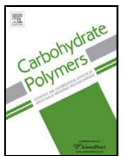
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Title: Qualitative and quantitative assessment of water sorption in natural fibres using ATR-FTIR spectroscopy

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1	Qualitative and quantitative assessment of water sorption in natural fibres
2	using ATR-FTIR spectroscopy
3	
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20 Abstract

21 In the field of composite materials, natural fibres appear to be a viable replacement for glass 22 fibres. However, in humid conditions, strong hydrophilic behavior of such materials can lead 23 to their structural modification. Then, understanding moisture sorption mechanisms in these 24 materials is an important issue for their efficient use. In this work, the water sorption on three 25 natural fibres (flax, hemp and sisal) was studied using Fourier Transformed InfraRed 26 spectroscopy. The spectral informations allowed both qualitative and quantitative analysis of 27 the moisture absorption mechanisms. The main chemical functions involved in the water 28 sorption phenomenon were identified. The absolute water content of the fibres was also 29 determined by using a Partial Least Square Regression (PLS-R) approach. Moreover, typical 30 sorption isotherm curves described by Park model were fitted as well as water diffusion 31 kinetics These last applications confirmed the validity of the FTIR spectra based predictive 32 models.

33 34

Keywords: natural fibres, water sorption, Fourier Transformed InfraRed spectroscopy,
 Partial Least Square Regression, diffusion kinetics

37 38

39 **1. Introduction**

40 Nowadays, natural fibres are considered to be a good alternative for glass fibres replacement

- 41 in the purpose of composite material reinforcement (particularly in automotive manufacturing
- 42 or sport and leisure sector) (Bledzki & Gassan, 1999; Suddell & Evans, 2005). These fibres

43 present a better environmental impact than glass fibres (recyclability, biodegradability) as 44 well as higher specific mechanical properties because of their low density. However, their 45 pronounced hydrophilic behavior -due to their particular structure- leads to high level of 46 moisture absorption in wet environments (Célino, Fréour, Jacquemin, & Casari, 2013). This 47 results in the structural modification of the fibres and an evolution of their mechanical properties together with the composites in which they are fitted in (Dakhal, Zhan, & 48 Richardson, 2007; Placet, Cisse, & Boubakar, 2012; Symington, Banks, David, & Pethrick, 49 50 2009). Thereby, the understanding of these moisture absorption mechanisms is of great 51 interest to get a better control of such new biomaterials.

52

53 Generally, one of the most important factors controlling the water diffusion phenomenon in 54 polymeric materials is the molecular interaction occurring between the diffusing compound 55 and the substrate. The diffusion phenomenon is subjected to the ability of the polymer 56 molecular sites to establish hydrogen bonds with the water molecules. Spectroscopic 57 techniques such as Nuclear Magnetic Resonance (NMR), dielectric or Fourier Transform 58 Infra-Red spectroscopy (FTIR) have been proved to be well adapted to study this 59 phenomenon since they allow to characterize molecular interactions involving potential sorption sites for water (Mijovic & Zhang, 2003; Popineau, Rondeau-Mouro, Suplice-Gaillet 60 61 & Martin, 2005). Among these approaches, FTIR spectroscopy has been widely used to study 62 water transport in polymer and particularly to study the water sorbed into epoxy resins 63 (Cotugno, Larobina, Mensitieri, Musto & Ragosta, 2001; Feng, Berger, & Douglas, 2004; 64 Fieldson & Barbari, 1993; Musto, Ragosta, & Mascia, 2000). Indeed, this technique provides 65 attractive features: i.e. the very high sampling rate, the sensitivity, the accuracy of the 66 quantitative analysis and the informations at the molecular level contained in the vibrational 67 spectra. Moreover, the development of Attenuated Total Reflectance FTIR spectroscopy 68 (ATR-FTIR) encouraged and facilitated the use of this non-invasive technique directly onto 69 solid materials (Chalmers & Dent, 1997).

70

71 On pure cellulosic polymers, the potential sorption sites for water were determined to be 72 hydroxyl and carboxyl groups which are particularly easily detected in FTIR spectroscopy 73 (Berthold, Olsson, & Salmén, 1998). However, few studies have been performed to 74 characterize water sorption by FTIR directly on raw lignocellulosic fibres. (Laity & Hay, 75 2000) demonstrated that it was possible to reproduce water sorption kinetics by recording 76 infrared spectra in reflexion mode on cellophane. More recently, (Olsson & Salmén, 2004) 77 examined the association of water on pulp paper using FTIR spectra acquired in transmission 78 mode. Their results indicated the existence of characteristic bands affected by water. 79 Moreover they determined a linear relationship between the absorbance of those bands and 80 the water content, using a univariate approach. This last work was of particular importance 81 since it highlighted major features helping the understanding of chemical groups involved in 82 water diffusion in pulp paper. However both approaches, based on univariate analysis of 83 FTIR signal failed to properly reproduce the sorption isotherms.

84

The aim of this work is the use of FTIR as an experimental tool to investigate the water sorption onto raw plant fibres, known to be good candidates for the reinforcement of

composite materials (i.e. flax, hemp and sisal). First FTIR spectral signature of each fibre was
investigated in order to describe the molecular effect of the water sorption mechanisms.
Secondly, a multivariate model linking water sorption and the whole FTIR spectra was
developed using partial least square regression (PLS-R). Finally this model was applied for
the accurate monitoring of the water diffusion in the tested biomaterials.

92 93

94 **2. Materials and methods**

95 2.1 Materials

Among the disposable plant fibres, hemp, flax, and sisal fibres were chosen because they presented the best mechanical properties regarding the replacement of glass fibres for the purpose of reinforcing the polymeric matrix (Summerscales, Dyssanayake, Virk & Hall, 2010; Wambua, Ivens, & Verpoest, 2003,). In our study, the bundle of the model fibres was investigated. Bundle of fibres are extracted from the stem (flax and hemp) or the leaf (sisal) of the plant. They are composed of about ten elementary fibres linked together by a pectic cement (i.e. the middle lamella). Their section is in the order of a few millimeters.

103

104 2.2 Moisture sorption protocol

105 First, all thefibre samples tested were dried in desiccators containing silicate gel for 48 hours 106 before controlled moisture sorption treatment. Then, samples were placed in different 107 hygroscopic conditions in a climatic chamber (supplied by Climats and assisted by Spiral 3 108 software) in order to get samples with different water content. The relative humidities tested 109 were 30, 50, 60, 75, 85 and 97 % at room temperature. Experimental sorption isotherms and 110 sorption kinetics were obtained by periodically weighting the samples. Dynamic Vapour 111 Sorption apparatus could also been used to plot sorption isotherms with more accuracy as 112 demonstrated by (Bessadok et al., 2009 or Xie et al., 2011). But in our case, gravimetric measurement provided necessary results. Data were read to 0.01 mg on a precision balance 113 114 (Sartorius – MC1 Analytic AC210P). The equilibrium moisture content Ms is considered to 115 be reached when the mass is stable according to relation (1).

$$M_{W}(\%) = \frac{M(t) - M_{0}}{M_{0}} \times 100 \tag{1}$$

116 where M_0 is the initial weight of the bulk specimens before moisture sorption (dry 117 conditions) and M(t) is the weight of the specimen at time t.

118

119 2.3 Infrared spectroscopic investigations

120 2.3.1 ATR-FTIR spectra acquisition

121 Infrared spectra were recorded in reflection mode directly on the single reflexion diamond

122 crystal of the ATR accessory loaded with bundles of the three fibres presented in section 2.1.

123 The Bruker tensor 27 FTIR spectrometer equipped with the ATR platinum module, with a

- 124 deuterated triglycine sulphate detector RT-DLaTGS and the OPUSv7.0.122 software (Bruker
- 125 Optics, Germany) was set up with the following parameters. The spectral resolution was fixed

ссертер м

- to 1 cm⁻¹, the number of scans to 32, the selected spectral range between 4000 and 400 cm⁻¹. 126
- The penetration depth in the fibres was 0.4 μ m at 4000 cm⁻¹ and 2.7 μ m at 600 cm⁻¹ according 127
- to equation (2) relating penetration depth (dp) to wavelength. 128

$$dp = \frac{\lambda}{2\Pi n_1 \sqrt{\sin^2 \theta - (n_2 / n_1)^2}}$$
(2)

Where λ is the infrared wavelength, n₁, the refractive index of the internal reflexion element 129 (IRE), n_2 the refractive index of the substrate and θ the angle of incidence of the IR beam 130 (Griffith & Haseth, 2007). In our case $n_1 = 2.41$, $n_2 = 1.5$ for the fibres assimilated as polymer, 131 $\theta = 40^{\circ}$, and $\lambda = 4000-600 \text{ cm}^{-1}$). 132

Background spectra were collected using the same instrument settings as those employed for 133 134 the samples and was performed against air. Spectra were recorded for 10 replicates per fibre 135 sample.

136

137 2.3.2 ATR-FTIR spectra acquisition in kinetic mode

138 For the monitoring of water diffusion, time dependent ATR-FTIR spectra were acquired 139 every 2 minutes in ambient conditions during drying of a saturated fibre sample. Each tested 140 fibres were systematically placed in a relative humidity atmosphere of 95 % before the 141 measurement. Recorded spectra were directly treated by Partial Least Square regression 142 (section 2.3.4) in order to determine the water content and to plot the drying kinetics of the 143 tested fibres. Then, the results were compared with the kinetics obtained by the gravimetric measurements. For the gravimetric measurements, the samples were placed on a precision 144 145 balance (0.01 mg) during the drying step and data were collected every 2 minutes.

146

147 2.3.3 ATR-FTIR spectra preprocessing

148 All the spectra were recorded to the background spectra. For the PLS-R model calibration 149 and validation the raw spectra were used since OPUS v 7.0.122 software was set up to 150 manage spectral corrections automatically, according to the value of its optimization criteria. 151 For the Kruskal Wallis analysis the raw spectra were corrected for ambient CO₂ only and 152 smoothed. For the multivariate analysis, the raw spectra were corrected for ambient CO_2 , 153 smoothed and a second derivative calculation was also performed. All these treatments were 154

- achieved using integrated functions of (Bruker Optics, Germany)
- 155 2.3.4 Qualitative Multivariate analysis of ATR-FTIR spectra

156 Pre-processed spectra or pre-processed second derivative spectra were exported as text files 157 for file format modification on Microsoft Excel v14.0.0 or statistical treatment on R 2.15.2. 158 Second derivative spectra were systematically used to improve the infrared band resolution and thus enhance the discrimination of vibrators contributing to the shape of raw FTIR spectra 159 160 (Mecozzi, Pietroletti, & Tornambe, 2011). The spectral data were centered and scaled (mean 161 subtracted and divided by standard deviation) to minimize the unit range influence before 162 performing unsupervised multivariate analysis (Principal Component Analysis (PCA) and 163 Hierarchical Clustering on Principal Components (HC-PC)) using the factominer R package 164 (Lê, Josse & Husson, 2008). The number of classes was determined without a priori by firstly

165 using the unsupervised multivariate analysis (i.e. PCA). A refined assessment of the clusters was then performed using a Partial Least Square Discriminant Analysis (PLS-DA), conducted 166 on all the wave numbers stemming from the ATR-FTIR spectra. This multivariate approach 167 168 was used in order to enhance the formation of clusters by using a model of class prediction 169 (for review see Wold, Sjöström, & Eriksson, 2001). The quality of the prediction was characterized by two parameters, $R^2(Y)$ (goodness of the fit) and Q^2 (predictive capability of 170 the model). The clustering capabilities of the PLS-DA model were considered as satisfactory 171 with both $R^{2}(Y)$ and Q^{2} values superior to 0.5 (Westerhuis et al., 2008). The generation of the 172 173 PLS-DA models was performed in two steps: first the calibration step ($R^2(Y)$ and Q^2), second the validation step named also the cross validation step ($R^2(Y)$)int and Q^2 int). For the cross 174 validation step, (Westerhuis et al., 2008) latest study indicated clearly that it was important to 175 176 notice that even with negative O^2 int values, it was still possible to obtain clear separation between classes. The PLS-DA analysis was conducted using the web server analysis pipeline 177 178 Metaboanalyst 2.0, dedicated to metabolomic data exploitation (Xia, Mandal, Sinenikov, 179 Broadhurst, & Wishart, 2012). For this supervised multivariate analysis ATR-FTIR spectral 180 data were Log10 [1+x] - transformed and Pareto scaled (divided by the square root of the 181 standard deviation) (Van den Berg, Hoefsloot, Westerhuis, Smilde, & Van Der Werf, 2006).

182 To assess the influence of the water content on the ATR-FTIR spectra of the tested fibres, a 183 Kruskal Wallis Test (non parametric ANOVA) was performed per wave number of the ATR-184 FTIR raw spectra recorded on a tested fibre for different water contents. As this non 185 parametric version of the ANOVA does not make any assumption on the nature of the 186 underlying distribution (it does not assume the distribution to be normal), it should be less 187 sensitive to the outliers commonly found in FTIR signatures (since the outliers are taken into 188 account). This allowed a better detection of the vibrators impacted by the water uptake (Di 189 Giambattista et al., 2011). P-value beyond significance threshold (0.05) were plotted on raw 190 spectra data in order to visualize impact of water sorption on ATR-FTIR bands.

191

192

193 2.3.5 Quantitative analysis of ATR-FTIR spectra

There are two steps to obtain PLS-R models: the calibration and the validation. Generally two kinds of validation can be performed: the cross validation and the external validation. Usually the calibration and the cross validation are performed in a single step since identical data set is used to perform calculation.

198

199 (i) Calibration and cross validation

For the calibration (cross validation) step, the procedure of calculation and selection of the best predictive model was iterative and comprised a loop formed of two steps. Firstly the preprocessed infrared spectra were regressed against the reference value (i.e. for a considered fibre: its water content determined gravimetrically). Secondly a full cross validation of the model was performed by omitting one sample and operating the test in the remaining spectral data set. This procedure allowed finding models with high correlations R^2 and low Root Mean Square Error of Cross-Validation (RMSECV). The infrared data sets used in our study

207 comprised 11 samples and 5 replicates per sample, i.e. 55 spectra (ASTM Committee E13,

208 2005).

209 *(ii) External validation*

An independent data set was used for the model evaluation (4 samples and 5 replicates per sample, i.e. 20 spectra) in order to obtain highest correlations (i.e. high R²) and lowest root mean square error of prediction (RMSEP).

213

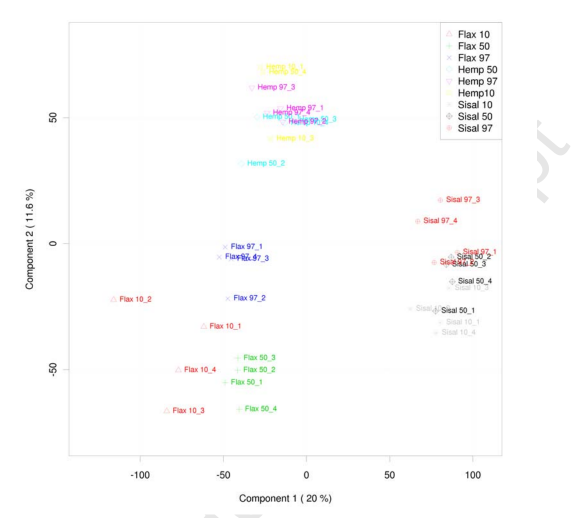
214 **3. Results and discussion**

- 215 3.1 FTIR qualitative analysis of the water uptake effect on the studied fibres
- 216 3.1.1 Each model fibre presented its own FTIR fingerprint

× C

The second derivative spectral data were qualitatively analysed using Partial Least Square Discriminant Analysis (PLS-DA) for each model fibres and for three different relative humidities (RH = 10, 50 and 97 %). On figure 1, results indicate that the three fibres are clustered into three distinct groups, clearly associated to each type of fibre (axis 1). Inside each cluster, it is moreover possible to distinguish subgroups according to relative humidity content (axis 2).

- 223 The clustering results clearly reveal that each type of fibre could be distinguished according to 224 its FTIR spectrum, exhibiting moreover differences for the three tested relative humidities. 225 Those observations suggest that each fibre presents its own FTIR signature, depending on its 226 proper chemical composition. Indeed, according to the work of Satyanarayana and colleagues 227 (Satyanarayana, Arizaga, & Wypych, 2009) this spectral signature could be directly related to 228 the amount of the constitutive macromolecules (cellulose, hemicelluloses, lignin and pectin) 229 and to the crystallization degree of the cellulose which are both fibre dependent. Moreover, 230 as the chemical make-up and the crystallinity degree of the fibres are known to influence the 231 moisture sorption, the PLS-DA results suggest that each fibre should present its own water 232 diffusion behavior. Taken together all those remarks indicate that the water quantification 233 models should be developed per type of fibres.
- 234



235

Figure 1: PLS-DA score plot of second derivative FTIR signatures of the three model fibres
 for three RH conditions.

The legend in the upright corner details the nature of the model fibre and the values of the tested RH % (as suffixes). The classification model used here presented the following characteristics i.e. $R^2(Y) = 0,989$ and $Q^2 = 0,869$ and was validated using 20 permutations $(R^2(Y)int = 0,555 \text{ and } Q^2int = -0,343).$

3.1.2 Water sorption molecular sites could be identified for the model fibres subjected to
increasing RH %

The spectral dependence on the water content is illustrated on figure 2. The development of the sisal FTIR spectra by increasing relative humidity and water content has been treated here solely for reason of clarity. Similar results were observed for hemp and flax fibres (data not shown). The results of the Kruskal Wallis analysis performed on each individual wave number of the raw spectra highlight several zones of the sisal FTIR fingerprint that were strongly impacted by the increasing water uptake. It helps the FTIR spectra bands assignation (using table 1) and allows the interpretation of the experimental observations.

251

Table 1: Assignment of the main absorption bands in FTIR spectra of sisal, flax and hemp fibres. Sisal FTIR data were interpreted according to investigations issued from (Adibi,

254 Cabrales, & Haigler, 2013; De Rosa, Kenny, Puglia, Santulli, & Sarasini, 2010; Liang &
255 Marchessault, 1959; Nelsson & O'Connor, 1964).

Wave number (cm ⁻¹)	Assignment	
3600-3100	Hydrogen bonded of OH stretching in cellulose	
	and/or hemicelluloses	Ι.
2935	CH stretching of cellulose and hemicelluloses	
2862	CH ₂ stretching of cellulose and hemicelluloses	
1735	C=O stretching vibration of carboxylic acid in pectin or ester group in hemicelluloses	
1635	OH bending vibration characteristic of sorbed water	
1595	Aromatic ring in lignin	
1502	Aromatic ring in lignin	
1425	Carboxylic acid of pectin and COO- vibration	
1375	CH bending of cellulose and hemicelluloses	
1335 - 1315	CH ₂ wagging of cellulose and hemicelluloses	
1275	Characteristic peak of lignin	
1240	C-O of acetyl in pectin or hemicelluloses	
1160	anti-symmetrical deformation of the C-O-C band	
1125-895	C-O stretching and ring vibrational modes	
895	Characteristic of β-links in cellulose	
700-650	O-H out of plane bending	

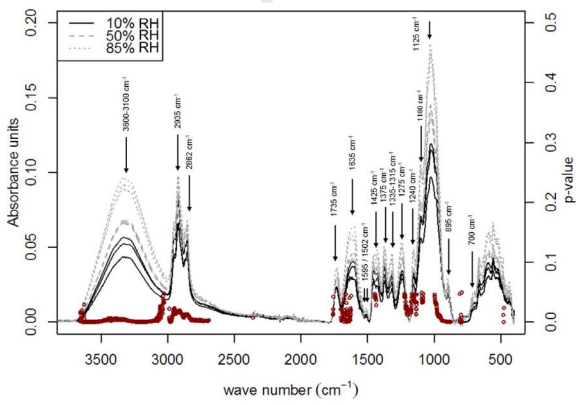


Figure 2: Infrared spectra bands impacted by increasing relative humidity for sisal fibre.

For clarity, only sisal fibre has been presented and only three relative humidities are shown
for three replicates (10 %; 50 % and 85 %). p-values scores, indicating significant impact of
the water uptake on the FTIR bands, were marked using red dots.

264

(i) Part of the water-fibre interactions could be associated to the formation of hydrogen bonds
between the water and the hydroxyl groups of the cellulose or hemicelluloses constitutive of
the raw material.

The broad and unresolved band situated between 3600 and 3000 cm⁻¹ was found to be 268 significantly impacted by the increasing relative humidity content. Potential molecular sites of 269 270 the polymers constituting the studied fibre could be identified here to explain the interactions 271 with the water molecules. Indeed, this band is generally associated to the OH stretching 272 vibrations and hydrogen bonds of hydroxyl groups found in cellulosic materials. However, the 273 absence of clear structured shape makes difficult the assignment of this absorption band 274 (Fengel, 1992; Marchessault, 1962). Recently, (Kondo, 1997) and (Hinterstoisser & Salmèn, 275 1999) helped its resolution by using fitting deconvolution model and DMA-FTIR coupled 276 analysis, respectively. The hydrogen bonds of hydroxyl groups in the 3600-3000 cm⁻¹ wave 277 number range were thus associated to general intramolecular and intermolecular hydrogen 278 bonding and to free hydroxyl in cellulose macromolecule. This peak is also representative of 279 the contribution of the free or the bound water linked to the substrate (Cotugno, Mensitieri, 280 Musto, & Sanguigno, 2005; Murphy & Pinho, 1995; Musto et al., 2000). Here it could be the 281 amorphous phase of the cellulose or the hemicelluloses of the material.

282

(ii) The water uptake could be also characterized by the monitoring of the interaction between the polymers constituting the raw fibre and the free water.

285 The interaction of the cellulose or the hemicelluloses molecular sites of sisal, with the free 286 water is also suggested by the significant impact monitored for the FTIR band situated at 1635 cm⁻¹. Indeed such modification was already observed in the case of water diffusion and 287 288 absorption on cellulosic material, and was found to be characteristic of the free water sorbed 289 on studied material (Laity &Hay, 2000; Murphy & Pinho, 1995). The existence of this 290 interaction should be normally reinforced by the presence of a peak situated at approximately 700 cm⁻¹ and assigned to the out of plane bending of the hydroxyl groups characteristic of the 291 free water molecule. However this last peak is not clearly highlighted by the Kruskal Wallis 292 293 analysis to be dependent of the water content, bringing therefore uncertainty about the 294 influence of free water contribution monitoring. This lack of characterization could be partially explained by the fact that in spectra recorded in attenuated total reflectance, the 295 definition between 700 cm⁻¹ and 400 cm⁻¹ could not be as good as in transmission mode, 296 precluding therefore the exploitation of the spectral information situated in this range of wave 297 298 numbers (Parker & Roy, 1966).

299

300 *(iii) Other molecular sites impacted by the increasing water content could be suggested.*

The FTIR bands situated in the wave number interval ranging from 1100 to 700 cm⁻¹ were also found to be affected by the water uptake. This signature could be associated to the stretching of the C-O-C group of the polysaccharide components of the studied fibres and

304 confirmed the previous observation of Olsson and colleagues (Olsson & Salmén, 2004). The evolution of the peak situated at 1425 cm⁻¹ could also be associated to the interaction of the 305 water to the carboxylic acid moiety of the pectin polymer as firstly depicted by Marchessault 306 307 in his exhaustive study on wood polysaccharides infrared spectra (Marchessault, 1962). 308 Surprisingly, according to the Kruskal Wallis p values assignment, the CH stretching bands situated at 2935 and 2900 cm⁻¹ were found to be significantly impacted by the water uptake. 309 This is difficult to interpret, since no clear chemical explanation could be envisaged. Several 310 311 hypotheses could however bring elements of understanding. It could be inherent to the 312 material which is known to be subject to chemical intrinsic variability or be assigned to non-313 specific experimental variation. Indeed, tests have been performed on different raw fibres for 314 the identical humidities and it effectively highlighted differences in this specific range of 315 wave numbers (data not shown). The way the spectra were processing could also bring 316 elements of explanation. Indeed they were analyzed with almost the same preprocessing 317 treatment than the one used for the PLS-R model generation. In particularly they were not 318 baseline corrected. The shoulder of the OH stretching vibration band clearly shifted up the 319 whole CH bands when the amount of water increases in the sample -whereas no significant 320 increase of the area was monitored (data not shown). This should perturb the Kruskal Wallis 321 test in this particular zone, since it was performed per wave numbers and not per groups of 322 wave numbers -i.e. a band. The combined use of a smoothing function or of moving average 323 performed onto the p values results could enhance the pertinence of the Kruskal Wallis test. 324 These last observations should be carefully considered in the forthcoming multivariate 325 analysis interpretations.

326

327 *3.1.3 Summary*

The results presented above showed that it was possible to discriminate the type of fibre thru its ATR-FTIR second derivative signature using a PLS analysis. Moreover, results revealed that the whole fibre infrared spectra were impacted by the moisture changes. These features strongly suggest that to monitor accurately the water diffusion in the tested biomaterials, a multivariate approach should be used. Indeed, it should improve the water content prediction compared to a univariate approach since the spectral information linked with an increasing water content is visible on the whole spectrum.

335

336 *3.2 FTIR quantitative analysis of the water content of the studied fibres*

337 3.2.1 It was possible to predict the water uptake of every tested fibre by developing material
338 dependent PLS-R models

339 The absolute water content of the tested fibres was determined by using a Partial Least Square Regression (PLS-R) approach. For each kind of fibre, the MIR spectra were used in the range 340 from 4000 cm⁻¹ to 400 cm⁻¹ with the corresponding water uptake measured gravimetrically, to 341 calculate multiple PLS-R models and to perform their validation. The models calculated from 342 the information contained in the spectral regions situated between 3600 cm⁻¹ to 3200 cm⁻¹ and 343 around 1650 cm⁻¹ were selected preferentially since they were associated to the main regions 344 345 characteristics of the interaction between the water and the polymers constituting each 346 studied fibres (3.1). Other regions were not excluded since they could be associated to the

material intrinsic variability (e.g. around 2900 cm⁻¹). The automatic pre-processing 347 treatments, the spectral regions and the quality indicators (i.e. the cross-validation and the test 348 set validation results) used to calculate the PLS-R models are summarized in the table 2. The 349 350 spectral regions selected without a priori are in very good agreement with the observations 351 performed in 3.1, especially for the sisal fibre which presents a wave number range 352 distribution correlated to the three main components known to interact with the water molecules. Moreover, the estimation of the errors associated to the predictive models, 353 354 highlights their pertinence since the RMSECV and RMSEP are very low (\leq to 1). The 355 average number of PLS components is inferior to 7 and the average coefficient of 356 determination greater than 96 %, confirming the quality of the selected models The 357 significance of these parameters has been well described in FitzPatrick and colleagues latest works (FitzPatrick, Champagne & Cunningham, 2012). The figure 3 illustrates the cross-358 359 validation and the test set validation results obtained for the sisal fibre. The dispersion 360 diagrams, displaying the relationships between predictive moisture content obtained by PLS-R quantitative model and gravimetric measurements show a very good correlation ($R^2 > 98$). 361

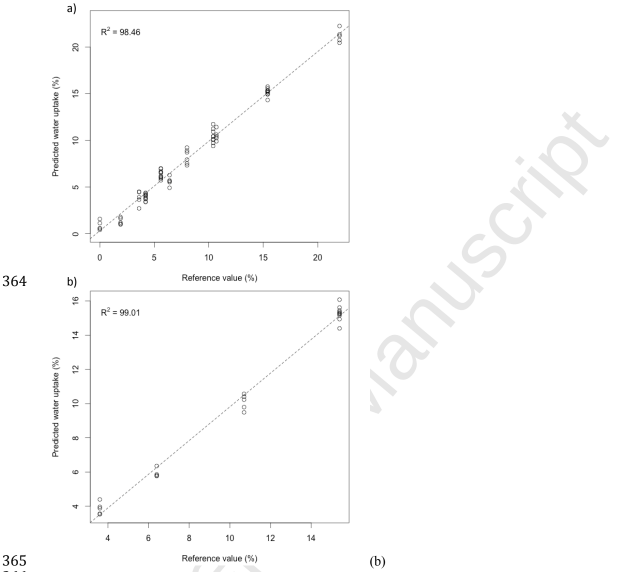


Figure 3: Cross-validation (a) and test set validation (b) results of water uptake prediction
for sisal fibre.

The PLS-R predictions of the water content obtained with the FTIR approach were compared here to the gravimetric measurements. The water uptake is expressed in % of the dry weight of the sisal fibre. The coefficients of determination (%) of the regression lines (indicated in dotted lines) are indicated in the upper left corner.

383 Table 2: PLS-R quality parameters for cross and test set validation for the models selected

384 *for the three studied fibres.*

385 RMSECV stands for Root Mean Square Error of Cross Validation. RMSEP stands for Root

386 *Mean Square Error of Prediction.* R^2 *is the coefficient of determination.*

387

		Hemp	Flax	Sisal
	Pre-processing	First derivative	Min/max normalization	Normalization by vector
	Spectral region (cm ⁻¹)	3998-1119 /760- 400	3998-3277 / 2559- 2198 / 1839-400	3998-3277 / 1839- 1479/ 1120-400
	RMSECV	1.14	1.07	0,72
Cross validation	Number of PLS components	4	9	6
	R ² (%)	95.26	94.92	98.46
	RMSEP	0.620	0.843	0,515
External validation	Number of PLS components	5	7	7
	R ² (%)	97.90	96.65	99.01

388

3.3 The calculated predictive models were applied to quantify the water sorption phenomenonon the three tested fibres

391 *3.3.1 Monitoring of the sorption isotherm using PLS-R models*

392 The estimation of the water uptake of each studied fibre according to its relative humidity 393 exposition was performed using FTIR spectral information and gravimetric measurements as 394 reference method. FTIR data were interpreted in two ways, one using information arising 395 from one band characteristic of the water sorption (univariate approach) and one exploiting 396 the data of the whole FTIR spectra (multivariate approach: PLS-R model). First, the fibre 397 relative water content was estimated to be solely representative of the absorbance variation of the hydroxyl groups situated in the 3600-3000 cm⁻¹ range as already depicted by Olsson & 398 399 Salmén (2004) (3.1.2). Second, the PLS-R models developed for each studied fibre were used 400 to predict the absolute water uptake taking into account the whole FTIR spectral information. 401 The sorption isotherms present a similar shape for the three studied fibres, whether they were 402 depicted by gravimetric measurements or deduced by FTIR data. The figure 4 illustrates the 403 case of the sisal and indicated that the water content is directly related to the relative 404 humidity by following a sigmoidal relation, as already described by (Alix et al., 2009; 405 Gouanvé, Marais, Bessadok, Langevin & Métayer, 2007). That kind of sorption isotherms are 406 in a good agreement with the Park's model (Park, 1986). This model assumes the association 407 of three mechanisms describing the three parts of the curve (figure 4). It is often used to 408 explain the sorption isotherms of hydrophilic and porous media, as cellulosic fibres (Bessadok 409 et al., 2009).

410 The first part of the curve could be related to Langmuir's mode (RH < 10 %, this mode is not 411 really visible on the figure 4). At these relative humidities, water is sorbed onto specific sites

by hydrogen bonding. As discussed in the section 3.1.2 (qualitative analysis of the bands

413 impacted by water) the specific sites could be hydroxyl functions of amorphous cellulose and

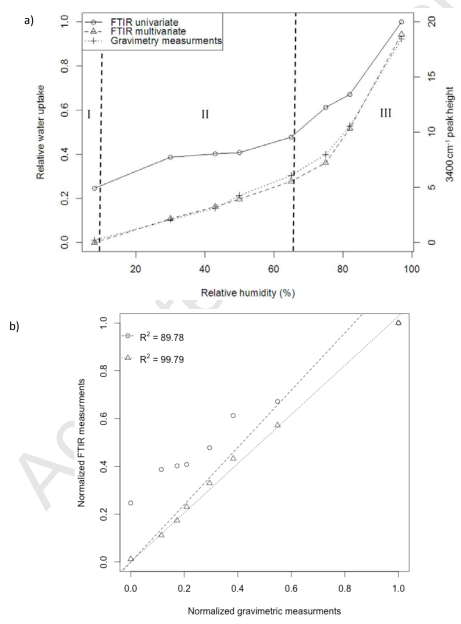
hemicelluloses or carboxylic function of pectin. When relative humidity increases, there is a

saturation of these specific sites of sorption. Then, the water concentration increases linearly with relative humidity as Henry's law describes (until RH = 65 %). This behavior could be explained by the porous structure of fibres where water is free to diffuse. The third part is well described by a power function that represents an aggregation phenomenon of water molecules. Indeed, at high relative humidity, water concentration is too important, and water molecules linked together to form clusters.

421 It is important to notice that the prediction arising from the FTIR PLS-R model fits very

422 well the experimental gravimetric measurements ($R^2 > 99$), highlighting the pertinence of the

- 423 FTIR multivariate approach compared to the FTIR univariate method ($R^2 < 90$).
- 424



425

428 *Figure 4*: Comparison of sorption isotherm for sisal fibre using FTIR and gravimetric
429 approaches. (a) Sorption isotherm plotted for the sisal fibre according to the measurement
430 method (FTIR univariate, FTIR multivariate, gravimetry). (b) Comparison performed between

431 normalized FTIR measurements and normalized gravimetric reference method measurements 432 (FTIR univariate-gravimetry: \bigcirc , FTIR multivariate-gravimetry : \triangle). R^2 stands for coefficient 433 of determination (%).

434

435 3.3.2 Monitoring of the diffusion kinetic using PLS-R models

436 The desorption kinetics of the water sorbed by the studied fibres were also monitored by 437 exploiting the multivariate FTIR approach. The fibre samples were first aged at RH = 95 % 438 until saturation was reached. Second, FTIR spectra were recorded in kinetic mode during their 439 drying in ambient relative humidity conditions i.e. $RH = 45 \pm 3$ %. FTIR spectral information 440 was treated using the developed PLS-R models for each kind of studied fibres and compared 441 to the gravimetric measurements, systematically performed in parallel. For reason of clarity, 442 the desorption kinetics of sisal fibre is solely represented on figure 5a. FTIR data, predict that 443 after moisture sorption in a wet environment at RH = 95 %, the relative mass gain reached 19 444 %. This value was correlated with our gravimetric measurements and previous works realized 445 on similar materials (Gouanvé et al., 2007; Stamboulis, Baillie & Peijs, 2001). The desorption 446 kinetic present a sigmoidal shape that could be interpreted as being the consequence of a 447 delay time in the establishment of a water concentration equilibrium at the surface of the 448 studied fibre (similar behaviour was monitored for the others studied fibres (data not shown). 449 At the end of the experiment the initial weight of the fibre was not recovered. A relative mass 450 loss of about 8 %, compared to the initial weight, was observed. Such mass loss could be 451 attributed to the existence of water content in fibres at ambient relative humidity (Baley, 452 Morvan, & Grohens, 2005).

453

454 Our experimental data could be well fitted by the Carter and Kibler model as described in 455 Célino and colleagues previous works (Célino et al., 2013). In this model, the moisture 456 absorption is described quantitatively by assuming that the absorbed moisture consisted of 457 both a mobile (i.e. free water) and a bound phase (i.e. bound water). Molecules of the mobile 458 phase should diffuse spontaneously with a concentration and stress independent diffusion 459 coefficient D_{γ} , and should be absorbed with a probability per unit time γ at certain molecular 460 sites of the polymer fibre. On the other hand, molecules of the bound phase should be 461 released, becoming thereby mobile, with a probability per unit time β . This assumption was 462 supported by the qualitative FTIR spectral data recorded during the diffusion kinetic (figure 463 5b). Indeed, during the monitoring of the water desorption phenomenon, the main impacted regions (diminishing absorbance) corresponded to the vibration regions of both the free and 464 bound water (3600-3200 cm⁻¹) and the free water uniquely (1635 cm⁻¹). Those experimental 465 466 observations strengthen the assumption of the Carter and Kibler model, suggesting that two 467 kind of water population coexisted in the fibres during absorption or desorption. Water should 468 bind to specific sites as hydroxyl or carboxyl functions and free mobile water could diffuse 469 inside the fibre porous structure.

470

The curve obtained by gravimetric measurement or determined by the PLS-R model based on ATR-FTIR measurement, are in a good agreement. A slight slope difference could be however observed during the transient step. This could be attributed to the fact that the two kinetics were not strictly monitored in identical conditions (section 2.3.2). In gravimetric

measurement, bundle of fibres were free of physical constraints whereas in ATR-FTIR
measurements, fibres were pressed above the measurement crystal. The pressure applied by
the ATR hammer could accelerate the diffusion of the water from the fibre sample, explaining
therefore the observed deviation.

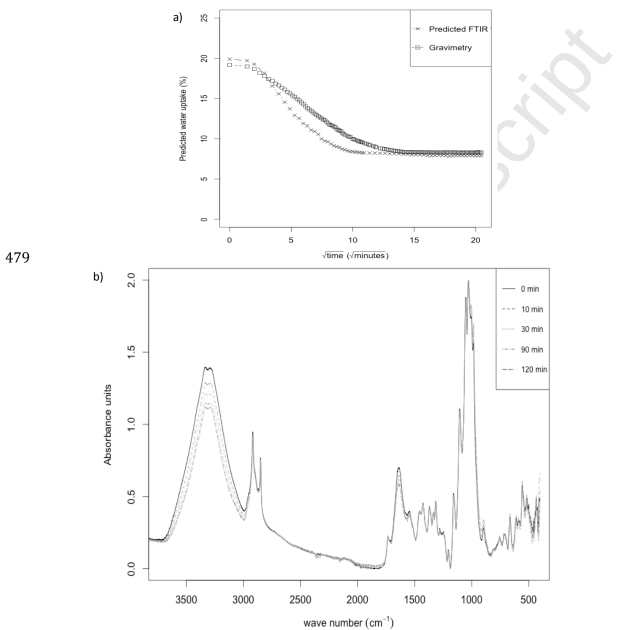




Figure 5: Quantitative and qualitative monitoring of the water diffusion kinetic for the sisal
fibre. (a) Desorption kinetics monitored by FTIR PLS-R prediction and gravimetric
approaches. (b) FTIR spectra recorded during kinetic monitoring for time =0, 10, 30, 90 and
120 min.

485 **4. Conclusion**

486 In the emerging field of composite biomaterials, natural fibres promise an immense potential 487 of application. However, their strong hygroscopic behavior requires the understanding of the 488 moisture sorption mechanisms. Their water content is also of great importance since it could

drive the final properties of the composites where they are fitted in. The aim of this work was
to develop a quick and easy method based on ATR-FTIR spectroscopy to characterize
qualitatively and quantitatively the water sorption phenomenon directly on raw fibres.

492

493 First, our results indicated that it was necessary to develop an approach per fibre since the 494 three tested biomaterials presented their own FTIR signature and therefore their own chemical 495 fingerprint. Hydroxyl and carboxyl molecular sites of the fiber polymer were found to be 496 directly impacted by the water uptake, and confirmed the results observed with less structured 497 cellulosic material (Berthold et al., 1998, Olsson & Salmén, 2004). Second, the FTIR PLS-R 498 models developed for each fibre, in order to quantify their water content, showed very good 499 predictive abilities when compared to the reference method (gravimetric measurements). 500 They were used to monitor accurately sorption isotherms whereas univariate FTIR models 501 could not fit it correctly as depicted by Olsson and colleagues (Olsson & Salmén, 2004). The 502 proposed approach could also supply readily interpretable kinetic data by monitoring 503 continuously the water desorption. The main advantage is the interpretation of kinetics at the 504 molecular level by studying the impacted spectral bands. More detailed analysis 505 (deconvolution of the OH valency band) could lead to the discrimination of the free and the 506 bound water as described in Cotugno and colleagues previous work on an epoxy system 507 (Cotugno et al., 2005).

508

Moreover, our results indicated that the information collected at the surface of the fibre could be efficiently correlated to a global method based on volumetric measurements (i.e. gravimetry). The multivariate approach used to characterize the water sorption dynamically or statically could explain the quality of the results and should encourage the use of such a direct and rapid approach for water sorption mechanism study on other cellulosic biomaterials.

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46, 479-489.

Figure 1: PLS-DA score plot of second derivative FTIR signatures of the three model fibres for three RH conditions. 679 680

The legend in the upright corner details the nature of the model fibre and the values of the tested RH % (as suffixes). The classification model used here presented the following characteristics i.e. $R^2(Y) = 0,989$ and $Q^2 = 0,869$ and was validated using 20 permutations ($R^2(Y)$ int = 0,555 and Q^2 int = -0,343).

685

Figure 2: Infrared spectra bands impacted by increasing relative humidity for sisal fibre.

For clarity, only sisal fibre has been presented and only three relative humidities are shown for three replicates (10 %; 50 % and 85 %). p-values scores, indicating significant impact of the water uptake on the FTIR bands, were marked using red dots.

690

Figure 3: Cross-validation (a) and test set validation (b) results of water uptake prediction forsisal fibre.

The PLS-R predictions of the water content obtained with the FTIR approach were compared here to the gravimetric measurements. The water uptake is expressed in % of the dry weight of the sisal fibre. The coefficients of determination (%) of the regression lines (indicated in dotted lines) are indicated in the upper left corner.

697

Figure 4: Comparison of sorption isotherm for sisal fibre using FTIR and gravimetric approaches. (a) Sorption isotherm plotted for the sisal fibre according to the measurement method (FTIR univariate, FTIR multivariate, gravimetry). (b) Comparison performed between normalized FTIR measurements and normalized gravimetric reference method measurements (FTIR univariate-gravimetry: \circ , FTIR multivariate-gravimetry : \triangle). R² stands for coefficient of determination (%).

704

Figure 5: Quantitative and qualitative monitoring of the water diffusion kinetic for the sisal fibre. (a) Desorption kinetics monitored by FTIR PLS-R prediction and gravimetric approaches. (b) FTIR spectra recorded during kinetic monitoring for time =0, 10, 30, 90 and 120 min.

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Table 1: Assignment of the main absorption bands in FTIR spectra of sisal, flax and hemp
fibres. Sisal FTIR data were interpreted according to investigations issued from (Adibi,
Cabrales, & Haigler, 2013; De Rosa, Kenny, Puglia, Santulli, & Sarasini, 2010; Liang &
Marchessault, 1959; Nelsson & O'Connor, 1964).

715

Table 2: PLS-R quality parameters for cross and test set validation for the models selected forthe three studied fibres.

718 RMSECV stands for Root Mean Square Error of Cross Validation. RMSEP stands for Root

- 719 Mean Square Error of Prediction. R² is the coefficient of determination.
- 720

720	HIGHLIGHTS
721	
722	- Effect of moisture sorption on plant fibres was investigated by FTIR spectroscopy.
723	- Hydroxyl and carboxyl moieties are specific sites for water sorption.
724	- Partial least squares regression quantified the water content of natural fibres.
725	- Models fitted the sorption isotherms and diffusion kinetics.
726	
727	
728	