

Prediction of soil organic and inorganic carbon concentrations in Tunisian samples by mid-infrared reflectance spectroscopy using a French national library

Cécile Gomez, Tiphaine Chevallier, Patricia Moulin, Imane Bouferra, Kaouther Hmaidi, Dominique D. Arrouays, Claudy C. Jolivet, Bernard

Barthès

▶ To cite this version:

Cécile Gomez, Tiphaine Chevallier, Patricia Moulin, Imane Bouferra, Kaouther Hmaidi, et al.. Prediction of soil organic and inorganic carbon concentrations in Tunisian samples by midinfrared reflectance spectroscopy using a French national library. Geoderma, 2020, 375, pp.114469. 10.1016/j.geoderma.2020.114469. hal-02890493

HAL Id: hal-02890493 https://hal.inrae.fr/hal-02890493

Submitted on 23 Aug 2022

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License

Version of Record: https://www.sciencedirect.com/science/article/pii/S0016706119323961 Manuscript_1c04d245da3ec17786ab0bceb7c80d8d

1	Prediction of soil organic and inorganic carbon concentrations in
2	Tunisian samples by mid-infrared reflectance spectroscopy using a
3	French national library
4	
5	Cécile Gomez ^{1,2} , Tiphaine Chevallier ³ , Patricia Moulin ⁴ , Imane Bouferra ³ , Kaouther
6	Hmaidi ^{3,5} , Dominique Arrouays ⁶ , Claudy Jolivet ⁶ , Bernard G. Barthès ³
7	
8	¹ LISAH, University of Montpellier, INRAE, IRD, Montpellier SupAgro, 34060 Montpellier,
9	France. Corresponding author. cecile.gomez@ird.fr
10	² Indo-French Cell for Water Sciences, IRD, Indian Institute of Science, Bangalore
11	560012, India
12	³ Eco&Sols, University of Montpellier, CIRAD, INRAE, IRD, Montpellier SupAgro, 34060
13	Montpellier, France
14	⁴ US IMAGO, IRD, BP1386, Dakar, Senegal
15	⁵ UR Pédologie, Faculté des Sciences de Tunis, El Manar Tunis, Tunisia
16	⁶ INRAE, US 1106 InfoSol, F45000, Orléans, France
17	
18	
19	Keywords: Soil Organic and Inorganic Carbon; Mid-Infrared reflectance spectroscopy;
20	natural logarithm transformation; Local and Global calibration; Partial least squares
21	regression (PLSR); National dataset.
22	
23	Abstract
24	Mid-infrared reflectance spectroscopy (MIRS, 4000-400 cm ⁻¹) is being considered to

25 provide accurate estimations of soil properties, including soil organic carbon (SOC) and

soil inorganic carbon (SIC) contents. This approach has mainly been demonstrated by 26 using datasets originating from the same area A, with similar geopedological conditions, to 27 28 build, validate and test prediction models. The objective of this study was to analyse how 29 MIRS performs when applied to predict SOC and SIC contents, from a calibration database collected over a region A, to predict over a region B, where A and B have no 30 31 common area and different soil and climate conditions. This study used a French MIRS 32 soil dataset including 2178 topsoil samples to calibrate SIC and SOC prediction models 33 with partial least squares regression (PLSR), and a Tunisian MIRS topsoil dataset 34 including 96 soil samples to test them. Our results showed that when using the French 35 MIRS soil database, *i*) the SOC and SIC of French validation samples were successfully predicted using global models ($R^{2}_{val} = 0.88$ and 0.98, respectively), *ii*) the SIC of Tunisian 36 37 samples was also predicted successfully both using a global model and using a selection of spectral neighbours from the French calibration database (R^{2}_{test} of 0.96 for both), *iii*) the 38 SOC of Tunisian samples was predicted moderately by global model (R^{2}_{test} of 0.64) and a 39 40 transformation by natural logarithm of the calibration SOC values significantly improved the SOC prediction of Tunisian samples (R^{2}_{test} of 0.97), and *iv*) a transformation by natural 41 logarithm of SOC values provided more benefit than a selection of spectral neighbours 42 43 from the French calibration database for predicting Tunisian SOC values. Therefore, in the future, MIRS might replace conventional physico-chemical analysis techniques, or at least 44 be considered as an alternative technique, especially when optimally exhaustive 45 46 calibration databases will become available.

47

48 **1. Introduction**

Soil is the largest reservoir of continental carbon and participates in the global carbon
cycle (Jacobson et al., 2000; Scharlemann et al., 2014). Soil emits carbon dioxide (CO₂)
through autotrophic and heterotrophic respiration and acts as a sink of atmospheric CO₂

through photosynthesis; then, organic decomposition products are integrated into the soil as organic matter, which is composed of approximately 50% to 58% of carbon (Gregorich et al., 1994; Pribyl, 2010). Moreover, soil organic carbon (SOC) has a long-acknowledged and key role in soil physical, chemical and biological fertility (Reeves, 1997). Thus, understanding the dynamics of soil carbon is a major issue for soil fertility and for climate change mitigation.

58 Soil carbon is not only found in organic form. At the global scale, approximately one-59 third of the total soil carbon is inorganic (soil inorganic carbon, SIC) (Batjes, 1996), and 60 calcareous soils cover more than 30% of the earth's land surface (Chen and Barak, 1982; 61 Romanyà and Rovira, 2011). SIC is mainly in the form of calcium carbonate (CaCO₃), and high SIC contents are often localised in dry areas where SOC stocks are low. In these 62 soils, SIC levels can be 2 to 10 times higher than SOC levels (Bernoux and Chevallier, 63 64 2014). SIC is made of primary minerals derived from the fragmentation of carbonate bedrock (lithogenic carbonates) or secondary minerals from inorganic carbon precipitation 65 66 in soil pores or around roots (pedogenic carbonates). Pedogenic carbonates have various forms-nodules, lamellae or crystals-and have varying solubility. 67

Several analytical methods have been developed to guantify SIC and SOC contents in 68 69 soils, but they are tedious and/or costly. In particular, SOC determination in carbonated 70 soils often requires hazardous reagents. SIC content has usually been measured by calcimetry (ISO, 1995b) but can also be measured by dry combustion with a CNH 71 72 elemental analyser equipped with a specific module (CO₃-C module) after phosphoric acid dissolution of the SIC (Mc Crea, 1950; Hannam et al. 2016). Quantifying SOC in 73 calcareous soils has either been carried out directly, by wet oxidation (Walkley and Black, 74 1934) or dry combustion after removing SIC by acid pretreatment (Harris et al., 2001), or 75 indirectly, by subtracting the SIC, measured by calcimetry, from the total carbon content 76 determined by dry combustion. Due to indirect determination, incomplete oxidation and the 77

use of hazardous reactants with both of these methods, alternative methods based on the
thermal lability of SIC and SOC have also been tested (Wang et al., 2012; Apesteguia et
al., 2018).

81 For one to two decades, visible near-infrared (Vis-NIR, 400-2500 nm) and midinfrared reflectance spectroscopy (MIRS, 4000-400 cm⁻¹), which measures diffuse 82 83 reflectance, have been proposed as alternative methods to these physico-chemical 84 analytical methods (e.g., Viscarra Rossel et al., 2006; Cécillon et al., 2008; Bellon-Maurel 85 and McBratney, 2011). MIRS is based on the study of absorption bands corresponding to 86 fundamental molecular vibrations, and NIRS is based on the study of absorption bands 87 corresponding to overtones and combinations of fundamental vibrations (Williams and Norris, 1987). Several studies highlighting the potential of Vis-NIR and/or MIRS for 88 89 predicting various soil attributes, including SIC and SOC, have been listed by Viscarra 90 Rossel et al. (2006) and then by Soriano-Disla et al (2014).

In the MIR range, carbonates may be identified by strong and numerous absorption 91 92 bands, for instance, bands at approximately 820-750 cm⁻¹, 1800 cm⁻¹, 2520 cm⁻¹ and 93 2900-2990 cm⁻¹ (e.g., Tatzber et al., 2007; Du and Zhou, 2009, Comstock et al., 2019). In the NIR range, carbonates may be identified by peaks at 2341 and 2480 nm (Lagacherie 94 95 et al., 2008; Barthès et al., 2016). In the MIR range, organic carbon may also be identified by numerous absorption peaks, for instance, peaks at approximately 2920 and 1230 cm⁻¹ 96 (Grinand et al., 2012). In the NIR range, organic carbon may be identified by absorption 97 98 peaks at approximately 1910 nm (Viscarra Rossel et al., 2006; Viscarra Rossel and 99 Webster, 2012) and 2050-2150 nm (Workman and Weyer, 2008). Finally, MIRS has 100 generally been reported to provide more accurate performance in terms of SOC and SIC 101 predictions compared to NIRS (e.g., McCarty et al., 2002; Reeves, 2010; Bellon-Maurel & McBratney, 2011; Clairotte et al., 2016). Nevertheless, better NIRS than MIRS predictions 102 of SOC have been reported in tropical and Mediterranean regions due to the overlap of 103

absorption regions related to metal oxides and organic compounds (Rabenarivo et al.,
2013; Barthès et al., 2016).

106 Following the emergence of infrared spectroscopy technologies for soil 107 characterisation, soil spectral libraries covering extensive areas have recently been 108 developed for estimating soil properties, especially SOC; most libraries are in the Vis-NIR range at the national scale (e.g., in Australia, Viscarra Rossel and Webster, 2012; in 109 110 Denmark, Knadel et al., 2012; in France, Gogé et al., 2014; in China, Shi et al., 2015), 111 continental scale (in Europe, Stevens et al., 2013) and even global scale (Brown et al., 112 2006; Viscarra Rossel et al., 2016), but libraries also exist in the MIR range (e.g., in 113 France, Grinand et al., 2012, and Clairotte et al., 2016; in the US, Wijewardane et al., 2018, Comstosck et al., 2019 and Dangal et al., 2019). All these libraries use spectra 114 115 collected using dried and ground samples in laboratory conditions.

116 Most of the studies dealing with large soil spectral databases (national or continental) have aimed to calibrate prediction models with samples from a region A to 117 118 predict the properties of soil samples from the same region A. Therefore, the soil and climate conditions are similar between the calibration and validation databases. McCarty et 119 al. (2002) calibrated SOC and SIC prediction models by using two-thirds of 257 soil 120 121 samples collected from 14 geographically diverse locations over eight states in the west 122 central US and validated the models by using the remaining one-third of soil samples, obtaining good accuracy and low bias. Stevens et al. (2013) used the LUCAS database, 123 124 which includes samples from 23 member states of the European Union, to predict SOC. In 125 their study, each calibration dataset and associated validation dataset was composed of 126 samples from a similar soil type (organic or mineral) and land use (cropland, grassland or 127 woodland), and the most spectrally representative samples were selected as calibration samples. Shi et al. (2015) used a Chinese database, which included samples from 20 128 provinces, to predict SOC. In their study, the calibration dataset was selected to minimise 129

both the spectral distance and geographical distance to each validation sample, which also belonged to the database (region *A*). Clairotte et al. (2016) successfully calibrated prediction models by using subsets of the French national database and then tested these models on a test subset (10% of this French database, either spectrally representative or not; samples were not selected based on their pedo-climatic context).

135 Some studies have focused on potential and limitation analysis of prediction models 136 calibrated with a large database composed of samples collected over a region A to predict 137 properties for soil samples collected from a small region b within A. McCarty et al. 138 (2002) calibrated SOC and SIC prediction models by using 257 soil samples collected 139 from 14 geographically diverse locations over 8 states in the west central US and obtained biased predictions when their models were applied to 16 independent soil samples 140 141 collected in another state (Nebraska) that was also in the west central US. Gogé et al. (2014) calibrated local prediction models from the French national database, collected 142 over 550 000 km² (region A), to predict soil properties in soil samples collected from a 143 144 small French area of 24 km² (Occitanie region, south of France; this region b was included in A but under-represented). Additionally, Comstosck et al. (2019) calibrated prediction 145 models by using a US national database (region A) to predict carbonate in soil samples 146 147 collected from two states (New York and Iowa, regions b) that were poorly represented in 148 the US database.

Finally, to the best of our knowledge, few studies have focused on potential and limitation analysis of prediction models calibrated with a large database composed of samples collected over a region *A* to predict properties for soil samples collected over a region *B*, where *A* and *B* have no common area and hence have potential differences in soil and climatic conditions. Only Jauss et al. (2017) and Ahmed et al. (2017) used MIR spectroscopy as a routine method for predicting pyrogenic carbon and for predicting Total

155 Carbon and SOC, respectively, on United States soils (region *B*), from a soil spectral 156 database of Australia (region *A*).

The objective of this study was to analyse how MIRS may be used to predict SOC 157 158 and SIC contents when using a national database collected over a region A to predict values for soil samples collected over a region *B*, where *A* and *B* have no common area 159 160 and have different soil and climate conditions. This study used a French MIRS soil dataset 161 (region A) including 2178 topsoil samples collected from an area of 550 000 km² (French 162 metropolitan territory, composed of temperate and Mediterranean soils) to calibrate SIC and SOC prediction models. These models were tested on a Tunisian MIRS soil dataset 163 164 (region *B*), including 96 soil samples collected from an area of approximately 80 000 km² 165 (northern half of Tunisia, mainly Mediterranean and arid soils).

166

167 **2. Materials and methods**

168

169 **2.1. Soil datasets**

170

2.1.1. The French national soil collection

The national soil collection provided by the French national soil quality monitoring network 171 172 (RMQS; Arrouays et al., 2002) and called DB RMQS was used in this study to calibrate the SIC and SOC models. This RMQS collection is composed of 2178 soil samples 173 representing all main soil types encountered over the sampled 552 000 km² of the French 174 metropolitan territory (Corsica included): Cambisols, Calcosols, Luvisols, Leptosols, 175 Andosols, Albeluvisols, Podzosols, etc. (IUSS Working Group WRB, 2014). The latitude of 176 sample sites ranges from 41 to 51°N, and their longitude ranges from 5.0°W to 9.5°E. The 177 178 sampling design was based on a square grid with 16-km spacing. At the centre of each 179 square, 25 individual core samples were taken from 0 to 30 cm depth using an unaligned

180 sampling design within a 20×20 m area. Core samples were bulked to obtain a composite 181 sample for each site (Arrouays et al., 2002).

182

183

2.1.2. The Tunisian soil samples

Ninety-six soil samples were used as the test set, called DB Tunisia. These samples were 184 185 collected from 45 localities, covering approximately 80 000 km² (from 35 to 37°N and 08 to 186 11°E), with the aim of representing the main soil types and land uses of the northern half of 187 Tunisia. This was done based on previous studies carried out at the Tunis El Manar 188 University, without particular design. Field samples within the same locality were 189 kilometres apart and under different land uses. Soil samples were collected at 0-10 cm 190 using a spade, and the sampling campaign was carried out within a few months in late 191 2010. This Tunisian set was previously studied in Barthès et al. (2016).

192

193

194

2.2.1. Physico-chemical analyses

2.2. Laboratory Analysis

The 2178 RMQS samples were air dried, 2-mm sieved and then finely ground (< 0.25 mm) 195 196 using mortar and pestle. The SIC content of the RMQS samples was calculated as 0.12 197 times the soil calcium carbonate content, which was determined using these finely ground (< 0.25 mm) air-dried samples using a Bernard calcimeter according to the standard 198 procedure ISO 10693 (ISO, 1995a). The carbonate content was calculated after calibration 199 200 with a pure calcium carbonate standard and was expressed as the equivalent calcium carbonate content. The SIC content of the RMQS samples (DB RMQS SIC) ranged from 201 0 to 103.9 g kg⁻¹, averaged 6.4 g kg⁻¹, and had a median of 0 g kg⁻¹ and a skewness value 202 203 close to 3.1 (Table 1).

Then, the SOC content of the RMQS samples was calculated as the difference between the total carbon (TC) and inorganic carbon contents. The TC content was

determined by dry combustion with an elemental analyser (Thermo Fisher Scientific CHN NA2000, Waltham, MA, US) using approximately 25–30-mg aliquots of finely ground (< 0.25 mm) air-dried soil samples that were sealed into tin capsules, according to the standard procedure ISO 10694 (ISO, 1995b). The SOC content of the RMQS samples (DB_RMQS_SOC) ranged from 0.6 to 411.3 g kg⁻¹, averaged 25.8 g kg⁻¹, and had a median of 19.6 g kg⁻¹ and a skewness value close to 4.9 (Table 1).

- 212
- 213

[Table 1]

- 214
- 215

216 The 96 Tunisian soil samples were also air-dried, sieved to 2 mm and then finely ground (< 0.2 mm) using mortar and pestle. The SOC content of the Tunisian samples was 217 analysed by dry combustion after decarbonisation using chlorhydric acid, following the 218 standard procedure ISO 10694 (1995b), with the same elemental analyser as that used for 219 220 RMQS samples but using silver capsules. Soils were decarbonated prior to SOC determination: 10 mL of water were added to 1 g of soil and 0.5 M HCl solution was then 221 dripped onto the sample until there was no more effervescence; then the samples were 222 223 washed in water until pH reached 7. The SOC concentration was then determined on finely ground 25–30 mg aliquots by dry combustion using an elemental analyser (Thermo 224 Fisher Scientific CHN NA2000, Waltham, MA, USA). The SOC content of the Tunisian 225 samples (DB Tunisia SOC) ranged from 2.0 to 121.0 g kg⁻¹, averaged 20.1 g kg⁻¹, and 226 had a median of 14.6 g kg⁻¹ and a skewness value close to 3 (Table 1). 227

The soil inorganic carbon content of the Tunisian samples was calculated as the difference between the TC (determined by dry combustion using the same CHN analyser as that used for the RMQS samples) and SOC contents. The SIC content of the Tunisian

samples (*DB_Tunisia_SIC*) ranged from 0.0 to 92.9 g kg⁻¹, averaged 43.3 g kg⁻¹, and had
a skewness value close to -0.2 (Table 1).

233 As the SIC contents of the RMQS and Tunisian samples were analysed by two 234 different methods, 29 test samples from the Tunisian set (i.e., 30% of the set) were reanalysed following the same method as that used for the RMQS samples (i.e., directly, by 235 236 calcimetry; ISO, 1995a). The Pearson correlation coefficient (R), root mean square error 237 (RMSE) and bias between the SIC values determined by both approaches (calcimetry vs. 238 difference between TC and SOC) were 0.997, 2.4 g kg⁻¹ and 1.1 g kg⁻¹, respectively 239 (Figure 1). The values of SIC calculated by both approaches could thus be considered 240 equivalent.

- 241
- 242

[Figure 1]

- 243
- 244
- 245

2.2.2. Mid-infrared spectroscopy

Mid-infrared spectroscopic analysis was performed following the same procedure for both 246 spectral libraries. First, air-dried, 2-mm sieved, and 0.2-mm ground samples were oven-247 248 dried at 40°C for twelve hours. Reflectance spectra were acquired using a Fourier 249 transform Nicolet 6700 spectrophotometer (Thermo Fischer Scientific, Madison, WI, US) in the MIR region. Reflectance was acquired at 934 wavenumbers between 4000 and 250 400 cm⁻¹ with a 3.86 cm⁻¹ spectral resolution. This spectrophotometer is equipped with a 251 252 silicon carbide source, a Michelson interferometer as a dispersive element, and a 253 deuterated triglycine sulfate detector. Soil samples were placed in a 17-well plate. The soil 254 surface was flattened with the flat section of a glass cylinder, and samples were then scanned using an auto-sampler (soil surface area scanned: ca. 10 mm²). Each spectrum 255 resulted from 32 co-added scans, and the body of the plate (next to the wells) was used as 256

a reference standard and scanned once per plate (i.e., every 17 samples). Twenty
 wavenumbers were removed due to frequent noise in the spectrum, so MIR spectra in the
 range from 4000 to 478 cm⁻¹, with 914 wavenumbers, were used.

260

261

2.3 PLSR model calibration

All procedures were performed using R software (R Core Team, 2012), and both the ade4 (Dray and Dufour, 2007) and pls packages (Mevik and Wehrens, 2007) were used.

- 264
- 265

2.3.1. Dataset preparation

266 Both DB RMQS SIC and DB RMQS SOC were divided into a calibration set (3/4 of the dataset) and a validation set (1/4 of the dataset). The samples of each dataset were 267 ranked according to ascending reference value (observed SIC or SOC). The sample with 268 the lowest reference value was put in the calibration set, the next sample was put in the 269 validation set, and then the next three samples were put in the calibration set. The 270 271 procedure was continued by alternately placing the next sample in the validation set and the following three samples in the calibration set. Following this process, the distributions 272 273 of the DB Calib RMQS SOC calibration and DB Valid RMQS SOC validation datasets 274 were similar. As well, the distributions of the DB Calib RMQS SIC calibration and *DB_Valid_RMQS_SIC* validation datasets were similar. 275

As the SOC values of the RMQS dataset followed a non-normal distribution (Table 1), the SOC values of the *DB_Calib_RMQS_SOC* calibration dataset were transformed with natural logarithm (In(SOC)) to reach a normal distribution, giving rise to new dataset for calibration *DB_Calib_RMQS_InSOC*. The SIC distribution in the French dataset was also non normal, but due to the very large number of null values, In-transformation was hardly possible (Bellon-Maurel et al., 2010; Terra et al., 2015).

The reflectance was converted into "absorbance" (log₁₀ [1/reflectance]), and a 282 standard normal variate correction was applied to remove additive and multiplicative 283 284 effects (Barnes et al., 1989). Spectral outliers, which are defined as samples spectrally 285 different than the rest of the samples (e.g., Pearson, 2002), were removed from the calibration datasets. The spectral outliers were identified by applying the Mahalanobis 286 287 distance (Mark and Tunnell, 1985) to data condensed by principal component analysis 288 (PCA). In the present study, a Mahalanobis distance of 3.5 was selected as the threshold 289 for the identification of spectral outliers.

- 290
- 291

2.3.2. Partial least squares regression

292 Partial least squares regression (PLSR) is a multivariate approach that specifies a linear 293 relationship between a dependent (response) variable (Y-variable, i.e., SIC or SOC 294 content in the present case), and a set of predictor variables (X-variables, i.e., MIR spectra in the present case; Tenenhaus, 1998). The general concept of PLSR is to extract a small 295 296 number of orthogonal variables (called the latent variables) that account for the maximum variation in the X-variables. A detailed description of the PLSR procedure can be found in 297 298 Wold et al. (2001). This method is commonly used for NIRS or MIRS prediction of soil properties (e.g., Viscarra-Rossel et al., 2006; Bellon-Maurel et al, 2010). 299

The maximum number of latent variables of PLSR was defined as 30. A leave-oneout cross-validation (LOOCV) procedure was adopted to verify the prediction capability of the PLSR model for the calibration set. Each time, n - 1 samples were used to build a regression model, which was applied to the sample not used in developing the model. This procedure was repeated for all n samples, resulting in predictions for all n samples.

305

306

2.3.3. Global models

307 Global calibration is a common calibration procedure where all calibration samples are used to build a unique prediction model that is applied identically to all validation or test 308 309 samples. One global prediction model (denoted GM_{SIC}) was built for SIC prediction based 310 on DB Calib RMQS SIC, validated on DB Valid RMQS SIC and tested on DB Tunisia SIC. As well, a global prediction model (denoted GM_{SOC}) was built for SOC 311 prediction based on DB Calib RMQS SOC, validated on DB Valid RMQS SOC and 312 313 tested on *DB Tunisia SOC*. Finally, a global prediction model (denoted *GM*_{InSOC}) was built 314 for SOC prediction based on DB Calib RMQS InSOC, applied to spectra of 315 DB Valid RMQS SOC and DB Tunisia SOC, and the output predictions In(SOC) were 316 back-transformed into SOC values using exp(In(SOC)).

The optimal number of latent variables of GM_{SIC} , GM_{SOC} and GM_{InSOC} was determined using prediction residual error sum of squares (PRESS) analysis of LOOCV results to avoid under- and over-fitting. Then, all calibration samples were used to build the prediction model with the appropriate number of latent variables, and this model was applied to validation and test sets.

322

323

2.3.4. Local models

A local regression approach was implemented based on PLSR to predict the SOC and SIC content of Tunisian samples. Given a sample *p_i* from *DB_Tunisia* to predict:

The Pearson coefficient of correlation between the spectrum of the Tunisian sample
 p_i and each RMQS calibration spectrum (from *DB_Calib_RMQS*) was calculated;

328 2- The *N* samples from *DB_Calib_RMQS* with spectra that correlated to the spectrum
329 of *p_i* beyond a cut-off value of 0.95 were considered spectral neighbours of the Tunisian
330 sample *p_i*, without maximum limit for *N*.

331 3- A PLSR model was built using the *N* spectral neighbours of the Tunisian sample p_i .

If a Tunisian soil sample had less than 30 spectral neighbours among the
 DB_Calib_RMQS set, this soil sample was not predicted.

334 One local prediction model (denoted LM_{SIC}) was built for SIC prediction based on 335 DB Calib RMQS SIC, validated on DB Valid RMQS SIC and tested on DB Tunisia SIC. One local prediction model (denoted LM_{SOC}) was built for SOC prediction 336 based on DB Calib RMQS SOC, validated on DB Valid RMQS SOC and tested on 337 338 DB Tunisia SOC. Finally, one local prediction model (denoted LM_{InSOC}) was built for SOC 339 prediction based on DB Calib RMQS InSOC, applied to spectra of 340 DB Valid RMQS SOC and DB Tunisia SOC, and the output predictions In(SOC) were 341 back-transformed into SOC values using exp(ln(SOC)).

342 As the calibration sets for SOC and SIC predictions did not include the same samples (DB Calib RMQS SOC and DB Calib RMQS SIC, respectively), the nearest 343 calibration neighbours of a given French validation and Tunisian sample were not the same 344 for SOC and SIC predictions. The optimal number of latent variables was finally 345 346 determined using PRESS analysis of LOOCV on the selected spectral neighbours to avoid under- and over-fitting. Regardless of the type of local model, the spectral outliers were not 347 348 investigated because the selection of nearest neighbours was considered an implicit 349 rejection of outliers.

350

351

2.4 PLSR model evaluation

The performance of global models was evaluated according to figures of merit described in
 Bellon Maurel et al. (2010), from cross-validation, validation and test databases.

The coefficient of determination of cross-validation (R^{2}_{cv}) and root mean square error of cross-validation (*RMSECV*) for *DB_Calib_RMQS* were used. R^{2}_{cv} was computed as 1-ESS/TSS, where ESS is the error sum of squares and TSS the total sum of squares.

The coefficient of determination and root mean square error of prediction for 357 DB Valid RMQS, R²val and RMSEval respectively, were used. R²val was also computed as 358 359 1-ESS/TSS. The ratio of performance to deviation in DB Valid RMQS (RPD_{val}), which is 360 the ratio between the standard deviation in DB_Valid_RMQS and RMSEval, was calculated. The ratio of performance to interquartile range of DB Valid RMQS (RPIQval), 361 362 which is the ratio between interguartile range (difference between the third and first 363 quartiles) of DB Valid RMQS and RMSE_{val}, was also calculated. This parameter has been 364 proposed for variables with non-normal distributions (Bellon-Maurel et al., 2010). And the bias, which is the mean difference between observations and predictions, was calculated 365 366 for DB Valid RMQS (biasval).

367 The coefficient of determination and root mean square error of prediction for DB Tunisia, R²test and RMSEtest respectively, were used. R²test was computed as 1-368 369 ESS/TSS. The ratio of performance to deviation in DB Tunisia (RPD_{test}), which is the ratio between the standard deviation in DB Tunisia and RMSE_{test}, was calculated. The ratio of 370 371 performance to interquartile range of *DB_Tunisia* (*RPIQ_{test}*), which is the ratio between the interguartile range of DB Tunisia and RMSE_{test}, was also calculated. And the bias, which is 372 373 the mean difference between observations and predictions, was calculated for DB Tunisia 374 (biastest).

Finally, a wavelength was considered a significant contributor in a global model when the values of both the regression coefficient and variable importance in the projection (VIP) were sufficiently large: the threshold for the VIP was set to 1 (Chong and Jun, 2005; Wold et al., 1993, 2001), and the thresholds for the regression coefficients were their standard deviations (Viscarra-Rossel et al., 2008).

380 The performances of local models were based on the same figures of merit as 381 those used in global calibration, calculated on *DB_Valid_RMQS and DB_Tunisia*.

382 Concerning the models built from the *DB_Calib_RMQS_InSOC*, independent validation 383 and test statistics were calculated from back-transformed data.

- 384
- 385

387

386 **4. Results**

4.1. Preliminary analysis of soil properties and spectra

388 The RMQS soil sampling covered the French territory, and ranges of SOC and SIC 389 contents in the DB RMQS SOC and DB RMQS SIC, respectively, are large (Table 1). 390 According to the French soil classification, 33 soil reference groups were sampled, with a 391 dominance of Cambisols (IUSS Working Group WRB, 2014; 27% of the sample set), 392 calcareous soils (Calcosols, 22%) and Luvisols (16%). High SIC values are mainly located 393 in three French areas: 1) the southeast (Prealps), mainly with Leptosols and Calcosols, 2) 394 the northeast (chalk Champagne) also mainly with Leptosols, and 3) a transect from west (the Aguitanian Basin) to south (Mediterranean Sea), mainly with Calcosols (Figure 2A1). 395 396 High SOC values are mainly located in 1) mountain areas (Alps in the southeast, Pyrenees in the extreme southwest, Massif Central in the south-centre, Jura in the centre-east), 2) 397 398 cool regions covered by forests and pastures (centre-east), and 3) intensive livestock 399 production areas (northwest; Figure 2B1).

The Tunisian soil samples covered the northern half of Tunisian territory, and the SIC contents range in the *DB_Tunisia_SIC* is as large as the one in the *DB_RMQS_SIC*, whereas the SOC contents range in the *DB_Tunisia_SOC* is lower than the one of the *DB_RMQS_SOC* (Table 1). The sampled Tunisian soils were mainly Calcaric Cambisols and Regosols, Kastanozems, and Chromic and Vertic Cambisols.

- 405
- 406

[Figure 2]

408 Principal component analyses were performed on pre-treated spectra of DB Calib RMQS SIC and DB Calib RMQS SOC, respectively, and pre-treated Tunisian 409 410 spectra were projected onto the plans made by the first and second components. Most 411 Tunisian spectra overlapped a subset of RMQS spectra for SIC (Figure 3a) and for SOC (Figure 3b). So most Tunisian soil samples had similar spectral signatures than a subset of 412 413 RMQS spectra used for calibrating prediction models.

[Figure 3]

- 414
- 415
- 416
- 417

418

4.2. Global models

a. Soil inorganic carbon content

419 For SIC prediction, 52 spectral outliers were identified within the initial calibration dataset, so 1582 RMQS samples were ultimately kept and constituted the DB Calib RMQS SIC 420 421 dataset. The SIC content of these 1582 RMQS samples contained in DB Calib RMQS SIC ranged from 0.0 to 103.9 g kg⁻¹, averaged 6.4 g kg⁻¹, and had a 422 skewness value close to 3.1 (Table 1). The SIC content of the 544 RMQS samples 423 contained in DB Valid RMQS SIC ranged from 0.0 to 95.2 g kg⁻¹, averaged 6.3 g kg⁻¹, 424 425 and also had a skewness value close to 3.1 (Table 1).

The GM_{SIC} was built from the DB Calib RMQS SIC dataset using an optimal 426 number of 15 latent variables, validated on the DB Valid RMQS SIC dataset and then 427 428 tested on the DB Tunisia SIC dataset. The performance of the GM_{SIC} prediction model was accurate, with an R^{2}_{cv} of 0.97 and *RMSECV* of 2.8 g kg⁻¹ in the calibration step (Table 429 2) and an R^{2}_{val} of 0.98 and $RMSE_{val}$ of 2.1 g kg⁻¹ in the validation step (Table 2, Figure 4b). 430 When applied to the DB Tunisia SIC dataset, this GM_{SIC} prediction model provided 431 accurate and unbiased predictions ($R^{2}_{test} = 0.96$, $RMSE_{test} = 5.2$ g kg⁻¹ and $bias_{test} = 0.2$ 432 433 g kg $^{-1}$; Table 3, Figure 4c).

A total of 96 spectral bands might be considered significant based on the analysis of 434 the VIP and regression coefficients of *GM*_{SIC} (Figure 4d). Among the 96 significant spectral 435 bands, those at approximately 2500, 1800 and 860 cm⁻¹ had regression coefficients higher 436 437 than 3 times the standard deviation and therefore might be considered the most significant 438 ones. 439 [Figure 4] 440 441 442 [Table 2] 443 [Table 3] 444 445 b. Soil organic carbon content 446 For SOC prediction, 48 spectral outliers were identified within the initial calibration dataset, 447 448 so 1586 RMQS samples were ultimately kept and constituted the DB Calib RMQS SOC and DB Calib RMQS InSOC datasets. The SOC contents of the 1586 RMQS samples of 449 DB Calib RMQS SOC ranged from 0.6 to 411.3 g kg⁻¹, averaged 25.6 g kg⁻¹, and had a 450 skewness value close to 5.5 (Table 1). The ln(SOC) values of the 1586 RMQS samples of 451 452 DB_Calib_RMQS_InSOC ranged from -0.5 to 6 ln(g kg⁻¹), averaged 3 ln(g kg⁻¹), had a median of 3 ln(g kg⁻¹) and a skewness value close to 0.3. The SOC content of the 544 453 454 RMQS samples contained in DB Valid RMQS SOC ranged from 1.5 to 159 g kg⁻¹, averaged 25.5 g kg⁻¹, and had a skewness value close to 2.5 (Table 1). 455

The GM_{SOC} was built from the $DB_Calib_RMQS_SOC$ dataset using 23 latent variables, validated on the $DB_Valid_RMQS_SOC$ dataset and then tested on the $DB_Tunisia_SOC$ dataset. The performance of the GM_{SOC} prediction model was modest, with an R^2_{cv} of 0.80 and RMSECV of 9.9 g kg⁻¹ in the calibration step (Table 2) and an R^2_{val}

of 0.88 and $RMSE_{val}$ of 7.2 g kg⁻¹ in the validation step (Table 2, Figure 5b). When applied to the Tunisian test set, this GM_{SOC} prediction model provided low accuracy ($R^{2}_{test} = 0.64$, $RMSE_{test} = 16.0$ g kg⁻¹) and biased predictions ($bias_{test} = -5.2$ g kg⁻¹) (Table 3, Figure 5c). A total of 92 spectral bands might be considered significant based on the analysis of

the VIP and regression coefficients of *GM_{SOC}* (Figure 5d). Nevertheless, among these 92
spectral bands, none was associated to very high regression coefficients.

- 466
- 467

[Figure 5]

468

469 A GM_{InSOC} prediction model was built from the DB Calib RMQS InSOC dataset using 10 latent variables and this model was applied to spectra of both DB Valid RMQS SOC and 470 471 DB Tunisia SOC datasets. Finally, the In(SOC) predictions were back-transformed into 472 SOC values for calculating the figures of merit. The performance of the GM_{InSOC} prediction model was accurate, with an R^{2}_{cv} of 0.89 and *RMSECV* of 0.2 g kg⁻¹ in the calibration step 473 474 (Table 2) and an R^{2}_{val} of 0.90 and $RMSE_{val}$ of 6.6 g kg⁻¹ in the validation step (Table 2, 475 Figure 6b). When applied to the Tunisian test set, the GM_{InSOC} prediction model provided high accuracy ($R^{2}_{test} = 0.97$, $RMSE_{test} = 4.2$ g kg⁻¹) and very slightly biased predictions 476 477 $(bias_{test} = 0.7 \text{ g kg}^{-1})$ (Table 3, Figure 6c).

A total of 95 spectral bands might be considered significant based on the analysis of the VIP and regression coefficients in the GM_{InSOC} (Figure 6d). Among these 95 significant spectral bands, those at approximately 2915, and 1800 cm⁻¹ had regression coefficients higher than 3 times the standard deviation and therefore might be considered the most significant ones.

483

- [New Figure 6]
- 485

4.3. Local models

487

a. Soil inorganic carbon content

All validation soil samples had more than 30 spectral neighbours within the $DB_Calib_RMQS_SIC$ dataset, so SIC could be predicted from LM_{SIC} for all samples of $DB_Valid_RMQS_SIC$. The LM_{SIC} provided very accurate and unbiased SIC predictions in validation ($R^2_{val} = 0.99$ and $RMSE_{val} = 1.8$ g kg⁻¹; Table 2). Therefore, this LM_{SIC} provided validation performance slightly better than that of GM_{SIC} (Table 2).

493 All Tunisian soil samples had more than 30 spectral neighbours within the 494 DB Calib RMQS SIC dataset, so SIC could be predicted from LM_{SIC} for all Tunisian 495 samples. The LM_{SIC} provided accurate and slightly biased SIC predictions on Tunisian 496 samples ($R^{2}_{test} = 0.96$, $RMSE_{test} = 5.6$ g kg⁻¹ and $bias_{test} = 1.7$ g kg⁻¹; Table 3, Figure 7a). Therefore, LM_{SIC} provided test performance slightly lower than that of GM_{SIC} , mainly due to 497 498 bias (Table 3). The number of latent variables selected for LM_{SIC} on Tunisian samples 499 varied depending on the sample predicted and followed a relatively normal distribution 500 centred at approximately 13, which was close to the optimal number of latent variables selected by the *GM*_{SIC} (Figure 7c). 501

The number of spectral neighbours of Tunisian samples varied from 65 to 1293 502 503 (Figure 7b). Only a slight trend was observed between the number of neighbours and the 504 prediction error, with a lower error when the number of neighbours increased (Figure 7b). This trend could be expected, as the use of a higher number of neighbour samples for 505 506 calibration should result in more accurate predictions. The spectra from DB Calib RMQS SIC used for building the 96 local individual Tunisian models LM_{SIC} 507 were selected from 0 to 87 times. So no spectrum from DB Calib RMQS SIC was 508 509 systematically selected, whereas only 1.6% of spectra from DB Calib RMQS SIC were never selected. The frequently selected samples were mainly located in SIC-richest areas 510 (Calcosols and Leptosols) such as the southeast (Prealps), the northeast (chalk 511

512 Champagne) and the transect from west (the Aquitanian Basin) to south (Mediterranean 513 Sea) (Figure 2A1 and 2A2).

[Figure 7]

- 514
- 515
- 516
- 517

b. Soil organic carbon content

518 As DB Calib RMQS SOC and DB Calib RMQS InSOC datasets contained same 519 predictors X-variables (MIR spectra), spectral neighbours of validation samples were the 520 same for *LM_{SOC}* and *LM_{InSOC}* models. All validation soil samples had more than 30 spectral 521 neighbours within the DB Calib RMQS SOC dataset and DB Calib RMQS InSOC datasets, so SOC and ln(SOC) could be predicted from LM_{SOC} and LM_{lnSOC} , respectively, 522 for all samples of *DB Valid RMQS SOC*. The *LM_{SOC}* provided accurate and very slightly 523 524 biased SOC predictions in validation ($R^{2}_{val} = 0.93$, $RMSE_{val} = 5.4$ g kg⁻¹ and $bias_{val} = -0.7$ g kg⁻¹; Table 2). Therefore, LM_{SOC} provided validation performance higher than that of 525 GM_{SOC} (Table 2). The LM_{InSOC} provided accurate and unbiased SOC predictions in 526 validation ($R^{2}_{val} = 0.92$, $RMSE_{val} = 5.7$ g kg⁻¹ and $bias_{val} = -0.1$ g kg⁻¹; Table 2). Therefore, 527 LM_{InSOC} provided validation performance higher than that of GM_{SOC} and almost similar to 528 529 that of *LM_{SOC}* (Table 2).

All Tunisian soil samples also had more than 30 spectral neighbours within the 530 DB Calib RMQS SOC and DB Calib RMQS InSOC datasets, so SOC and In(SOC) 531 could be predicted from LMSOC and LMINSOC, respectively, for all Tunisian samples. The 532 *LM_{SOC}* provided accurate and very slightly biased SOC predictions on Tunisian samples 533 $(R^{2}_{test} = 0.89, RMSE_{test} = 6.9 \text{ g kg}^{-1} \text{ and } bias_{test} = 0.5 \text{ g kg}^{-1}$; Table 3, Figure 8a). Therefore, 534 535 this LM_{SOC} provided test performance markedly higher than that of GM_{SOC} (Table 3). The number of latent variables selected by this LM_{SOC} depending on the sample followed a 536 bimodal distribution centred at approximately 16 and 22 (Figure 8c). The second peak of 537

number of latent variables, approximately 22, was close to the number of latent variables
selected by the *GM*_{SOC} (Figure 8c).

The LM_{InSOC} model provided accurate and very slightly biased SOC predictions on Tunisian samples ($R^{2}_{test} = 0.93$, $RMSE_{test} = 5.8$ g kg⁻¹ and $bias_{test} = -0.6$ g kg⁻¹; Table 3, Figure 9a). The LM_{InSOC} provided test performance markedly higher than that of GM_{SOC} but lower than that of GM_{InSOC} (Table 3). The number of latent variables selected by this LM_{InSOC} depending on the sample followed a relatively normal distribution centred at approximately 11 (Figure 9c), which was close to the optimal number of latent variables selected by the GM_{InSOC} .

547 As DB Calib RMQS SOC and DB Calib RMQS InSOC datasets contained same predictors X-variables (MIR spectra), spectral neighbours of Tunisian samples were the 548 549 same for LM_{SOC} and LM_{InSOC} models. The number of spectral neighbours of Tunisian 550 samples varied from 65 to 1292 (Figure 8b and 9b), and no clear trend was observed between the number of neighbours and the prediction error obtained by LM_{SOC} and 551 552 LM_{InSOC} models (Figure 8b and 9b, respectively). As for SIC prediction with LM_{SIC} , the spectra from DB Calib RMQS SOC and DB Calib RMQS InSOC datasets used for 553 building the 96 individual local Tunisian models were selected from 0 to 87 times. So no 554 spectrum from DB Calib RMQS SOC and DB Calib RMQS InSOC datasets was 555 systematically selected, whereas 1.7% of spectra from DB_Calib_RMQS_SOC and 556 DB Calib RMQS InSOC datasets were never selected. The frequently selected samples 557 558 were mainly located in SOC-poor areas (Calcosols and Leptosols) in the northeast (chalk 559 Champagne) and on the transect from west (Aquitanian Basin) to south (Mediterranean 560 Sea) and in soils richer in SOC in the southeast (Prealps) (Figure 2B1 and 2B2).

561

562

[Figure 8]

563

[New Figure 9] 564 565 566 567 5. Discussion 5.1. Global models built on region A for application to region A. 568 569 Before being applied to the Tunisian database, the global models were calibrated with an DB Calib_RMQS_SOC 570 RMQS subset (DB Calib RMQS SIC, and 571 DB Calib RMQS InSOC) and validated on an RMQS subset (DB Valid RMQS SIC and 572 DB Valid RMQS SOC), which means that the global models were calibrated using soil 573 samples collected over a region A to predict values for soil samples collected over this 574 same region A. 575 The validation performance of GM_{SIC} was in accordance with results reported in the literature. Grinand et al. (2012) obtained similar performances using the same RMQS 576 MIRS database, with R^{2}_{val} and RPD_{val} values of 0.97 and 7.6, respectively, when 3/4 of the 577 578 set, selected at random, was used for calibration and 1/4 was used for validation. Barthès et al. (2016) obtained similar performances for SIC using the Tunisian MIRS database for 579 both calibration and prediction, with R^{2}_{cv} and RPD_{cv} values of 0.98 and 7.8, respectively. 580 581 Mc Carty et al. (2002) obtained similar performances using another MIRS database 582 collected in the US, with $R^{2}_{val} = 0.98$ (RPD_{val} was not mentioned and could not be 583 calculated). The most significant spectral bands for GM_{SIC} , located at 2500, 1800 and 860 584 cm⁻¹ (Figure 4d), might be attributed to stretching or bending vibrations in carbonate 585 molecules, as suggested by Du and Zhou (2009) and then by Grinand et al. (2012). 586 The validation performance of the GM_{SOC} was also in accordance with some 587 literature results. Clairotte et al. (2016) obtained very similar performances using the full

588 RMQS MIRS database (including two depth layers, 0-30 and 30-50 cm, instead of one in 589 the present study), with R^{2}_{val} and RPD_{val} values of 0.88 and 2.7, respectively. Barthès et al.

590 (2016) obtained better performances using the Tunisian MIRS database with cross-591 validation, with R^{2}_{cv} and RPD_{cv} values of 0.95 and 4.3, respectively. Moreover, Mc Carty et 592 al. (2002) obtained slightly higher performances using a MIRS database collected in the 593 US for both SOC calibration and validation, with $R^{2}_{val} = 0.94$ (RPD_{val} was not mentioned 594 and could not be calculated).

Following previous researches dealing with non-normal distribution of soil properties (e.g., Waruru et al., 2014; Dangal et al., 2019), natural logarithm transformation was applied to the highly skewed SOC values of the RMQS database to reach a normal distribution in the calibration dataset ($DB_Calib_RMQS_InSOC$). Thanks to this normal distribution, the performance of the GM_{InSOC} on $DB_Valid_RMQS_SOC$ was slightly better than the one of the GM_{SOC} .

Finally, validation performance was higher for SIC with GM_{SIC} (R^{2}_{val} and RPD_{val} values of 0.98 and 7.6, respectively; Table 2) than for SOC prediction with GM_{SOC} (R^{2}_{val} and RPD_{val} values of 0.88 and 2.7, respectively; Table 2) and GM_{InSOC} (R^{2}_{val} and RPD_{val} values of 0.90 and 2.9, respectively; Table 2), confirming that MIRS allows markedly more accurate predictions of SIC than SOC as also shown by McCarty et al. (2002), Grinand et al. (2012) and Barthès et al. (2016).

607

608 5.2. Models built on region A for application to region B.

The models were calibrated by using a RMQS subset ($DB_Calib_RMQS_SIC$, $DB_Calib_RMQS_SOC$ and $DB_Calib_RMQS_InSOC$) and tested on the Tunisian dataset, which means that the models were calibrated by using soil samples collected over a region *A* to predict values for soil samples collected over a region *B*, where *A* and *B* had no common area, so the soil and climate conditions were different between the calibration and test datasets. Our results showed that GM_{SIC} provided accurate test performance (R^{2}_{test} and RPD_{test} values of 0.96 and 4.9, respectively; Table 3), which was however lower

when applied to region *B* than to region *A* (Figure 4c, Tables 2 and 3). The *GM*_{SOC} also provided markedly lower performance (R^{2}_{test} and RPD_{test} values of 0.64 and 1.3, respectively; Table 3) when applied to our region *B* than to our region *A* (Figures 5c, Table 2 and 3).

The RMQS spectra used for SIC and SOC predictions by local models were 620 621 selected using a similarity measure (Pearson's coefficients of correlation), following the 622 same approach than Shenk et al. (1997) and Nocita et al. (2014). So the driver of 623 neighbours selection was the spectral similarity between Tunisian and French spectra. The 624 LM_{SIC} did not improve the SIC prediction accuracy compared to the GM_{SIC} (Table 3). 625 Therefore, *GM_{SIC}* seemed robust and did not need to be adjusted to spectral particularities of region B. So rather than spectral similarity between French and Tunisian samples, the 626 627 main reason for accurate SIC predictions in region *B* seemed to be the strong spectral 628 features of SIC in the MIR region, as suggested by Gogé et al. (2014). The LM_{SOC} improved SOC prediction accuracy compared to the GM_{SOC} (Table 3). Therefore, GM_{SOC} 629 630 seemed poorly robust and the calibration over region A needed to be adjusted to spectral particularities of region B using spectral neighbours (e.g., Shenk et al., 1997; Nocita et al., 631 2014). The increase in the performance of MIRS-based SOC prediction when shifting from 632 633 global to local PLSR is in accordance with literature (e.g. Ramirez-Lopez et al., 2013; Shi et al., 2015; Clairotte et al., 2016 and Dangal et al., 2019). Finally, the frequently selected 634 spectral neighbours of Tunisian samples by the LM_{SOC} were mainly located in SOC-poor 635 areas in the northeast (chalk Champagne) and on the transect from west (Aquitanian 636 Basin) to south (Mediterranean Sea) and in soils richer in SOC in the southeast (Prealps) 637 638 (Figure 2B1 and 2B2). So these frequently selected spectral neighbours of Tunisian 639 samples by SOC local model were not located only over the more similar climatic and 640 pedological contexts such as the Mediterranean context (southeast of France).

642 **5.3. Impact of the SOC In-transformation on models built on region A for application**

643 to region **B**.

The global model calibrated on region A with log-transformed SOC values (GM_{InSOC}) 644 645 provided accurate performance on region B (R^{2}_{test} and $RMSE_{test}$ values of 0.97 and 4.9 g kg⁻¹, respectively; Figure 6c, Table 3). The local model provided better performance when 646 calibrated on region A with log-transformed SOC values (LM_{InSOC}) than with SOC values 647 648 (*LM*_{SOC}), but less accurate than GM_{InSOC} on region B (R^{2}_{test} and $RMSE_{test}$ values of 0.93 649 and 3.6 g kg⁻¹, respectively; Figure 9c, Table 3). So whatever the model using log-650 transformed SOC data in calibration database (GM_{InSOC} or LM_{InSOC}), the SOC predictions 651 on region B were improved compared to models using highly skewed SOC values in 652 calibration database (GM_{SOC} or LM_{SOC} ; Table 3), as showed by Jaconi et al. (2019).

Finally, and unexpectedly, GM_{InSOC} provided better performance than LM_{SOC} when applied on Tunisian samples. So a transformation of calibration SOC values improved SOC model performance more clearly than spectral selection of calibration samples (neighbours). Therefore SOC model performance was more sensitive to the distribution of the explained variable of the calibration samples than to spectral similarity between calibration and test spectra.

659

660 5.4. Perspectives

This study, which used MIRS to predict SOC and SIC contents by using a database collected over a region *A* to predict values over a region *B*, where *A* and *B* have no common area, could be continued with a study to develop predictions based on selected spectral bands. Indeed, spectral band selection remains to be explored, as several studies testing such an approach have obtained different results. Viscarra Rossel and Lark (2009) successfully used wavelets and a variable selection technique to improve SOC calibration using Vis–NIR and MIRS data. Additionally, Volhand et al. (2016) outperformed SOC

predictions based on Vis-NIR spectra by using band selection. However, Stevens et al. 668 (2013) tested recursive feature elimination based on the random forest approach and 669 670 obtained no overall increase in the accuracy of soil property prediction using the LUCAS 671 (European) Vis-NIR soil database compared to models using all spectral bands. In addition, Yang et al. (2019) tested a generic algorithm for spectral band selection but 672 673 obtained no overall increase in SOC prediction accuracy. As previously tested by, e.g., 674 Guerrero et al. (2014) and Guy et al. (2016), spiking could be another useful approach to 675 improve prediction accuracy when applying large-scale calibrations to small regions. 676 Spiking consists of adding a small subset of samples from region B (spiking subset) to the 677 dataset from region A to recalibrate a model.

678 As well, this study could be continued with an impact analysis of the selection of 679 spectral neighbours. Both the number of spectral neighbours and the procedure to select 680 them could be analysed. Several approaches are available for selecting representative calibration samples (Shetty et al., 2012) and could also be tested. For example, to analyse 681 682 the spectral similarity between calibration and test spectra, the Pearson correlation coefficient between spectra could be replaced by the Mahalanobis distance between 683 684 spectra (Nocita et al., 2014) or the Pearson correlation coefficient distance based on Fast Fourier Transform of spectra (Gogé et al., 2012). Finally, some covariates could be added 685 in local regression to improve prediction accuracy, as previously tested by Nocita et al. 686 (2014) who used clay contents of samples as covariates to predict SOC content. 687

688

689

690 **6. Conclusion**

This work highlighted that, as expected, the SOC and SIC contents of French samples were successfully predicted from the French MIRS soil database using a global model based on PLS regression. Predictions of SIC and SOC are accurate when the calibration

694 and validation samples come from same pedologic and climatic contexts. This work also highlighted that when the calibration and validation samples come from different pedologic 695 696 and climatic contexts, the SOC prediction performance over validation samples decreases, 697 whereas the SIC prediction performance remains accurate. Finally, this work showed that 698 prediction models were more sensitive to the distribution of the explained variables of 699 calibration samples than to the spectral similarity between calibration and test spectra. 700 This study confirmed the very high applicability of MIRS for SIC determination and the 701 robustness of SIC prediction models, even when the calibration and validation samples 702 come from different contexts.

- 703
- 704

705 Acknowledgment

706 RMQS soil sampling and physico-chemical analyses were supported by the GIS Sol, which is a scientific group of interest on soils involving the French Ministry for ecology and 707 708 sustainable development, the French Ministry of agriculture, the French National institute for geographical and forest information (IGN), the French Environment and Energy 709 710 Management Agency (ADEME), the French agency for biodiversity, French Institute for 711 Research and Development (IRD) and the French Institute for Agronomic Research (INRA, 712 which is a French public research organisation dedicated to agriculture, food and environment). We thank all the people involved in sampling RMQS sites and in samples 713 714 preparation. D.A. is coordinator of the Research Consortium GLADSOILMAP, supported 715 by LE STUDIUM Loire Valley Institute for Advanced Studies. And we acknowledge the 716 contribution of three anonymous reviewers for their help in improving the manuscript.

- 717
- 718
- 719 **References**

- Apesteguia, M., Plante, A.F., Virto, I., 2018. Methods assessment for organic and
 inorganic carbon quantification in calcareous soils of the Mediterranean region.
 Geoderma Regional 12, 39–48.
- Ahmed, Z.U., Woodbury, P.B., Sanderman, J., Hawke, B., Jauss, V., Solomon, D.,
 Lehmann, J., 2017. Assessing soil carbon vulnerability in the Western USA by
 geospatial modeling of pyrogenic and particulate carbon stocks. Journal of
 Geophysical Research: Biogeosciences 122, 354–369.
- Arrouays, D., Jolivet, C., Boulonne, L., Bodineau, G., Saby, N., Grolleau, E., 2002. A new
 initiative in France: a multi-institutional soil quality monitoring network. Comptes
 Rendus de l'Académie d'Agriculture de France 88, 93–105.
- Barnes, R.J., Dhanoa, M.S., Lister, S.J., 1989. Standard normal variate transformation and
 de-trending of near-infrared diffuse reflectance spectra. Applied Spectroscopy. 43,
 772–777.
- Barthès, B.G., Kouakoua, E., Moulin, P., Hmaidi, K., Gallali, T., Clairotte, M., Bernoux, M.,
 Bourdon, E., Toucet, J., Chevallier, T., 2016. Studying the physical protection of soil
 carbon with quantitative infrared spectroscopy. Journal of Near Infrared Spectroscopy
 24, 199–214.
- Batjes, N. H., 1996. Total carbon and nitrogen in the soils of the world. European Journal
 of Soil Science 47, 151–163.
- Bellon-Maurel, V., Fernandez-Ahumada, E., Palagos, B., Roger, J.M., McBratney, A., 2010.
 Prediction of soil attributes by NIR spectroscopy. A critical review of chemometric
 indicators commonly used for assessing the quality of the prediction. Trends in
 Analytical Chemistry (TRAC) 29(9), 1073–1081.
- Bellon-Maurel, V., McBratney, A.B., 2011. Near-infrared (NIR) and mid-infrared (MIR)
 spectroscopic techniques for assessing the amount of carbon stock in soils Critical
 review and research perspectives. Soil Biology and Biochemistry 43, 1398–1410.

- Bernoux, M., Chevallier, T., 2014. Carbon in Drylands. Multiple Essential Functions. Les
 Dossier Thématiques du CSFD. N°10. CSFD/Agropolis International, Montpellier,
 France.
- Brown, D.J., Shepherd, K.D., Walsh, M.G., Dewayne Mays, M., Reinsch, T.G., 2006.
 Global soil characterization with VNIR diffuse reflectance spectroscopy. Geoderma
 132, 273–290.
- Cécillon, L., Barthès, B.G., Gomez, C., Ertlen, D., Genot, V., Hedde, M., Stevens, A., Brun,
 J.J., 2009. Assessment and monitoring of soil quality using near-infrared reflectance
 spectroscopy (NIRS). European Journal of Soil Science 60(5), 770–784.
- Chen, Y., Barak, P., 1982. Iron nutrition of plants in calcareous soils. Advances in
 Agronomy 35, 217–240.
- Chong, I.G., Jun, C.H., 2005. Performance of some variable selection methods when
 multicollinearity is present. Chemometrics and Intelligent Laboratory Systems 78,
 103–112.
- Clairotte, M., Grinand, C., Kouakoua, E., Thébault, A., Saby, N.P.A., Bernoux, M., Barthès,
 B.G., 2016. National calibration of soil organic carbon concentration using diffuse

infrared reflectance spectroscopy. Geoderma 276, 41–52.

- 763 Comstock, J.P., Sherpa, S.R., Ferguson, R., Bailey, S., Beem-Miller, J.P., Lin, F., Lehmann,
- J., Wolfe, D.W., 2019. Carbonate determination in soils by mid-IR spectroscopy with
 regional and continental scale models. PLoS ONE 14(2): e0210235.
- Dangal, S.R.S., Sanderman, J., Wills, S., Ramirez-Lopez, L., 2019. Accurate and precise
 prediction of soil properties from a large mid-infrared spectral library. Soil Systems 3,
 11.
- Dray, S., Dufour, A.B., 2007. The ade4 package: implementing the duality diagram for
 ecologists. Journal of Statistical Software 22, 1–20.

- Du, C., Zhou, J., 2009. Evaluation of soil fertility using infrared spectroscopy: a review.
 Environmental Chemistry Letters 7, 97–113.
- Gogé, F., Joffre, R., Jolivet, C., Ross, I., Ranjard, L., 2012. Optimization criteria in sample
 selection step of local regression for quantitative analysis of large soil NIRS
 database. Chemometrics and Intelligent Laboratory Systems 110, 168–176.
- Gogé, F., Gomez, C., Jolivet, C., Joffre, R., 2014. Which strategy is best to predict soil
 properties of a local site from a national Vis–NIR dataset? Geoderma 213, 1–9.
- Gregorich E.G., Carter M.R., Angers D.A., Monreal C.M. et Ellert B.H., 1994. Towards a
 minimum data set to assess soil organic matter quality in agricultural soils. Canadian
 Journal of Soil Science 74, 367–385.
- Grinand, C., Barthès, B.G., Brunet, D., Kouakoua, E., Arrouays, D., Jolivet, C., Caria, G.,
 Bernoux, M., 2012. Prediction of soil organic and inorganic carbon contents at a
 national scale (France) using mid-infrared reflectance spectroscopy (MIRS).
 European Journal of Soil Science 63(2), 141–151.
- Guerrero, C., Stenberg, B., Wetterlind, J., Viscarra Rossel, R.A., Maestre, F.T., Mouazen,
 A.M., Zornoza, R., Ruiz-Sinoga, J.D., Kuang, B., 2014. Assessment of soil organic
 carbon at local scale with spiked NIR calibrations: effects of selection and extraweighting on the spiking subset. European Journal of Soil Science 65, 248–263.
- Guy, A.L., Siciliano, S.D., Lamb, E.G., 2015. Spiking regional vis-NIR calibration models
 with local samples to predict soil organic carbon in two High Arcticpolar deserts using
 a vis-NIR probe. Canadian Journal of Soil Science 95, 237–249.
- Hannam, K.D., Kehila, D., Millard, P., Midwood, A.J., Neilson, D., Neilson, G.H., Forge, T.
 A., Nichol, C., Jones, M.D., 2016. Bicarbonates in irrigation water contribute to
 carbonate formation and CO₂ production in orchard soils under drip irrigation,
 Geoderma 266, 120–126.

- Harris, D., Horwath, W.R., van Kessel, C., 2001. Acid fumigation of soils to remove
 carbonates prior to total organic carbon or carbon-13 isotopic analysis. Soil Science
 Society of America Journal 65, 1853–1856.
- ISO (International Organization for Standardisation), 1995a. ISO 10693:1995 –
 Determination of Carbonate Content Volumetric Method. ISO, Geneva.
- ISO (International Organization for Standardisation), 1995b. ISO 10694:1995 Soil Quality
 Determination of Organic and Total Carbon after Dry Combustion (Elementary
 Analysis). ISO, Geneva.
- IUSS Working Group WRB, 2014. International Union of Soil Sciences, Working Group
 World Reference Base for Soil Resources. World Reference Base for Soil Resources
 2014. International Soil Classification System for Naming Soils and Creating Legends
- 807
 of Soil Maps. FAO, Rome.
- Jacobson, M., Charlson, R.J., Rodhe, H., Orians, G.H., 2000. Earth System Science: From
 Biogeochemical Cycles to Global Changes. Academic Press, London.
- Jaconi, A., Poeplau, C., Ramirez-Lopez, L., Van Wesemael, B., Don, A., 2019. Log-ratio
 transformation is the key to determining soil organic carbon fractions with near infrared spectroscopy. European Journal of Soil Science, 70, 127–139.
- Janik, L.J., Skjemstad, J.O., Merry, R.H., 1998. Can mid infrared diffuse reflectance
 analysis replace soil extractions? Australian Journal of Experimental Agriculture 38,
 681–696.
- Jauss, V., Sullivan, P.J., Sanderman, J., Smith, D.B., and Lehmann, J., 2017. Pyrogenic
 carbon distribution in mineral topsoils of the northeastern United States. Geoderma
 296, 69–78.
- Knadel, M., Deng, F., Thomsen, A., Greve, M.H., 2012. Development of a Danish national
 vis-NIR soil spectral library for soil organic carbon determination. In: Minasny, B.,

- Malone, B.P., McBratney, A.B. (Eds.), Digital Soil Assessments and Beyond. CRC
 Press, Boca Raton, FL.
- Lagacherie, P., Baret, F., Feret, J.-B., Madeira Netto, J., Robbez-Masson, J.-M., 2008. Estimation of soil clay and calcium carbonate using laboratory, field and airborne hyperspectral measurements. Remote Sensing of Environment 112(3), 825–835.
- McCrea, J.M., 1950. On the isotopic chemistry of carbonates and a paleotemperature scale. Journal of Chemical Physics 18, 849–857.
- Mark, H.L., Tunnell, D., 1985. Qualitative near-infrared reflectance analysis using
 Mahalanobis distances. Analytical Chemistry 57, 1449–1456.
- McCarty, G.W., Reeves III, J.B., Reeves, V.B., Follett, R.F., Kimble, J.M., 2002. Mid infrared and near-infrared diffuse reflectance spectroscopy for soil carbon
 measurement. Soil Science Society of America Journal 66: 640–646.
- Mevik, B.-H., Wehrens, R., 2007. The pls Package: Principal Component and Partial Least
 Squares Regression in R. Journal of Statistical Software 18, 1–24.
- Nocita, M., Stevens, A., Toth, G., Panagos, P., van Wesemael, B., Montanarella, L., 2014.
- 836 Prediction of soil organic carbon content by diffuse reflectance spectroscopy using a
- 837 local partial least square regression approach. Soil Biology & Biochemistry 68, 337–
 838 347
- Pearson, R.K., 2002. Outliers in process modeling and identification. IEEE Transactions on
 Control Systems Technology 10(1), 55–63.
- Pribyl, D.W., 2010. A critical review of the conventional SOC to SOM conversion factor.
 Geoderma 156, 75–83.
- R Development Core Team, 2012. R: A Language and Environment for Statistical
 Computing. R foundation for Statistical Computing, Vienna. http://www.R-project.org/
 Rabenarivo, M., Chapuis-Lardy, L., Brunet, D., Chotte, J.-L., Rabeharisoa, L.,
 Barthès, B.G., 2013. Comparing near and mid-infrared reflectance spectroscopy for

- determining properties of Malagasy soils, using global or LOCAL calibration. Journal
 of Near Infrared Spectroscopy 21(6): 495–509.
- Ramirez-Lopez, L., Behrens, T., Schmidt, K., Stevens, A., Demattê, J.A.M., Scholten, T.
 2013. The spectrum-based learner: A new local approach for modeling soil vis–NIR

spectra of complex datasets. Geoderma 195–196, 268–279.

- Reeves, D.W., 1997. The role of soil organic matter in maintaining soil quality in
 continuous cropping systems. Soil & Tillage Research 43, 131–167.
- Reeves III, J.B., 2010. Near- versus mid-infrared diffuse reflectance spectroscopy for soil analysis emphasizing carbon and laboratory versus on-site analysis: where are we and what needs to be done? Geoderma 158 (1-2), 3–14.
- Romanyà, J., Rovira, P., 2011. An appraisal of soil organic C content in Mediterranean
 agricultural soils. Soil Use and Management 27, 321–332.
- Scharlemann, J.P., Tanner, E.V., Hiederer, R., Kapos, V., 2014. Global soil carbon:
 understanding and managing the largest terrestrial carbon pool. Carbon
 Management 5, 81–91.
- Shenk, J., Westerhaus, M., Berzaghi, P., 1997. Investigation of a LOCAL calibration
 procedure for near infrared instruments. J. Near Infrared Spectrosc. 5 (1), 223-232.
- Shetty, N., Rinnan, A., Gislum, R., 2012. Selection of representative calibration sample
 sets fornear-infrared reflectance spectroscopy to predict nitrogen concentration in
 grasses. Chemometrics and Intelligent Laboratory Systems 111, 59–65.
- Shi, Z., Ji, W., Viscarra-Rossel, R.A., Chen, S., Zhou, Y., 2015. Prediction of soil organic
 matter using a spatially constrained local partial least squares regression and the
 Chinese vis–NIR spectral library. European Journal of Soil Science 66, 679–687.
- Soriano-Disla, J.M., Janik, L.J., Viscarra Rossel, R.A., MacDonald, L.M., McLaughlin, M.J.,
 2014. The performance of visible, near-, and mid-infrared reflectance spectroscopy

- for prediction of soil physical, chemical, and biological properties. Applied
 Spectroscopy Reviews 49(2), 139–186.
- Stevens, A, Nocita, M, Tóth, G, Montanarella, L, van Wesemael, B., 2013. Prediction of
 soil organic carbon at the European scale by visible and near infrared reflectance
 spectroscopy. PLoS ONE 8(6): e66409.
- Tatzber, M., Mutsch, F., Mentler, A., Leitger, E., Englich, M., Gerzabek, M., 2010.
 Determination of organic and inorganic carbon in forest soil samples by mid-infrared
 spectroscopy and partial least squares regression. Applied Spectroscopy
 64(10):1167–75.
- Tenenhaus, M., 1998. La Régression PLS. Editions Technip, Paris.
- Terra, F.S., Demattê, J.A.M., Viscarra Rossel R.A. (2015). Spectral libraries for quantitative
 analyses of tropical Brazilian soils: Comparing vis–NIR and mid-IR reflectance data.
 Geoderma 255–256 (2015) 81–93
- Viscarra Rossel, R.A., Jeon, Y.S., Odeh, I.O.A., McBratney, A.B., 2008. Using a legacy soil
 sample to develop a mid-IR spectral library. Soil Research 46, 1–16.
- Viscarra Rossel, R.A., Walvoort, D.J.J., Mc Bratney, A.B., Janik, L.J., Skjemstad, J.O.,
 2006. Visible, near-infrared, mid-infrared or combined diffuse reflectance
 spectroscopy for simultaneous assessment of various soil properties. Geoderma 131,
 59–75.
- Viscarra Rossel, R.A., Webster, R., 2012. Predicting soil properties from the Australian soil
 visible-near infrared spectroscopic database. European Journal of Soil Science 63,
 848–860.
- Viscarra Rossel, R.A., Behrens, T., Ben-Dor, E., Brown, D.J., Demattê, J.A.M., Shepherd,
 K.D., Shi, Z., Stenberg, B., Stevens, A., Adamchuk, V., Aïchi, H., Barthès, B.G.,
 Bartholomeus, H.M., Bayer, A.D., Bernoux, M., Böttcher, K., Brodsky, L., Du, C.W.,
 Chappell, A., Fouad, Y., Genot, V., Gomez, C., Grunwald, S., Gubler, A., Guerrero,

C., Hedley, C.B., Knadel, M., Morras, H.J.M., Nocita, M., Ramirez-Lopez, L., Roudier,
P., Campos, E.M. Rufasto, Sanborn, P., Sellitto, V.M., Sudduth, K.A., Rawlins, B.G.,
Walter, C., Winowiecki, L.A., Hong, S.Y., Ji, W., 2016. A global spectral library to
characterize the world's soil. Earth-Science Reviews 155, 198–230.

Viscarra Rossel, R.A., Lark, R.M., 2009, Improved analysis and modelling of soil diffuse
 reflectance spectra using wavelets. European Journal of Soil Science 60, 453–464.

- Vohland, M., Ludwig, M., Harbich, M., Emmerling, C., Thiele-Bruhn, S., 2016. Using
 variable selection and wavelets to exploit the full potential of visible–near infrared
 spectra for predicting soil properties. Journal of Near Infrared Spectroscopy 24, 255–
 269.
- Walkley, A., Black, I.A., 1934. An examination of the Degtjareff method for determining soil
 organic matter, and a proposed modification of the chromic acid titration method. Soil
 Science 37(1), 29–38.
- Wang, X., Wang, J., Zhang, J. (2012) Comparisons of three methods for organic and
 inorganic carbon in calcareous soils of northwestern China. PLoS ONE 7(8): e44334.
- Wijewardane, N.K., Ge, Y., Wills, S., Libohova, Z., 2018. Predicting physical and chemical
 properties of US soils with a mid-infrared reflectance spectral library. Soil Science
 Society of America Journal 82, 722–731
- Williams, P.C., Norris, K.H., 1987. Qualitative applications of near-infrared reflectance
 spectroscopy. In: Williams, P., Norris, K. (Eds.), Near-Infrared Technology in the
 Agricultural and Food Industries. American Association of Cereal Chemists, St. Paul,
 MN, pp. 241–246.
- Wold, S., Johansson, E., Cocchi, M., 1993. PLS partial least squares projections to latent
 structures. In: Kubinyi, H. (Ed.), 3D-QSAR in Drug Design, Theory, Methods, and
 Applications. ESCOM Science Publishers, Leiden, pp. 523–550.

- Wold, S., Sjöström, M., Eriksson, L., 2001. PLS-regression: a basic tool of chemometrics.
 Chemometrics and Intelligent Laboratory Systems 58, 109–130.
- 925 Workman, J. Jr.,, Weyer, L., 2008. Practical Guide to Interpretive Near-Infrared 926 Spectroscopy. CRC Press, Boca Raton, FL.
- Waruru, B.K., Shepherd K.D., Ndegwa G.M., Kamoni P.T., Sila A.M. 2014. Rapid
 estimation of soil engineering properties using diffuse reflectance near infrared
 spectroscopy. Biosystems Engineering, 121, pp. 177-185.
- Yang, M., Xu, D., Chen, S., Li, H., Shi, Z., 2019. Evaluation of machine learning
 approaches to predict soil organic matter and pH using vis-NIR spectra. Sensors 19,
 263.





















Number of latent variables

Table 1: Soil datasets statistics. The SIC values set to zero correspond to values under the laboratory quantification limit (< 0.1 g kg^{-1}).

Dataset	Number of soil samples	Min g kg⁻¹	Max g kg⁻¹	Mean g kg⁻¹	Median g kg⁻¹	SD ª g kg⁻¹	Skewness
DB_RMQS_SOC	2178	0.6	411.3	25.8	19.6	21.8	4.9
DB_RMQS_SIC	2178	0.0	103.9	6.4	0.0	16.1	3.1
DB_Tunisia_SOC	96	2.0	121.0	20.1	14.6	21.0	3.0
DB_Tunisia_SIC	96	0.0	92.9	43.3	48.5	25.6	-0.2
DB_Calib_RMQS_SOC	1586 ^b	0.6	411.3	25.6	19.4	22.3	5.5
DB_Calib_RMQS_SIC	1582°	0.0	103.9	6.4	0	16.1	3.1
DB_Valid_RMQS_SOC	544	1.5	159.0	25.5	19.6	19.5	2.5
DB_Valid_RMQS_SIC	544	0.0	95.2	6.3	0.0	15.7	3.1

^a SD: standard deviation

^b after removing 48 spectral outliers

° after removing 52 spectral outliers

Modele	R² _{cv}	RMSECV	D 2 /	RMSE _{val}	bias _{val}		<i>RPIQ_{val}</i>
Models		(g kg⁻¹)	n val	(g kg⁻¹)	(g kg ⁻¹)		
GMsic	0.97	2.8	0.98	2.1	0.0	7.6	0.5
LMsic	nd	nd	0.99	1.8	0.0	8.8	0.6
GM soc	0.80	9.9	0.88	7.2	-0.4	2.7	2.4
LMsoc	nd	nd	0.93	5.4	-0.7	3.6	3.2
GM insoc	0.89	0.2*	0.90	6.6	-0.1	2.9	2.6
LMInSOC	nd	nd	0.92	5.7	-0.1	3.4	3

Table 2: Figures of merit obtained with global and local models over the French calibration and validation databases.

*RMSEcv calculated on In(SOC)

nd: Not determined.

Table 3: Figures of merit obtained with global and local models over the Tunisian soil samples.

Prediction model	R ² test	RMSE _{test} bias _{test}		RPD _{test}	<i>RPIQ_{test}</i>	
		(g kg ⁻¹)	(g kg⁻¹)			
GMsic	0.96	5.2	0.2	4.9	8.9	
LMsic	0.96	5.6	1.7	4.6	8.3	
GMsoc	0.64	16.0	-5.2	1.3	0.8	
LMsoc	0.89	6.9	0.5	3.0	1.9	
GMinsoc	0.97	4.2	0.7	4.9	3.1	
LMinsoc	0.93	5.8	-0.6	3.6	2.3	