NMR*

385 The aromaticity of isolated fulvic acid samples was calculated as the ratio of the area of
386 aromatic hydrogen region to the total area of the ¹H-NMR spectrum (% aromaticity). An
387 analogous method was used to calculate the ratio of the area of aliphatic hydrogen and
388 carbohydrate hydrogen regions.

389 As it can be seen in Figure 2, the aromatic hydrogen region is situated in a chemical
390 shift of 6-8ppm by frequency. On the other hand, the carbohydrate hydrogen and
391 aliphatic hydrogen regions are situated in a chemical shift of 3-4.2 ppm and 0.5-3 ppm
392 respectively. For Figure 2, the two soils with the highest % aromatics were chosen for
393 representation purposes (GS-UK-1 and CC-UK-1).

394 Results of these measurements can be found in Table 3, along with the results from the
395 optical techniques. Correlation analyses were conducted comparing the results from all
396 of them (Table 4).

397 The largest carbon fraction of the UK soils was carbohydrates. The same trend was
398 found in the CC-ES soils, but not in the OG-ES soils, where the largest carbon fraction
399 was aliphatics.

400 The proportion of aromatics of the UK soils ranged from 3.91%-5.98%, whereas for the
401 CC-ES and OG-ES soils, it ranged from 2.40-3.47% and 3.90%-4.33% respectively. As
402 these were taken from samples in the first horizon of each soil, the low level of
403 aromaticity can be explained by the fact that plant residues may have accumulated in the
404 surface, while microorganisms did not have enough capacity to decompose them. As
405 Cardoza *et al.* (2004) point out, NMR is a powerful tool for investigating humic
406 substances interactions at the molecular level.

407 Aromatics have positive correlations for SUVA₋₂₅₄, CWC and HWC but are inversely
408 correlated to FI. Correlations are high for aromatics compared with SUVA₋₂₅₄ and
409 HWC (Figures 3 and 4. R_s of 0.95 and 0.90 respectively; $p < 0.001$), but not as good for
410 FI (Figure 5). The correlation between aromatics and CWC, although significant, is not
411 as strong as the one with HWC ($R_s = 0.74$, $p < 0.05$). Weishaar *et al.* (2003) found that
412 SUVA₋₂₅₄ is a good predictor of the humic fraction (higher SUVA₋₂₅₄ indicates higher
413 humic acid content and molecular weight in DOM solutions) and the general chemical

414 properties of HWC, although it does not predict the reactivity of HWC from different
415 types of source materials nor information of the individual molecules of samples unless
416 these are humic substances (Chin *et al.*, 1997; Weishar *et al.*, 2003). They obtained a
417 good correlation between SUVA₋₂₅₄ and aromaticity determined by ¹³C-NMR (R=0.97),
418 which was slightly higher than ours. Yeh *et al.* (2014) also found a high correlation
419 between SUVA₋₂₅₄ measurements and the aromatic fraction of samples of organic
420 matter of river sediments of Taiwan. Jamieson *et al.* (2014) found that higher SUVA₋₂₅₄
421 implied higher aromaticity in biochar from sugar maple, thus demonstrating that this
422 effect is currently studied when characterising organic material with a high recalcitrance
423 and that is able to retain carbon for thousands of years, due to its stability (Lehmann,
424 2007).

425 According to McKnight (2001) and other authors (Rodríguez *et al.*, 2014; Wei *et al.*,
426 2014), there is a correlation between the FI values and those from methods that indicate
427 the aromaticity of humic substances such as the ratio of the aromatic carbon region area
428 in the total NMR spectra and SUVA₋₂₅₄. This pattern was observed in our soils, with
429 negative correlations between FI and these 2 parameters (Rs=-0.63 and Rs=-0.52
430 respectively), although only the trend with the aromatic carbon region area in the total
431 NMR spectra was statistically significant (p<0.05).

432 Kim *et al.* (2006) and Rodríguez *et al.* (2014) indicated that differences in the FI higher
433 than 0.1 would imply significant differences in the aromaticity of the samples. If this
434 assumption is applied to our case, CC-UK-2 has differences in its aromaticity with GS-
435 UK-1 but not with CC-UK-1. CC-ES-2 has a different aromaticity than CC-ES-1 and
436 CC-ES-3. In the same way, OG-ES-1 has differences with OG-ES-2 and OG-ES-3.
437 When all the types are compared, there are combinations of differences in the
438 aromaticity but a single pattern cannot be obtained.

439 Given the results above, the regressions between the aromatic carbon region measured
440 with $^1\text{H-NMR}$ and SUVA_{-254} and HWC are powerful. The equations we obtained can
441 explain very significantly the relationship between these variables ($p < 0.001$ for the
442 results of $^1\text{H-NMR}$ predicted with SUVA_{-254} and $p < 0.01$ if they are predicted using
443 HWC). These equations (Figure 3 and Figure 4) could therefore be used to predict the
444 aromaticity of water-extractable carbon.

445 The regression of the aromaticity measured with $^1\text{H-NMR}$ with FI, although significant
446 ($p < 0.05$), did not have the same level of confidence as the one with HWC and SUVA_{-254} .
447 As it can be seen in Figure 5, an outlier was identified in the Regression analysis (FI
448 value of CC-ES-3). Figure 5 showed how the regression model improved when the
449 outlier was removed from the analysis ($R_s = -0.98$). The linear regression equation that
450 resulted after the elimination of the outlier is able to explain the $^1\text{H-NMR}$ values using
451 FI with a very high significance ($p < 0.001$).

452 However, according to McKnight (2001), the FI would not be sufficient to estimate
453 aromaticity and other techniques are necessary (e.g. SUVA_{-254}), as geological processes
454 can alter aromaticity without changing the FI.

455 SUVA_{-254} did not demonstrate a particularly strong correlation with FI ($R_s = -0.52$, Table
456 4). The correlation was negative (higher SUVA_{-254} led to lower FI) which coincides
457 with authors like Kothawala *et al.* (2012) with minerals soils collected across Canada
458 although their trend is less clear than in the study of other authors such as Williams *et*
459 *al.* (2010), where the slope is more pronounced. The correlation of Williams *et al.*
460 (2010) was similar to ours ($r = -0.57$ in their study and $R_s = -0.52$ in ours), although the
461 statistical analysis in their case was the Pearson correlation, as their data followed a
462 normal distribution. Hassouna *et al.* (2012) found a negative correlation as well, with
463 slightly higher correlation in Mediterranean calcareous soils ($r = -0.69$). As in the case

464 described in Figure 5, if the same outlier was removed, the correlation improved
465 significantly ($R_s=-0.98$) and the equation of the linear regression explained with a very
466 high degree of significance ($p<0.001$) the relation between SUVA₋₂₅₄ and FI.
467 Analogously, the same outlier could be removed in the analysis of HWC vs. FI,
468 obtaining a significant correlation between them ($R_s=-0.88$). A linear regression model
469 explains significantly ($p<0.01$) the relationship between HWC and FI and therefore
470 similar conclusions can be stated as in the case of the correlation of SUVA₋₂₅₄ to FI.
471 Given the strong intrinsic relationships between the results of SUVA₋₂₅₄, HWC and FI;
472 and how they are able to predict the proportion of aromaticity measured with ¹H-NMR,
473 we can conclude that UV-Vis absorbance and fluorescence spectroscopy can be used to
474 characterise the aromaticity of carbon and may be a plausible substitute for ¹H-NMR,
475 given that they are more cost-effective. Other authors such as Zornoza *et al.* (2008,
476 2015) demonstrated the effectiveness of spectroscopy (near infrared reflectance
477 spectroscopy in their study) as an accurate, cost- and time-effective method for
478 predicting and/or estimating soil biogeochemical properties and other soil parameters.
479 Correlations between aliphatics and the data from the optical measures and CWC/HWC
480 were poor (Table 4), although slight trends can be observed for SUVA₋₂₅₄, FI and CWC.
481 In these cases, the correlation was positive for FI and CWC but negative for SUVA₋₂₅₄.
482 Correlations were also poor between carbohydrates and the data from the optical
483 measures and HWC/CWC, with only one trend detected. This was the inverse
484 correlation ($R_s=-0.35$) between HWC and carbohydrates.
485 BIX and HIX did not correlate well with any of the other parameters (Table 4), with
486 very few trends detected. Of the trends detected, HIX was positively correlated with FI
487 ($R_s=0.52$). It is worth noting that BIX was positively correlated with carbohydrates but
488 inversely correlated to aliphatics ($R_s=0.31$ and -0.31 respectively), whereas the trend

489 was the contrary for HIX ($R_s=-0.28$ and 0.22). A similar trend than ours was also found
490 by Kalbitz *et al.* (2003) between HIX and carbohydrates in forest soils, arable soils and
491 a fen area; although theirs was stronger ($r^2=0.81$). Williams *et al.* (2010) studied the
492 correlation between HIX and SUVA₋₂₅₄ in watershed from mixed land use in Canada.
493 They obtained a positive correlation ($r=0.74$), whereas our correlation was negative. On
494 the other hand, Yeh *et al.* (2014) did not find a clear trend between SUVA₋₂₅₄ and HIX.
495 The significance of our comparison between SUVA₋₂₅₄ and HIX values is extremely
496 low and therefore robust conclusions on this issue cannot be stated. Bu *et al.* (2010)
497 compared HWC and HIX in various soils with different vegetation types in Wuyi
498 Mountain (SE China), obtaining a poor negative correlation, just like ours. However,
499 Kalbitz *et al.* (2003) found a strong correlation between the aromatic fraction of DOM
500 and HIX ($r^2=0.80$).
501 Yeh *et al.* (2014) calculated BIX as well, obtaining that higher SUVA₋₂₅₄ values implied
502 lower BIX values, which coincides with our comparison in this case, although its
503 significance is not relevant enough to establish a robust conclusion.
504 Plotting the results from Birdwell and Engel (2010) for FI, HIX, and BIX did not led to
505 significant correlations, which is similar to our results and, along with the authors
506 commented above, demonstrate that a number of studies have found difficulties when
507 correlating HIX and BIX to other parameters.

508 **4. Conclusions**

509 Different techniques to measure the quantity and quality of SOM were tested in
510 cambisols from very different climatic locations and under different cropping systems.
511 HWC extracted a higher amount and carbon than CWC, and correlated better with the
512 %SOM (LOI) than CWC.

513 SUVA₋₂₅₄ and HWC correlated significantly with the proportion of aromatics measured
514 with ¹H-NMR, demonstrating their complementary nature. Linear regression models
515 fitted to the data were able to explain the relationship between the specific fraction of
516 aromatics measured with ¹H-NMR and SUVA₋₂₅₄ and HWC, and therefore allow the
517 aromatic fraction to be estimated at lower cost using SUVA₋₂₅₄ and HWC.
518 A linear regression model was also able to explain the relationship between ¹H-NMR
519 measurements and FI, after an outlier was removed.
520 BIX and HIX were not useful indexes for our purposes, as they did not correlate well to
521 the carbon fractions measured with ¹H-NMR or with FI, SUVA₋₂₅₄, or HWC.
522 The EEM spectra showed a difference in the peak types that appeared in the CC-UK
523 soils and in the CC-ES soil. The fact that figures 1.a and 1.b showed a stronger peak C
524 and figure 1.c. showed a stronger peak A indicated subtle qualitative differences in their
525 humic fractions. This could have been caused by the different soil management (more
526 intense agricultural activity in CC-ES-1). More work will be needed to confirm this.
527 In view of our results, optical measures have a potential, in combination, to predict the
528 aromatic fraction of SOM without the need of expensive and time consuming
529 techniques like ¹H-NMR; which could be very useful when the equipment is not
530 available or in instances when a high number of samples need to be analysed
531 simultaneously.

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534 **6. References**

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791 **Figure captions**

792 **Figure 1.** EEM plots from: a) GS-UK-1; b) CC-UK-1; c) CC-ES-1

793 **Figure 2.** Liquid state ^1H -NMR spectra from: a) GS-UK-1; b) CC-UK-1

794 **Figure 3.** Correlation between % Aromatics and SUVA₋₂₅₄. A linear regression model
795 was calculated to explain the relationship between the variables. The equation was y [%
796 Aromatics] = $0.250 + 0.950 \times [\text{SUVA}_{-254}]$, $p < 0.001$

797 **Figure 4.** Correlation between % Aromatics and HWC. A linear regression model was
798 calculated to explain the relationship between the variables. The equation was y [%
799 Aromatics] = $0.500 + 0.900 \times [\text{HWC}]$, ($p < 0.01$)

800 **Figure 5.** Correlation between % Aromatics and FI. A linear regression model was
801 calculated to explain the relationship between the variables. The equation was y [%
802 Aromatics] = $8.17 - 0.633 \times$ [FI], ($p < 0.05$), represented by the solid line. The removal
803 of an outlier resulted in the equation: y [% Aromatics] = $8.89 - 0.976 \times$ [FI], which
804 explained the relationship with a higher significance ($p < 0.001$). The second equation is
805 represented by the dashed line