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1 **Combination of 3D Fluorescence/PARAFAC and UV-Vis Absorption for the characterization of** 2 **agricultural soils from Morocco**

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- 9

10 **Abstract**

11 The present study, combining UV-Visible absorption and 3D fluorescence supported by PARAFAC
12 chemometric analysis, focused on the characterization of soil water extractable organic matter (WEOM) in the 12 chemometric analysis, focused on the characterization of soil water extractable organic matter (WEOM) in the 13 zone of Doukkala located near the Atlantic coast of Morocco. The extracts, in water, of a set of 30 samples zone of Doukkala located near the Atlantic coast of Morocco. The extracts, in water, of a set of 30 samples 14 covering the four main types of agricultural soils in the region (commonly labeled Tirs, Faid, Hamri and R'mel) 15 were investigated. E_2/E_3 and E_4/E_6 absorbance ratios, $S_{275-295}$ and $S_{350-400}$ spectral slopes, along with their
16 ratios S_P. as well as the fluorescence FI and humification HIX indices were calculated an 16 ratios S_R , as well as the fluorescence FI and humification HIX indices were calculated and interpreted. In the four
17 soil types these parameters revealed, on the one hand, organic materials of terrigenous origin wi 17 soil types these parameters revealed, on the one hand, organic materials of terrigenous origin with a certain 18 biological component, and showed on the other hand that these materials are in similar stages of humificat 18 biological component, and showed on the other hand that these materials are in similar stages of humification
19 with an important humic character. The 3D fluorescence crossed with PARAFAC chemometrics highlighted the 19 with an important humic character. The 3D fluorescence crossed with PARAFAC chemometrics highlighted the
20 absence of any protein component and revealed the prevalence of the FAs fraction in the OM humic material in 20 absence of any protein component and revealed the prevalence of the FAs fraction in the OM humic material in all the soils investigated. all the soils investigated.

22 **Keywords** Morocco . Agricultural Soils . WEOM . UV-Vis. Absorption . Total fluorescence & PARAFAC.

23 **Introduction**

24 Water extractable organic matter (WEOM), like indigenous dissolved organic matter (DOM) passing a filter pore
25 size of 0.4 – 0.7 µm, is composed of an array of molecules generally reflecting the composition of total s 25 size of $0.4 - 0.7$ μ m, is composed of an array of molecules generally reflecting the composition of total soil
26 organic matter [1]. Although it generally comprises < 2% of total soil organic matter, its turnover r 26 organic matter [1]. Although it generally comprises $\langle 2\%$ of total soil organic matter, its turnover rate and 27 solubility mean that its role is determinant in many chemical and biological processes in the soil fate 27 solubility mean that its role is determinant in many chemical and biological processes in the soil fate [2].
28 Furthermore, while the frontier between DOM and WEOM is not always obvious [3], it is reported that WEOM 28 Furthermore, while the frontier between DOM and WEOM is not always obvious [3], it is reported that WEOM
29 may likely to contain further material in comparison with indigenous DOM, being thus of larger magnitude [1]. 29 may likely to contain further material in comparison with indigenous DOM, being thus of larger magnitude [1].
20 Therefore, the properties of WEOM are important indicators of soil quality, namely its physical-chemical a 30 Therefore, the properties of WEOM are important indicators of soil quality, namely its physical-chemical and microbial community characteristics [4]. microbial community characteristics [4].

- 32 Regarding the optical properties, depending on the origin of the OM precursor soil, different fluorescence peaks
- 33 can be observed on the WEOM fluorescence landscape; they may originate from humic fractions (Humic Acids 34 and Fulvic Acide), protein fractions (mainly tryptophan and tyrosine) or even from persistent contaminants [5,
- 34 and Fulvic Acide), protein fractions (mainly tryptophan and tyrosine) or even from persistent contaminants [5, 6]. Therefore, WEOM fluorescence landscape is considered as a reliable indicator of the quality of the whol
- 35 6]. Therefore, WEOM fluorescence landscape is considered as a reliable indicator of the quality of the whole soil 36 organic matter.
- 37 The geographical area of Doukkala, at the Atlantic coast of Morocco, is known for its important agricultural
38 activity and the diversity of its soils. However, an anterior study of these soils, relating to a 10 years 38 activity and the diversity of its soils. However, an anterior study of these soils, relating to a 10 years follow-up
39 between 1987 and 1997, revealed an OM depletion [7]. This is why physical-chemical investigations f 39 between 1987 and 1997, revealed an OM depletion [7]. This is why physical-chemical investigations focusing
40 on soils of this highly agricultural region are required. Additionally, during a previous study on the fluore 40 on soils of this highly agricultural region are required. Additionally, during a previous study on the fluorescence 41 analysis of pesticide remains in these soils [6, 8], the WEOM fluorescence was submerging fluorescence signals 42 of pesticides emitting in the same spectral range. This interference made it difficult to discriminate fluorescence 43 signals originating from pesticide remains having a low fluorescence yield.
- 44 The objective of the present investigation is the analysis of WEOM in the principal soil types in the agricultural
45 zone of Sidi Bennour, in the region of Doukkala. The approach consists in combining UV-Visible absorp zone of Sidi Bennour, in the region of Doukkala. The approach consists in combining UV-Visible absorption and 46 3D fluorescence supported by PARAFAC chemometrics. The goal is to calculate and take mean of the spectroscopic parameters E_2/E_2 and E_4/E_6 absorbance ratios. $S_{275-295}$ and $S_{250-400}$ spectral slopes and slope spectroscopic parameters E_2/E_3 and E_4/E_6 absorbance ratios, $S_{275-295}$ and $S_{350-400}$ spectral slopes and slope 48 ratios S_R , as well as fluorescence FI and humification HIX indices. On another side, exploring 3D fluorescence 49 landscapes by visual peak-picking and trough PARAFAC chemometrics will permit the resolution of the landscapes by visual peak-picking and trough PARAFAC chemometrics will permit the resolution of the
- 50 underlying fluorescing components.

51 Such optical spectrometric characteristics should make it possible the characterization of the WEOM in these
52 soils, and the discrimination of fluorescence signals originating from emitting organic pollutants persisti soils, and the discrimination of fluorescence signals originating from emitting organic pollutants persisting in the 53 soil.

54 **Soil Sampling and Analysis**

55 **Soils classification and samples preparation**

- 56 The investigated soils are classified into four main types according to two previous studies [9, 10]:
- 57 Vertisol, commonly labeled Tirs, (52%): deep and clayey soil, with a low permeability, characterized by a high 58 water retention and considered of good productivity.
- 59 Slightly evolved, commonly labeled Faid, (17%): deep soil, little to moderately permeable and characterized by
60 a good permeability. a good permeability.
- 61 Isohumic, commonly labeled Hamri, (16%): deep to moderately deep soil, with a sandy clay texture and 62 characterized by a good permeability.
- 63 Sandy / fersiallitic, commonly labeled R'mel, (13%): sandy in surface and coarse-textured soil, characterized 64 by a high permeability and a low water retention capacity.
- 65 These soils are neutral to moderately basic [10].

66 Thirty representative soil samples were collected in the province of Sidi Bennour in non-cultivated and non-67 treated areas neither with pesticides nor with fertilizers. The samples were taken at soil depths ranging from 5 to 68 20 cm, stored in plastic bags and then kept cold.

69 For each soil type, the samples were mixed and shared into several sub-samples of 20g. Avoiding the use of 70 organic solvents, each sub-sample was mixed with 40 ml of water, as an extractor medium, under 1 hour stirring 71 to improve the dissolution of the organic material. The extraction was made at room temperature (at about 20-
72 24°C), protected from light. The water-soil mixture was then stored in dark bottle in the fridge for 24 to 24°C), protected from light. The water-soil mixture was then stored in dark bottle in the fridge for 24 to 48 h to
73 decant; the final extracts showed pH values in the range 7.5–8.0. As no significant fluorescence intensi 73 decant; the final extracts showed pH values in the range 7.5–8.0. As no significant fluorescence intensity
74 difference has been observed between the spectra of filtered trough 0.45um size pores and non-filtered 74 difference has been observed between the spectra of filtered trough 0.45µm size pores and non-filtered 75 supernatants, the measurements were recorded directly on top-supernatants without filtration. These top-75 supernatants, the measurements were recorded directly on top-supernatants without filtration. These top-
76 supernatants were transferred into a quartz cell of 1 cm optical path for absorption and fluorescence analyses. 76 supernatants were transferred into a quartz cell of 1 cm optical path for absorption and fluorescence analyses.

77 **UV-Vis. absorption and fluorescence measurements**

- 78 The absorption spectra were measured on the top-supernatants using a JASCO UV/VIS/ NIR V-570 spectrophotometer; the scans covered the spectral range 200-800 nm. spectrophotometer; the scans covered the spectral range 200-800 nm.
- For each type of soil, the average values of the absorbance ratios $\frac{E}{E}$ $\frac{E_2}{E_3} = \frac{A}{A}$ $\frac{A_{250}}{A_{365}}$ and $\frac{E_4}{E_6} = \frac{A}{A}$ 80 For each type of soil, the average values of the absorbance ratios $\frac{E_2}{E_3} = \frac{A_{250}}{A_{365}}$ and $\frac{E_4}{E_6} = \frac{A_{465}}{A_{665}}$, the slopes
- $S_{275-295}$ and $S_{350-400}$ as well as slopes ratio $S_R = \frac{S}{6}$ 81 $S_{275-295}$ and $S_{350-400}$ as well as slopes ratio $S_R = \frac{S_{275-295}}{S_{350-400}}$ were calculated. A_{250} , A_{365} , A_{465} and A_{665} denote
- 82 respectively the measured absorbances at 250, 365, 465 and 665 nm wavelengths. The spectral slopes over the 83 intervals 275-295 nm and 350-400 nm were calculated using a linear regression of Ln(a_{λ}), where a_{λ} is the
- absorption coefficient calculated in m⁻¹ according to the equation $a_{\lambda} = \frac{2.303 \times A}{l}$ 84 absorption coefficient calculated in m⁻¹ according to the equation $a_{\lambda} = \frac{2.303 \times A(A)}{1}$, with A(λ) and l represent the
- 85 measured absorbance and the quartz cell thickness (i.e. 0.01 m) [11].

86 The total fluorescence spectra, or excitation-emission matrices also called TEEMs matrices, were measured

- 87 using a Shimadzu RF-5301PC spectrofluorimeter controlled by LabSolution software. Continuous fluorescence
- 88 scans were collected between 250 and 700 nm, varying the excitation wavelength from 220 to 550 nm at a 5 nm
- 89 step. The excitation and emission slits were set at 3 nm (exc.) / 3 nm (em.) or 5 nm (exc.) / 5 nm (em.) depending
- 90 on the sample.
- 91 The fluorescence matrix representation adopted in this work is the 2D one where the wavelengths $\lambda_{\text{emission}}$ and 92 $\lambda_{\text{avcitation}}$ appear explicitly on the X and Y axes, and the fluorescence intensity is coded by colors ra $\lambda_{\text{excitation}}$ appear explicitly on the X and Y axes, and the fluorescence intensity is coded by colors ranging from 93 blue to red in rising order of values. The fluorescence maxima correspond to peaks, or foci, identified by their 94 wavelengths coordinates λexc./λem. Each TEEM requires the concatenation of an average of 57 conventional 95 fluorescence spectra (excitation from 220 to 500 nm at a 5 nm step). LabSolution package crossed with Origin 96 data analysis and graphing software, allowed the representation and manipulation of the measured TEEMs 97 matrices.

98

99 **PARAFAC chemometric analysis**

 The PARAFAC "Parallel Factor Analysis" algorithm is a tool which consists in resolving the measured fluorescence matrices into individual sub-matrices corresponding to the fluorophores coexisting in the samples 102 analyzed (supernatants). This chemometric analysis makes it possible to separate the different spectral contributions associated with the different types of fluorophores in these samples. This means that the positions contributions associated with the different types of fluorophores in these samples. This means that the positions $\lambda_{\text{exc}}/\lambda_{\text{em}}$ of the various underlying signals in a TEEM fluorescence matrix will be determined with a minimal error; the PARAFAC algorithm is considered by several authors to be a mathematical chromatography. Water and soil total fluorescence analysis is the analytical method where PARAFAC is being used the most [12 - 14]. The algorithm has been largely elucidated by various authors [15, 16].

108 **PARAFAC algorithm principle**

109 The algorithm consists in the decomposition of the of the experimental TEEMs data cube, measured on all 110 samples, into a set of trilinear and residual terms as following.

111 The fluorescence intensity of the sample i for the excitation and emission wavelength pair (k, j) is given by the 112 equation:

$$
X_{i,j,k}=\sum_{f=1}^F a_{i,f}b_{j,f}c_{k,f}+\epsilon_{i,j,k}
$$

- 113 with $i = 1, 2, ...$ $i = 1, 2, ...$ and $k = 1, 2, ...$ K. I, J and K designate respectively the numbers of samples, 114 emission wavelengths and excitation wavelengths.
- 115 $f = 1, 2, \dots F$. F is the number of components in the model i.e. the presumed number of the fluorophores coexisting in the samples analyzed. coexisting in the samples analyzed.
- 117 $X_{i,j,k}$ represents the fluorescence intensity of sample i for the pair of excitation and emission wavelengths (k, j) .
- 118 a_{if} represents the fluorescence factor (the concentration times the fluorescence quantum yield); each PARAFAC 119 component f has I values of a (scores).
- 120 b_{i f} represents the intensity of the emission spectrum of the fluorophore f at the emission wavelength j.
- 121 c_{k} represents the value of the absorption spectrum of the fluorophore f at the excitation wavelength K 122 (proportional to the specific absorption coefficient).
- 123 $\epsilon_{i,j,k}$ is the residual representing the variability not accounted for by the model, and which the calculation searches to minimize. This term includes instrumental noises and other non-modellable variations. searches to minimize. This term includes instrumental noises and other non-modellable variations.
- 125 In this work, the number of components (fluorophores) was varied between 2 and 4. This choice of a reduced number of components is inspired by the measured TEEMs matrices showing a single fluorescence peak, large 126 number of components is inspired by the measured TEEMs matrices showing a single fluorescence peak, large and extended from 370 to 550 nm. The "split-half analysis" method integrated into the algorithm was chosen 127 and extended from 370 to 550 nm. The "split-half analysis" method integrated into the algorithm was chosen
128 during the model validation step. The version of the PARAFAC algorithm used is the online available one [17 128 during the model validation step. The version of the PARAFAC algorithm used is the online available one [17];
129 the graphics program used to create the matrices is Matlab. the graphics program used to create the matrices is Matlab.
- 130 The successful decomposition of a multi-way data system using the PARAFAC algorithm relies on assumptions 131 about variability, tri-linearity and additivity [15, 16]. In this work, the fluorescence spectra were measured on 132 samples revealing absorbances at 254 nm lower than 0.5 as can be observed on figure 1; no apparent internal 133 filter effect was observed. The experimental matrices were corrected from all of the instrumental response, the 134 internal filter effect and the Raman scattering [16 - 18].

135 **Results and discussion**

136 **UV-Visible Absorption: absorbance and slope ratios**

In Table 1 are reported, for each type of soil, the average values of the absorbance ratios $\frac{E_2}{E_3} = \frac{A}{A}$ 137 In Table 1 are reported, for each type of soil, the average values of the absorbance ratios $\frac{E_2}{E_3} = \frac{A_{250}}{A_{365}}$ and $\frac{E_4}{E_6}$ A $\frac{A_{465}}{A_{665}}$, the slopes $S_{275-295}$ and $S_{350-400}$ as well as the ratios of these slopes $S_R = \frac{S}{S}$ 138 $\frac{A_{465}}{A_{665}}$, the slopes $S_{275-295}$ and $S_{350-400}$ as well as the ratios of these slopes $S_R = \frac{5275-295}{S_{350-400}}$

- 139 E_A/E_f ratio is generally related to the degree of condensation and aromaticity of the organic material in the soil
- 140 [19, 20]; this ratio varies inversely with respect to the degree of condensation / aromaticity. The values found in
- 141 the case of the four soil types studied, ranging between 3.26 and 4.82, suggest that these soils contain organic 142 materials in close humification stages. The highest value, 4.82, corresponding to the soil Faid is indicative of an
- 143 organic material with an aliphatic prevalence of this soil compared to the other ones [20].This result is also
- 144 confirmed by the values obtained for E_2/E_3 absorbance ratios varying from 3.77 to 4.33, similar to those reported for other soils [21]. E_2/E_3 ratio is associated with the average weight and molecular size of the reported for other soils [21]. E_2/E_3 ratio is associated with the average weight and molecular size of the organic
- 146 material contained in the soil [22], and reflects the relative contents of humic acids (HAs) and fulvic acids (FAs).
- 147 In the present case the values found for E_2/E_3 ratios, all greater than 3.5, indicate organic materials with FAs)
- 148 contents greater than those of HAs [23, 24].
- 149 Concerning the slopes $S_{275-295}$ and $S_{350-400}$, the average values found for all soil types range respectively in the 150 intervals 0.0101- 0.0141 nm⁻¹ and 0.0122 - 0.0152 nm⁻¹, and are consistent with those reported for other soils 151 [11, 25]. Moreover, the obtained S_R slope ratios, all lower than 1, are reasonably characteristic of terrigenous 152 organic materials generally characterized by $S_{275-295}$ lower than $S_{350-400}$. than $S_{350-400}$. The difference, of 153 about 20%, observed in S_R between Tirs and R'mel may be probably related to a significant difference in 154 chromophoric DOM (CDOM) in these soils which are respectively the darker and the lighter ones. Furthermore, as slopes and slope ratios are commonly related to the average molecular weight of the organic matter, the as slopes and slope ratios are commonly related to the average molecular weight of the organic matter, the 156 similarity of these ratios reveals that the four soil kinds contain organic materials with similar molecular weights 157 [11]. Subsequently, these soils are very likely in similar humification levels as indicated before by E_2/E_3 and 158 E_4/E_6 absorbance ratios.

160 **Fig. 1** UV-Vis. Absorption spectra measured on top-supernatants of water-soil mixtures.

- 159
-
- 161
- 162
- 163
- 164

165 **Table 1** Averaged spectroscopic parameters calculated from absorption and fluorescence spectra*

166

167 * More details on the shown parameters are in the text.

168 **Fluorescence: FI and HIX indices**

The fluorescence index $FI = \frac{I}{I}$ 169 The fluorescence index $FI = \frac{r_{\text{II.}(A_{\text{em}}=450 \text{ nm})}}{I_{\text{II.}(A_{\text{em}}=500 \text{ nm})}}$, calculated as the fluorescence intensities ratio at the excitation 170 wavelength $\lambda_{\text{exc}} = 370$ nm, provides information on the origin of the soil organic material; it highlights in 171 particular the relative importance of the terrigenous material with respect to the microbial biomass one [26].

172 An FI index above 1.8 corresponds to a material with a microbial precursor, while an FI index close to 1.2 is 173 indicative of a material with a rather terrigenous precursor [27]. In the case of the investigated soils, the FI 174 indices range in the interval 1.52-1.68. This order of magnitude indicates organic materials from terrigenous

175 sources with a certain microbial biological component. A similar result was observed in soils from China [28].

The humification index HIX calculated as the ratio of fluorescence intensity areas HIX = $\frac{\Sigma_{\Lambda}^{\lambda}}{\Sigma_{\Lambda}}$ 176 The humification index HIX calculated as the ratio of fluorescence intensity areas HIX = $\frac{2\lambda_{\text{em}}-435^{-11}}{\Sigma_{\lambda_{\text{em}}-345}}$, while

177 excitation wavelength is set at 254 nm, is a parameter positively correlated with the humification / maturation 178 degree of the soil organic matter [26]. It is often taken as an indicator of DOM variation in agricult 178 degree of the soil organic matter [26]. It is often taken as an indicator of DOM variation in agricultural soils [29].
179 In the present case, the HIX indices vary from 9 to 12. In comparison with the reference interv

In the present case, the HIX indices vary from 9 to 12. In comparison with the reference intervals given by

180 various authors [30, 31] the investigated soils can be considered as materials with an important humic character

181 as a consequence of an advanced maturation stage.

182 **Total fluorescence and PARAFAC chemometrics**

183 **Experimental TEEMs**

184 All soil extracts in water, prepared as described above, were subjected to total fluorescence measurements. In

185 Figure 2 are presented four examples of the collected raw experimental TEEMs matrices. Likewise, these typical

186 matrices show a single large fluorescence peak extended, in emission, from 370 to 550 nm. This signal is 187 attributable to humic substances in the soil organic matter [32, 33]. However, no signal attributable to any protein fraction has been observed.

protein fraction has been observed.

190 **Fig. 2** Examples of experimental total excitation emission matrices (TEEMs).

191 **PARAFAC Analysis**

192 The PARAFAC chemometric analysis has made it possible to discriminate the fluorescent components 193 previously overlapping each other in the experimental TEEMs matrices; the procedure is described above. For all soils, the validated model corresponds to a number of components equal to two, C_1 and C_2 . This mode all soils, the validated model corresponds to a number of components equal to two, C_1 and C_2 . This model is 195 entirely consistent with the experimental TEEMs that did not reveal any fluorescence signal outside the spectral 196 emission range 370-550 nm. Figure 3 shows the individual matrices corresponding to the PARAFAC components obtained for the four soil types. In all cases the component C_1 appearing around 340exc./442em. 197 components obtained for the four soil types. In all cases the component C_1 appearing around 340exc./442em.
198 corresponds to the fulvic acid (FAs) fraction while the component C_2 appearing around 385exc./465em. 198 corresponds to the fulvic acid (FAs) fraction while the component C_2 appearing around 385exc./465em.
199 corresponds to the humic acid (HAs) one of the soils WEOM. Indeed, these fractions are generally observed in 199 corresponds to the humic acid (HAs) one of the soils WEOM. Indeed, these fractions are generally observed in
1200 the respective intervals (320-350 nm) exc./ (400-450 nm) em. and (370-390 nm) exc./ (460-500 nm) em. [34 the respective intervals (320-350 nm) exc./ (400-450 nm) em. and (370-390 nm) exc./ (460-500 nm) em. [34, 35]. 201

202 In all soils, the ordering of FAs fraction as component C_1 and HAs fraction as component C_2 is entirely consistent 203 with the obtained values of E2/E3 absorbance ratios which are greater than 3.5 indicating an 203 with the obtained values of E2/E3 absorbance ratios which are greater than 3.5 indicating an organic material 204 with FAs prevalence. Hence, in the whole of the investigated soils, these findings reflect a dominance o 204 with FAs prevalence. Hence, in the whole of the investigated soils, these findings reflect a dominance of the FAs
205 fraction with respect to the HAs one [36]. Furthermore, the absence of any PARAFAC component compati 205 fraction with respect to the HAs one [36]. Furthermore, the absence of any PARAFAC component compatible 206 with a protein fraction, such as tyrosine or tryptophan, reveals highly humified organic materials as previous 206 with a protein fraction, such as tyrosine or tryptophan, reveals highly humified organic materials as previously
207 highlighted by the HIX humification indices ranging between 9 and 12. The absence of any protein-type 207 highlighted by the HIX humification indices ranging between 9 and 12. The absence of any protein-type fluorescence signal indicates the absence or, at least, the weakness of proteins microbiological activity in the soi fluorescence signal indicates the absence or, at least, the weakness of proteins microbiological activity in the soil 209 lands investigated; this is a consequence of the revealed advanced stage of humification. Probably, in the present 210 soils, the formation of humic subtances may be governed by the lignin mechanism putting forward the 211 hypothesis of the transformation of HAs into FAs [37]. Soil humification means the process leading to the 212 increasing of the humic material content (namely FAs and HAs); an advanced humification stage may be 213 associated with an increase in FAs concentration with a decrease in HAs concentration [38]. The formation of 214 humic substances (HAs, FAs and Humins) is one of the least understood aspects of humus chemistry and one of 215 the most intriguing.

219

216

220 **Conclusion**

221 The present study focused on the WEOM in a set of thirty agricultural soil samples collected from the zone of 222 Doukkala in Morocco. The set has covered the four types of agricultural soil (Tirs, Faid, Hamri and R'mel),
223 while the analysis was performed on the soils extracts in water by means of UV-visible absorption and tota while the analysis was performed on the soils extracts in water by means of UV-visible absorption and total 224 fluorescence combined to PARAFAC chemometrics.

- 225 E_4/E_6 and E_2/E_3 absorbance ratios, ranging respectively in the intervals 3.26 4.82 and 3.77- 4.33, revealed that
- 226 all soils contain organic materials in similar humification stages, with a predominance of FAs fractions with
- 227 respect to HAs ones. The average spectral slopes $S_{275-295}$ and $S_{350-400}$ varying respectively in the intervals
- 228 $0.0101 0.0141$ nm⁻¹ and $0.0122 0.0152$ nm⁻¹, with slope ratios S_R lower than 1, are characteristic of terrigenous
- 229 organic materials. In addition, the FI fluorescence indices found between 1.52 and 1.68 revealed organic materials from terrigenous sources, having some microbial biological component. In parallel, HIX humification
- 230 materials from terrigenous sources, having some microbial biological component. In parallel, HIX humification indices ranging from 9 to 12 indicated that the soils are in advanced humification stages. indices ranging from 9 to 12 indicated that the soils are in advanced humification stages.
- 232 3D fluorescence did not detect any protein fraction in the soils WEOM studied, highlighting thus poor fresh
233 organic matter soils. The PARAFAC chemometric analysis, in full agreement with experiments, revealed only 233 organic matter soils. The PARAFAC chemometric analysis, in full agreement with experiments, revealed only
234 two fluorescent components in the humic material with FAs fraction classified as component 1 while HAs one two fluorescent components in the humic material with FAs fraction classified as component 1 while HAs one 235 classified as component 2.
- 236 Consequently, the four types of agricultural soils are compartments in similar humification stages, containing 237 organic materials dominated by FAs fractions and very poorly provided in fresh organic matter.
- 238 Considering the issue of pesticide remains, the accomplished 3D fluorescence/PARAFAC characterization of the 239 WEOM in the investigated soils will make it possible to discriminate additional fluorescence signals originating 140 from any fluorescent contaminants persisting in the soil. from any fluorescent contaminants persisting in the soil.
- 241 **Authors' contributions.** Hassan Ba-Haddaou: (Experimental investigations, first draft preparation and submission). Hicham Hassoun: (Sampling, material preparation and experimental investigations). Salim Foudeil: 242 submission). Hicham Hassoun: (Sampling, material preparation and experimental investigations). Salim Foudeil: 243 (Data collection, first draft preparation and submission). Saadia Ait Lyazidi: (Supervision, methodology, final 244 draft reviewing and project administration). Mustapha Haddad: (Discussion, validation and funding acqu 244 draft reviewing and project administration). Mustapha Haddad: (Discussion, validation and funding acquisition).
245 Matthieu Masson: (Discussion and validation). Marina Coquery: (Discussion and validation). Christelle 245 Matthieu Masson: (Discussion and validation). Marina Coquery: (Discussion and validation). Christelle 246 Margoum: (Experimental logistics, discussion and validation). Margoum: (Experimental logistics, discussion and validation).
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- 250 **Data Availability** Not applicable.
- 251 **Declarations**
- 252 **Conflicts of interest/Competing interests** The authors declare they have no conflicts of interest and no 253 competing interests.
- 254 **Ethics Approval** Not applicable as the study does not include any use of animals and humans.
- 255 **Consent to Participate** All authors consent to participate in the research.
- 256 **Consent for Publication** All authors consent to participate in the publication of the research.

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