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1 Evolution of the flax cell wall composition during development

2 and after gravitropism by synchrotron fluorescence imaging

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16 Abstract

- 17 Flax (*Linum usitatissimum*) lodging is an issue of great interest for industrial producers
- 18 due to its economic impact; despite a strong varietal selection over around one century,
- 19 this plant remains sensitive to lodging which represents a main technico-economic issue.
- 20 To better understand lodging effects at the cell wall and stem scale, the cell wall
- 21 composition dynamics during cell wall development and after a 90° tilt bending stress is
- 22 reported. Deep-Ultra Violet fluorescence emission (DUV) dynamics recorded at the
- 23 Synchrotron SOLEIL-DISCO beamline by multichannel autofluorescence imaging is
- 24 addressed for five cellular wall types of flax stems after an artificially induced gravitropic
- 25 reaction. The quantitative fluorescent profile intensities were computed after image

26 analysis, and compared to the control flax stems, we reported a systematically higher 27 average intensity fluorescence (probability >95%) for the 90° tilted plants. Moreover, the 28 average stem fluorescence intensities were significantly different among the 3 29 developmental stages, with the youngest stage (VS) exhibiting on average 30% and 20% 30 less fluorescence than the medium (FG) and mature (M) stages, respectively. The flax 31 stem response to tilt impacted the xylem cellular type, while the bast fibres were arguably 32 less affected by the protein, and hydroxycinnamate contents. A complementary 33 investigation was carried out on bast fibres by infrared microspectroscopy to explore the 34 polysaccharide components not seen in deep UV fluorescence, and significant 35 modifications were monitored. 36 37 **Key words:** Flax; Infrared microspectroscopy; Hydroxycinnamate; Proteins; Lignin; 38 Polysaccharide 39 40 1. Introduction 41 One of the oldest fibre plants used by humans is a flax plant (*Linum Uusitatissimum* L.); 42 traces of flax as far as back the prehistoric civilization in Egypt, ancient Babylonia, and 43 Europe (Heer, 1873) have been verified. Flax, as industrial crop of economic interest, has 44 been used for textile applications, for new developments in the field of high-performance 45 composites in terms of the mechanics (Bourmaud et al., 2018; Mohanty et al., 2018), and 46 for additional functionalities such as acoustic properties, e.g., for violin design (Viala et 47 al., 2018).

Flax belongs to the dicotyledonous group (Ray F. Evert, 2006), and its development has
been well documented and recently reviewed (Goudenhooft et al., 2019a). Basically, this
temperate crop follows four identifiable growth stages and can be schemed on a
cumulative growing degree-day scale, starting from 0°C as the sowing date and reaching
2

52 1,000°C at flax stem fibre maturity. From a description point of view, the beginning is the 53 emergence of the plant, followed by the vegetative stage (VS). At 15–20 days of the VS, 54 the flax plant reaches approximately 15-20 cm (approximately 250 growing degree days). 55 Then, for approximately three weeks, FG takes place. At that time, the flax plant 56 elongates up to several centimetres per day (Gorshkova et al., 1996; Heller et al., 2015), 57 reaching 80–90 cm over this fast FG period (approximately 500 growing degree days). 58 The flax stem reaches a final length of approximately 1 m (Gorshkova et al., 2003) at 59 fibre maturity (M) (approximately 1000 growing degree days).

60 Some external stresses may impact the bast fibre yield and benefits, such as the lodging 61 phenomenon (Goudenhooft et al., 2019b), a stress of economic importance due to its 62 high impact on farmer incomes. From a fundamental view, it is fascinating to note that 63 flax stems can be restored from lodging through a gravitropism reaction under specific 64 conditions, such as the plant response to gravity after tilting (lbragimova et al., 2017). 65 Goudenhooft et al. (Goudenhooft et al., 2019b) reported a decrease into mechanical 66 properties of single fibres between the not-tilted flax fibres and tilted flax fibres, especially 67 for young plants, with almost no difference between the not-tilted /tilted flax fibres at plant 68 maturity. Among the stem cell types, bast fibres are actors in the plant gravitropic reaction 69 through the adjustability of their cell wall performance (Goudenhooft et al., 2019b), but 70 the implication of other cellular types, such as xylem, has also been demonstrated, with a 71 significant evolution of the xylem cell morphology and structure after the tilt stage 72 (Petrova et al., 2021). At the scale of the flax stem, the focus has often been on the bast 73 fibres, which is the reinforcement element of the flax stem (Goudenhooft et al., 2018; 74 Réquilé et al., 2018). In addition, substantial work have been done on the bast fibre from the pulling side of the flax stems tilted at the VS, those cells even displayed an increased 75 76 cell diameter and an enlarged lumen (Ibragimova et al., 2017).

Flax polysaccharides arguably represent the main component of the stem in terms of the
mass, with cellulose being the major component in both xylem (Beaugrand et al., 2014;
Nuez et al., 2020) and bast fibres (Bourmaud et al., 2019). For instance, the stem
epidermis covered by a cuticle, the cambial layer and the primary and secondary xylems
have been less documented (Lion et al., 2017).

82 In the flax stem, some biochemical compounds are naturally fluorescent, including 83 aromatic substances or compounds containing conjugated double bonds, and this 84 property can be used for detection via fluorescence imaging techniques. For example, 85 lignins and hydroxycinnamate exhibit intense fluorescence in the UV and visible regions. 86 In proteins, the dominant fluorophore is the indole group of tryptophan, which absorbs 87 near 280 nm and emits approximately 340 nm. In particular, synchrotron radiation 88 provides deep-UV illumination (DUV), enabling multipurpose molecular fluorescence 89 identification (Giuliani et al., 2009). Using a synchrotron source, a spatial resolution 90 compatible with the sizes of the cell and cell wall $(10-50 \mu m \text{ and } 1-5 \mu m, \text{ respectively})$ 91 can be attained with a good signal-to-noise ratio (Allouche et al., 2012). The excitation at 92 275 nm coupled with an emission in the range 327-353 nm makes it possible to visualize 93 the proteins thanks to the natural fluorescence of tryptophan and tyrosine, which they 94 harbour in their amino acid sequence (Jamme et al., 2014). Using the same excitation 95 wavelength, phenolic compounds exhibit natural fluorescence and can be detected at the 96 cell wall scale in lignocellulosic plants at an emission over 380 nm (Devaux et al., 2018a). 97 Because of fluorescence emission bandpass filters, hydroxycinnamate acids can be 98 partially distinguished from lignin-polymerized phenolic components. Indeed, 99 fluorescence emission bandpass filters at 420-460 nm and 480-550 nm can be used to 100 highlight hydroxycinnamate acids and lignin differentially (Jamme et al., 2013). 101 Since polysaccharides are not autofluorescent, mid-infrared (FTIR) spectroscopy has 102 been used to track polysaccharide compositions. Indeed, FTIR microspectroscopy has

103 proven its value in monitoring hemicellulose metabolism in flax (Chabi et al., 2017).

104 Additional work has highlighted the interest of this approach for flax, examining growth-

105 induced modifications of cells (Stewart et al., 1995). In this study, Stewart et al. reported

106 the spectra of bast fibre cell walls with a predominance of cellulose absorbance. In

107 addition, this work suggested the presence of acetylated hemicelluloses (xylan and/or

108 glucomannan).

109 In addition to the polysaccharidic components, protein and phenolic compounds play also

110 a major role on metabolism and defence mechanisms of flax cell walls. Major

111 achievements have been reported regarding flax protein expression (Corbin et al., 2013),

as well as genome identification and evolution patterns of the main protein family

113 according to differential regulation (Corbin et al., 2018). In addition, phenolic compounds

such as hydroxycinnamates are most often esters linked to polysaccharides and esters or

115 ethers linked to lignin monolignols. These compounds can act as interpolymer coupling

agents, for example, between polysaccharides and lignin or polysaccharides and

117 structural proteins containing tyrosine amino acids. Hydroxycinnamates, such as lignans,

also play an important role in plant defence (Mnich et al., 2020). An attempt to identify

and localize phenolic compounds in cell walls by microscopy was made using a laccase

120 tagged with colloidal gold; this enzyme was able to bind to phenolic compounds

121 (Gorshkova et al., 2000). The authors revealed several gold particle distribution patterns,

122 with compact labelling in xylem and bast fibres; a large proportion of bast fibre wall

123 phenolics consisted of phenolic acids rather than lignin. Lignin is often described as a 3D

124 phenolic polymer (Liao et al., 2020). In flax, lignin is mainly concentrated in xylem, and

the bast fibre lignin content is very low. The spatial regulation of constitutive lignin under

development or during stress is well-documented (Le Roy et al., 2017). The xylem

127 histological part of the stem was particularly well investigated with an elaborate chemistry

128 strategy coupled with a creative microscopic investigation (Lion et al., 2017).

129 The general basics regarding flax plants, their developmental stages and bast fibre

130 characteristics is address in the literature with some recommended papers (Goudenhooft

et al., 2019a; Petrova et al., 2019), but the study of the impact of stress on the cell wall

132 biochemistry or ultrastructure of other flax stem cell types is sparse.

133 In the current study, FTIR microspectroscopy was performed on only bast fibres. This

134 choice is motivated by the economic interest of that fibre and because it is well

135 documented in the literature in comparison to the others type of cell.

136 The purpose of this research is to monitor when, where and with what amplitude the

137 biochemical components in flax varied using FTIR (bast fibre only) and deep UV

138 fluorescence after a gravitropic event. Two conditions were chosen: i) during the normal

139 growth of the flax plant as a control plant and ii) following severe stress, namely, 90°

140 bending, which generates gravitropism. These two growing conditions were combined

141 with the three main stages of flax development (vegetative, fast growth and mature

stage), and for each, the proteins, hydroxycinnamate and lignin distributions and relative

143 amounts were studied for the five stem cell types investigated.

144

145 2. Material and methods

146 <u>2.1 Flax stems, sampling protocol and growing conditions</u>

147 Flax plants (*Linum usitatissimum* L., Bolchoï, seeds provided by Terre de Lin, France)

148 were studied. This variety is this variety is dedicated to the production of fiber fibre. The

149 protocol followed is described in (Goudenhooft et al., 2019b). Briefly, seeds were sown in

150 pots and cultivated outdoors (Lorient, France) in 2017 at a conventional seeding rate of

151 approximately 1800 seeds/m² (Bourmaud et al., 2016).

152 A control (C) batch containing plants that never tilted was used as a reference. Three

additional pots were used for the experiments, consisting of pots inclined at

154 approximately 90° at different key stages of plant development (Fig 1).

155	Figure 1
156	The first pot was tilted at the beginning of the VS, when plants reached approximately 25
157	cm, 29 days after sowing and a cumulative temperature of 370°C. The second pot was
158	inclined during the FG period, when plants were approximately 50 cm, 40 days after
159	sowing and 510°C of cumulated temperature. Finally, the third pot was tilted at plant
160	maturity (M), in the present case when plants were approximately 80 cm, 77 days after
161	sowing and when the cumulative temperature reached the expected 1000°C
162	(Goudenhooft et al., 2017).
163	At each stage of tilting, some plants were sampled. Immediately after sampling, stem
164	portions were fixed overnight in a solution of 4% v/v formaldehyde in 0.1 M phosphate
165	buffer (pH 7.2) at 4°C.
166	
167	2.2 Deep UV Fluorescence imaging at the SOLEIL synchrotron DISCO beamline
168	The DISCO beamline and the imaging microscope setup have been fully described in
169	Giuliani et al.(Giuliani et al., 2009) and in Jamme et al. (Jamme et al., 2013). The so-
170	called TELEMOS microscope is a modified full-field microscope (Axio Observer Z1, Carl
171	Zeiss GmbH, Germany) coupled to the monochromatized synchrotron beam. The power
172	delivered on the sample at the DISCO beamline imaging branch is limited to a few
173	microwatts (1.1*10-6 W at 280 nm). This value avoids beam photodamage by deep UV $$
174	absorption. The synchrotron beam aligned and focused on the sample using an Ultrafluar
175	40x (NA 0.6) objective provides a field of view of 250*250 μ m ² with a pixel size of 244 nm.
176	The microscope is equipped with a 16-bit back-illuminated CCD camera (Pixis BUV,
177	Princeton Instrument, USA) that allows 65536 grey levels to code the fluorescence
178	intensity and an image size of 1024×1024 pixels. The acquisition of images is driven by
179	the ImageJ plugin μManager (Edelstein et al., 2010).

180 The excitation wavelength was set at 275 nm. A dichroic mirror at 300 nm and three 181 bandpass filters were used for the acquisition of multispectral images. The emission filter 182 at 327-353 nm selected the fluorescence of tryptophan and tyrosine found in the proteins, 183 and the emission filter at 420-460 nm made it possible to acquire images of phenolic 184 compounds corresponding to both the hydroxycinnamic acids and lignin. The emission 185 filter at 480-550 nm was retained as a more specific lignin (Frédéric et al., 2013) however 186 more recent investigations has shown that other phenolics compounds like flavonol, 187 condensates tannins, anthocyanins can fluorescence in the same range as lignin do 188 (Vidot et al. 2019). Therefore, the signal collected between 480-550 nm cannot guarantee 189 the solely the lignin emission. Image acquisition was performed at room temperature, 190 and the acquisition time was set to 10 s for the three filters. Finally, visible images in 191 transmission mode were acquired in the sequence of multispectral acquisition. The acquisition time was set to 100 ms for the transmission images. 192

193 The reference images for background and illumination heterogeneities were acquired 194 once per day, and the run of the entire acquisition covered 3 days. The background 195 images corresponded to the signal recorded by the camera without any illumination and 196 any sample. Using the current camera, the background level was approximately 700 and 197 depended on the filter, acquisition time and location in the image. One background image 198 per bandpass filter was acquired by setting the acquisition time accordingly to 10 s. The 199 illumination field depends on the synchrotron beam alignment and the focal plane. The 200 images of the illumination were acquired using a luminescent reference compound (Nd-201 YAG crystal) using only the dichroic mirror without any filter and setting the acquisition 202 time to 250 ms. A Z-stack of 101 images was acquired with a 2 µm step between each 203 image.

Flax stem whole sections with a 45 μm thickness were obtained using a HM650V
vibratome (Microm International GmbH, Waldorf, Germany) and were deposited on a 1"

206 diameter guartz coverslip (R525000, Esco Optics). Just before the acquisition, the 207 sections were mounted in distilled water and sandwiched with a second circular guartz 208 coverslip. A 10x (NA 0.2) Ultrafluar Zeiss (Carl Zeiss GmbH, Germany) lens was used to 209 provide a field of view of 1.116 \times 1.116 μ m² with a pixel size of 1.092 μ m. Stem sections 210 were approximately 3-4 mm large, and a single image was not sufficient to observe the 211 whole section. A series of images were acquired to cover the whole surface of the section 212 after defining a region of interest in the centre of the images that corresponded to the 213 region illuminated by the synchrotron beam. Depending on the section, between 20 and 214 30 individual images were acquired to scan the whole section. 215 216 2.3 Fluorescence image preprocessing and analysis 217 The fluorescence intensity around the stem section was examined for the 5 cell types 218 including the epidermis (ep), the bast fibre (bf), the cambium (ca), the primary (px) or the 219 secondary (sx) xylem shown in Fig 2f.

220

Figure 2

The specific cell type intensity measurements were obtained by manual selection of the pixels (Fig 4B). The fluorescence intensity of the cell types was measured for the three filters as the average intensity of the selected pixels. This measurement was repeated for each of the 4 quarters of the stem.

The fluorescence intensities were averaged over the 2 repetitions of measures in the four quarters of the stem (= 8 measures of fluorescence intensity itself corresponding to the average value along the selected pixels as shown in figure 4 and explained in paragraph 2.3 Fluorescence image preprocessing and analysis). The average values shown in figure 5 and 6 are computed during the variance analysis and show the effect of the two factors development and tilting conditions.

Fluorescence images were preprocessed in Devaux et al. (Devaux et al., 2018b) using
the principles described in Tomazevic work (Tomazevic et al., 2002) for shading
correction. The goal was to remove the camera background and compensate for the
illumination inhomogeneity using eq 1:

235
$$IMC(\lambda) = \frac{(IM(\lambda) - BKG(\lambda))}{ILL(z)}$$
 eq (1)

236 where IM(λ) is the raw image acquired using filter λ , BKG(λ) is the additive background, ILL(z) is the zth image of the illumination reference Z-stack and IMC(λ) is the 237 238 preprocessed image. In the present work, the *z*th illumination image was obtained by 1) 239 calculating an illumination image as the maximum intensity of all individual images of the 240 section and 2) selecting the z^{th} image ILL(z) in the illumination reference Z-stack as the 241 most correlated to the estimated illumination image. The main steps of image 242 preprocessing are illustrated in the Supplementary Data section. 243 Supplementary Data section 244 Images of the whole section were finally built for the three emission filters by merging the

preprocessed images into three mosaic images, called composite images, using the

information provided by the moving stage. The intensities at the stitching interfaces were

set to the maxima between the two adjacent individual images.

248 Fluorescence image analysis included image observation and fluorescence intensity

249 measurement for each given cell type.

250

Table 1

False colour images were built from the three emission filters, as described in Table 1. In the resulting colour images, the regions rich in proteins are identified as mainly blue regions, the regions rich in hydroxycinnamates are identified as green regions, and the regions rich in the assumed lignin signal are identified with a colour varying between

255 yellow and red (see Fig 2).

256 Fluorescence intensities were measured in the images for each filter by manually 257 selecting pixels corresponding to different cell types: epidermis (ep), bast fibres (bf); 258 cambium (ca), secondary xylem (xs) and primary xylem (xp). For each cell type, two sets 259 of pixels were selected from each of the four quarters of the section. The intensities were 260 averaged over pixels, resulting in 5 cell types x 4 guarters x 2 repetitions x 3 filter 261 measures per image. The intensities were compared using variance analysis by 262 considering three experimental factors: development stage, tilting conditions, cell types 263 and their interactions.

Image processing and variance analyses were performed using MATLAB 2019a with the
commercial 'Statistics and Machine Learning' toolbox' (The MathWorks, France).²
Fluorescence profiles were extracted using a home made Matlab software allowing the
multispectral analysis of the images. Variance analysis was performed using Matlab

anova function. The methods allow to recover the mean values and the standarddeviations of the effect tested.

270

271 <u>2.4 Fourier transform infrared microspectroscopy - spectral analysis</u>

Small cylinders (approximately 5 mm in length) of flax stems from the 3 development
stages (see section 2.1 and Fig 1), under normal conditions and under stress with a 90°
tilt, were embedded in paraffin (Automate Tissue Tek VIP 3000). Sixteen thick cross
sections were obtained using an HM355S microtome from Microm and were collected on
ZnS windows. The sections were submitted to a Histochoice clearing agent (Sigma
H2779) solution to remove the paraffin.

278 Localized spectra were acquired by mid-infrared microspectroscopy using a Tensor 27

279 (Bruker Optics) spectrometer equipped with a Hyperion 2000 microscope (Bruker Optics).

Visible images were obtained using a Sony camera (Exwave HAD, SSC-DC80P). A 36x

objective was selected, and depending on the cell sizes and stage of development, an

aperture of 400 to 1600 μ m² was set. Infrared spectra were collected in the 4000-700 cm⁻ ¹ range at a resolution of 8 cm⁻¹. Each spectrum summed 700 scans for the background and 500 scans for the samples.

All spectra were baseline corrected and normalized using the acquisition software

286 OPUS7.5 (Bruker optics). The spectra were cropped to the 2000-700 cm⁻¹ range,

highlighting the polysaccharide region of interest for the bast fibre components. Norris

gap second-derivative spectra were assessed using Unscrambler software (version 10.1,

289 Camo). The second-derivative spectra were thereafter multiplied by (-1).

290

291 **3. Results and discussion**

292 <u>3.1. Qualitative analysis of the fluorescence images according to development stages</u> 293 and tilting conditions

294 Fig 2 shows the fluorescence images of the flax stem cross sections at the young 295 development stage for the control plant for the three filters 327-353 nm (Fig 2a), 420-460 nm (Fig 2b) and 480-550 nm (Fig 2c) corresponding to proteins, hydroxycinnamate and 296 297 lignin, respectively, and the composite view (Fig 2d). One can visually observe that the 298 intensities are different between channels and that the fluorescence signal seems uniform 299 all around the section circumference. The protein fluorescence is mainly located in the 300 secondary xylem. The hydroxycinnamate fluorescence is intense in the cambial tissue 301 and more diffuse in the xylem areas (Fig 2b). However, due to the size resolution of the 302 beam and the very small size of the cells and walls, the signals recovered in the cambium 303 (ca) is likely to mix the fluorescence of the cytosol and the wall. The lignin fluorescence is 304 stronger in the xylem area. The epidermis fluorescence is the most intense in the 305 hydroxycinnamate (Fig 2b) and lignin (Fig 2c) channels, whereas only a slight signal is 306 visible in the protein (Fig 2a) channels. Although we selected carefully bandpass filters to 307 collect the fluorescence emission of the hydroxycinnamate and lignins separately, we

308 cannot exclude a partial overlap of the emission spectra of these compounds, and 309 caution should be taken when assigning the signals in absolute values. Finally, one can 310 notice that in the inner part of the stem, the parenchymatic cells do not fluoresce much 311 regardless of the channel used. The three images were combined to obtain a composite 312 colour image that reveals the distribution of the three fluorescence signals (Fig 2d, 313 composite). The proteins are identified in the blue channel, while the hydroxycinnamates 314 are given by the green channel, and the lignin is depicted by the red channel (Fig 2e). 315 The interpretation of the colours obtained is sometimes tricky and requires the 316 quantification of the individual fluorescence intensity, which will be presented below. A 317 quarter of the stem section was stained with phloroglucinol and zoomed in the picture f 318 (Fig 2f). One can follow from the outer part of the stem to the inward epidermis (ep). 319 Then, the thick bast fibre wall is arranged in bundles (bf), followed by an area of several 320 layers of small and active cells where the cambial area is located. The cambial cells are 321 small and full of cytoplasm, it is expected that the fluorescence intensity signal recorded 322 there will mix the fluorescence of the cytosol and the wall. Further on, we can see the 323 xylem stained in red due to its high content of lignin, the xylem divided into (xs) as the 324 youngest secondary xylem, and finally the oldest xylem as (xp) is the primary xylem build-325 up by the plant.

326

Figure 3

327 Fig 3 shows the composite fluorescence images of the flax stem cross sections using the

328 same spatial scale for different maturity stages and the control versus 90° tilted

329 conditions obtained following the procedure illustrated in Fig 2.

330 Evolution according to the development stage

The first qualitative information is that during the development of the control plant (Fig 3

a-c), a global intensity decrease is observed. The mature stage M_C (Fig 3 c) displays a

lower fluorescence intensity, and the protein signal is almost extinct at this oldest stage.

334 These observations also apply to bast fibres, with some traces of hydroxycinnamate and 335 proteins at the MS. For FG and M stages, the protein signal is intense in the cambial area 336 (Fig 3b) and in an active epithelium if any. The fluorescence intensity of the xylem area 337 increases as the stem matures with a shift from blue green to green-yellow fluorescence. 338 This shift is attributed to a decrease in proteins involved in cell wall metabolism as the 339 stem ages (Fig 3c). The outer xylem part, located close to the cambium, always shows 340 more intense fluorescence. The inner region of xylem shows green-yellow fluorescence. 341 corresponding to the hydroxycinnamates and lignin channels, regardless of the 342 considered developmental stage. Previous work has shown that during the starting stage 343 of lignification, the structure of lignins can evolve; for herbaceous plants such as flax, 344 lignin mainly originates from the polymerization of trans-para-coumaryl alcohol (Day, 345 2004). Lignins can also be linked to cell wall proteins (Monties, 1989) and to phenolics, 346 ferulic or *p*-coumaric acids (Scalbert et al., 1985) (Monties, 1989). Thus, during the 347 implementation of lignin, its structure can take different configurations related to the 348 different fluorescence responses.

349 Evolution related to the tilted effect

350 For the 90° tilted plant, the evolution is not as linear as that during the development stage 351 For the plant inclined at 90°, the evolution is more contrasted than during the development phase of the control plants. VS 90 presented more green fluorescence than 352 353 FG 90 and M 90. More blue fluorescence is seen in FG 90 than in the two other stages. 354 When the gravitropic response takes place during the VS, the cambial region is more 355 intense than the control (see Fig 3a&d), suggesting greater cambial activity metabolism 356 with tilting. In the VS, the main difference is a more intense green ring visible in the 357 innermost stem tissue, corresponding to the primary xylem. One hypothesis to explain 358 this modification could be a consequence of a modification of the transport in the plant, 359 probably of the sap after the curving of the stem, which could induce a mobilization or

360 translocation of hydroxycinnamates. It is generally admitted that primary metabolic

361 activity is reflected by the presence of proteins (Buchanan et al., 2000). These proteins

362 are cell effectors, structural proteins, etc. involved in the major processes of living

363 organisms and cells. Arguably, hydroxycinnamate is a secondary metabolite. Their

- 364 localization and modifications therefore reflect metabolic activity but are not as strong as
- 365 the primary metabolism of proteins.
- 366 The restoration of the plant vertical position also occurred in the FG period (Fig 3 b&e).

367 During this step, the cambial region of tilted samples (Fig 3e) also seemed to be more

368 fluorescent than the control samples (Fig 3. d). The protein signal areas and intensities

369 (blue) were more pronounced, with the bast fibres being greener fluorescent

370 (hydroxycinnamate) than the control plants.

371 During the gravitropic reaction (Fig 3 c&f), a much higher fluorescence intensity was

observed at M_90 than at the control M_C. Despite the presence of well-mature cell

373 walls, especially in the primary tissues, the cambial activity fluorescence of tilted plants

374 (Fig 3f) was still present but more limited than that of tilted plants at the younger stages,

as shown by the reduction in the blue xylem "ring" in contact with the cambium (Fig 3

d&e). Moreover, the area and intensities of the signals coming from hydroxycinnamates

are much greater than those from any other samples, proving that mature cells are still

378 able to react to inclination bending. This reaction most likely affects the lignification

379 process or lignin cellular types such as xylem and hydroxycinnamate, which are

380 precursors of this cell wall component.

381 It seems by eye that mature fibres are not strongly involved in the gravitropic reaction,

382 whereas the xylem secondary part is. In addition, regarding the bast fibre, the signal

383 intensities of M_90 seemed lower than those of tilted plants from previous growth stages

and were similar to those of the control M_C plants.

385	Finally, no polarisation of the signal in response to tilting is analysed 'by eye' observation.				
386	This comment is mandatory because a polar gravitropic response is possible, visible at				
387	the scale of the stem between the opposite and bending sides (Ibragimova et al., 2017).				
388	Indeed, it has been reported that some morphological modifications in response to				
389	gravitropism can take place in the bast fibre but only when bending takes place at a				
390	juvenile plant stage.				
391	This qualitative observation of the signals by eye offers information, but quantitative				
392	information would provide more information. Therefore, we measured the fluorescence				
393	intensity on the cross sections, as explained later below.				
394					
395	3.2. Fluorescence quantitative analysis according to the development stages and tilting				
396	conditions				
397	For a quantitative fluorescence analysis, an arbitrary virtual division of the stem section				
398	into four quarters, called 1 to 4, was performed, as illustrated in Fig 4 A.				
399	Figure 4				
400	The fluorescence intensity collected for each filter allows quantitative evaluations in the				
401	fluorescence count unit.				
402	From those average fluorescence intensity values, an analysis of variance (ANOVA) was				
403	performed for each filter (Fig 5).				
404	Figure 5				
405	The developmental stage, tilting, cell types and all interactions were found to be				
406	significant. Because proteins, hydroxycinnamates and lignin arguably do not fluoresce in				
407	the same manner and in the same environment, a higher fluorescence does not				
408	necessarily mean higher molar amounts; therefore, it is not possible to compare the				
409	different components as molar amounts based on their respective signals. If we focus first				
410	on the development stage effect, then the average fluorescence intensity for the three 16				

411 emission channels corresponding to proteins (Fig 5a), hydroxycinnamates (Fig 5b) and 412 lignin (Fig 5c) varied between the 3 development stages. The highest intensity for 413 proteins and hydroxycinnamates was observed in the fast growing stage for both the 414 control and 90° tilted stems (FS C or FS 90), and the lowest intensity was observed in 415 the vegetative stage (VS C or VS 90°). This trend is different for lignin: an increase in 416 the intensity occurred between stages VS and FG before reaching a plateau. In contrast, 417 the lignin fluorescence seemed equal for the three developmental stages of the 90° tilted 418 stems. If we focus now on the tilt effect, then the gravitropic response systematically 419 induced an increase in the fluorescence intensity in the three channels compared to the 420 control plant stems, by 51, 28 and 23% for proteins, hydroxycinnamate and lignin, 421 respectively. The 90° tilt gives rise to an average increase of 32% of the fluorescence 422 (Fig 5), somehow confirming the qualitative observation (section 3.1, Fig 2). This result 423 suggests a global boost in plant metabolism following lodging stress. 424 425 3.2.1 At the cell type scale, evolution according to development stage 426 To go further in the analysis, we looked at the fluorescence intensity according to the cell 427 types. 428 Figure 6 429 Fig 6 presents the average fluorescence intensity for the 5 cell types related to their 430 developmental stage and tilt condition. The epidermal protein intensity signal was weak 431 and did not change according to the VS and the FG and M stages of the control plants 432 (Fig 6). The hydroxycinnamate fluorescence signal is more intense and remains stable. 433 The highest fluorescence intensity was measured for lignin, but the standard deviation of 434 the average values was so large that no significant trend could be deduced. Here, the 435 very thin layer of the epidermis is arguably a technical limitation with respect to the pixel

resolution, which might explain in part the importance of the standard deviation (SD)observed.

438 Bast fibres are thicker, and the SDs are very small regardless of the constituent under 439 consideration (Fig 6, second line of histogram). The intensity of protein fluorescence 440 increased between the VS and FG stage and then decreased between the FG and M 441 stages in the control plants. The intensity of the hydroxycinnamate fluorescence 442 remained almost unchanged between the VS and FG stage and then decreased at the M 443 stage. The lignin signal evolves accordingly, with only a decrease at stage M. Scarce 444 literature deals with the quantification or localization of lignin in bast fibres, and a recent 445 report based on safranin ratiometric image investigations on flax stem cross sections has 446 presented a slight increase in the lignin content of mutant plants during cell development 447 (Baldacci-Cresp et al., 2020).

448 Due to the economic importance of flax bast fibres, more literature is available regarding 449 their composition (Akin, 2013; Bourmaud et al., 2018; Bourmaud et al., 2019) and 450 evolution during plant growth and in response to stress (Goudenhooft et al., 2019a). In 451 connection with Fig 3, the colour of the bast fibre evolves with maturity; in the FG stage, 452 the blue colour representing mainly proteins is evident with a homogenous colour within 453 the fibre cell wall, indicating probably the achievement of the transformation of the 454 immature Gn wall into the mature G at this stage (Gorshkova et al., 2003). This 455 gualitative observation is supported by the protein fluorescence peak observed in Fig 6 456 (see FG-C, 'Bast Fibre Proteins'). The structure of bast fibres has been demonstrated to 457 be progressive, with a first Gn layer composed of cellulose and long galactan chains 458 evolving towards a more structured G layer. In this new and mature layer, the length of 459 galactan chains is reduced under the action of galactosidase enzymes, inducing their 460 entrapment between cellulose microfibrils (Gorshkova et al., 2015) and a more cohesive 461 and stiff (Arnould et al., 2017) structure. Here, protein family identification is not possible

since we indifferently detected all tryptophan that belongs to the whole protein class.

463 However, an increase in AGP and extensin during flax fibre development is more likely

464 (Day et al., 2013).

465 The cambium fluorescence signals (third line of histogram, Fig 6) for proteins,

466 hydroxycinnamates and lignin show similar trends to the bast fibre, i.e., an increase

467 between the VS and FG stage, followed by a slight decrease in the M stage. The

468 fluorescence intensity in cambium is rather high, and the variations between

469 developmental stages are marked.

470 The secondary xylem (xs) cells demark with the other cell wall types. Indeed, for 471 fluorescence signals attributed to hydroxycinnamates or lignin, there was a significant 472 decrease from the VS to the M stage. The protein fluorescence is higher than that of the 473 other cell types, and signals for proteins fall drastically between the FG and M stages. In 474 contrast, for the signals attributed to hydroxycinnamates and lignins, the decrease is 475 progressive between the VG and M stages. Regarding lignin, a recent report using an 476 elegant tag of lignin specifies the active areas of lignification in the secondary xylem of 477 flax (Lion et al., 2017). The authors showed temporal lignification differences among the 478 xylem vessel, ray parenchyma cells and tracheid cells, as well as revealing the 479 polarization at the scale of a single-cell lignification dynamics. In this work, such 480 resolutions could not be achieved, but we observed a slight decline in the fluorescence 481 intensity in stages FG and M compared to the VS (Fig 6). 482 Finally, the cell walls of the xylem primary seem to display a unique dynamic over 483 developmental stages, as no significant variations could be detected from the VS to the M

stage, regardless of the three biochemical components targeted (Fig 6).

485

484

486 *3.2.2* At the scale of the cellular type, evolution according to the 90° tilt stress

The epidermis (ep) fluorescence signals of the proteins, hydroxycinnamates and lignin did not seem to change in stems after 90° tilt stress (Fig 6). The gravitropic response may also be difficult to detect in this thin layer due to an important SD that could mask a moderate physiological plant response.

For the cambial area, all 3 biochemical families exhibited a stronger intensity after tilt,
regardless of the maturity stage considered. This finding highlighted the impact of
gravitropism on the cambial response and the preponderant function of this tissue in the
plant reaction.

The xylem secondary cell wall type is affected by the gravitropic response that follows the 90° tilt (Fig 6). Regarding the protein, signal intensities Indeed, the intensities in FG and M stages were significantly higher than those in the controls. The hydroxycinnamate signals were more intense in the VS and the FG and M stages than in the control stems. The lignin signal follows the same trend but with a smaller contrast between controlled and tilted plants.

The xylem primary cell wall type revealed only an increase in the protein signal at the FG stage. In contrast, for hydroxycinnamate, the FG stage is the only stage unaffected by the 90° tilt stress. Compared to control stems, the lignin signal was higher in both the VS and the M stage, but the FG stage was unaltered.

Finally, the bast fibre after 90° tilt could be expected to be involved in the response of the plant. Tilting is known to disrupt normal fibre thickening and lignification in tension wood fibre walls (Ghislain and Clair, 2017). Here, the fluorescence intensity recorded for proteins does not support a modification of the protein within the cell wall if tilt occurs in the VS or FG stage. The protein signals were unchanged between the control and tilted plants, in accordance with both our visual observations (Fig 3) and with the findings of Goudenhooft et al. (Goudenhooft et al., 2018). However, a slight protein signal increase

- 512 was observed if tilt occurred in the M stage. Tilting induces an increase in the
- 513 hydroxycinnamate signal, and the highest difference between tilting and control plants
- 514 was observed during the FG stage. For lignin, all tilted plants harbour a slight increase in
- the fluorescence compared to the control plant, and lignin is a well-known polymer
- 516 involved in plant stress mechanisms.

517 <u>3.3 FTIR on the bast fibre</u>

- 518 To supplement the deep UV fluorescence examination of the 5 cell types, a study was
- 519 performed by FTIR microspectroscopy to investigate the polysaccharide composition; this
- 520 method has been successfully performed on wood fibres (Gierlinger et al., 2008) and flax
- 521 mutant plants (Dymińska et al., 2014). We focused on the most targeted cell type of the
- 522 flax stems in the literature, namely, the bast fibres.

523 FTIR and the development effect on bast fibre

524

Figure 7

- 525 A micro-FTIR investigation was performed on the bast fibre cellular type.
- 526 Fig 7A illustrates the location of micro-FTIR spectra in a flax stem section. An average
- 527 spectrum was calculated, and the assignment of the main absorption bands to
- 528 biochemical compounds is presented in Fig 7B. The main mass at 1200-700 cm⁻¹ is
- 529 assigned to polysaccharides, such as cellulose, and heteropolysaccharides, such as
- 530 hemicelluloses and pectin.
- 531 The band at approximately 1460 cm⁻¹ is assigned to the CH₂ bending of lipids when
- 532 associated with the C-H₂ stretching bands at approximately 2850 and 2920 cm⁻¹ (not
- 533 shown). The bands at 1735 and 1245 cm⁻¹ correspond to esters, while the absorption
- 534 bands at approximately 1640 and 1540 cm⁻¹ are signatures of the presence of proteins.
- 535 The 1640 cm⁻¹ band also refers to fibre internal water.
- 536

Figure 8

537 The second-derivative spectra were processed to increase spectral differences in the 1800-700 cm⁻¹ region (Fig 8). Cellulose, the major constituent of flax fibres, is identified by 538 539 the spectral bands at 898, 995, 1010, 1030, 1060, 1105 and 1160 cm⁻¹ (Maréchal and 540 Chanzy, 2000). The hemicelluloses, O-acetylated glucomannan, xylan and xyloglucan 541 are also present in flax fibre absorbed in the same spectral region as cellulose but can be 542 revealed by the shoulder at approximately 940 and 815 cm⁻¹ (Himmelsbach et al., 1998). 543 The youngest development stage (Fig 8) differs from the later stages by bands assigned 544 to lipids (1735 and 1460 cm⁻¹) and adsorbed water (1640 cm⁻¹). The peak at 1010 cm⁻¹ is 545 associated with two shoulders at approximately 1600 and 1430 cm⁻¹, assigned to 546 carboxylate symmetric and antisymmetric COO⁻ bands, which correspond to residual 547 pectin. Its methylated or acetylated form is also present by bands at 1735 and 1245 cm⁻¹. 548 The variation in band intensities assigned to cellulose and hemicelluloses at 1010 and 549 1050 cm⁻¹ mainly arises from the change in the pectins/cellulose or cellulose 550 conformation ratio. Regarding the non-cellulosic polysaccharides, pectic galactan, as well 551 as their processing enzymatic machinery (Goubet and Morvan, 1993), are known to be 552 part of the bast fibre and evolve according to cell wall modelling (Mikshina et al., 2012), 553 while galactan and and proteins capable of modifying it play a role with cellulose within 554 Gn to G development (Gorshkova et al., 2004; Roach et al., 2011). 555 FTIR and tilt effect on the bast fibre 556 Figure 9 557 When comparing FTIR signals from the control bast fibre and after tilting (Fig 9), the 558 spectra of the youngest VS bast fibre from 90° tilted plants presented absorption bands

from 1105-1050 cm⁻¹ with a higher intensity (Fig 9A). This result suggests an increase in

the cellulose and hemicellulose contents for the bast fibre of tilted plants. Comparable

trends have already been reported with some genetically modified flax lines, with

562 modification of the cellulose ultrastructure and allomorphs in particular (Dymińska et al.,

563 2014). For the FG stage, only slight differences were observed between the control and 564 the 90° tilted plant fibres, corresponding to variations in the absorbance intensity bands 565 assigned to cellulose and hemicelluloses (995, 1030, 1050 and 1060 cm⁻¹) (Fig 9B). At 566 the MS, most of the signals of the 90° tilted FG spectra are close to those obtained for 567 control plants (Fig 9C), but the two bands drastically differ at 1105 and 1050 cm⁻¹, 568 indicating that there is also a change in cellulose and a broader panel of polysaccharides 569

570

between these two conditions.

571 <u>3.4 Seeking a stem polar effect on bast fibres by quantitative analysis of the fluorescence</u> 572 To confirm the gualitative conclusion of the absence of polarity between the stem sides 573 (Fig 3), we quantified the fluorescence according to the localization of the pixel selection 574 in the whole circumference of the stem, with a more continuous position resolution (Fig. 575 10). For that purpose, in the stem sections of control (M C) and tilted plants (M 90), we 576 measured the fluorescence from an initial '0' point arbitrarily located.

577

Figure 10

578 Then, we performed a full walk on bast fibres where pixels were selected at different 579 positions all around the stem circumference (Fig 10 a), with at least one set of measurements every 30°. No significant variations in any of the three channel 580 581 fluorescence intensities can be observed, as seen in the average value linearized 582 according to the angle of measurement (Fig 10 c). Indeed, although an up and down 583 trend can be observed for each of the fluorescence channels, there is no trend for a 584 higher or upper side of the stem with more proteins, hydroxycinnamates or lignins. The 585 tilted stem section showed the highest fluorescence intensity, confirming the 586 guantification of the bast fibre in Fig 6 and the ANOVA (Fig 5). Under our experimental 587 conditions, the 10 cm stem samples collected (starting from 2 cm above the cotyledons), which correspond to the lasting curvature zone when recovery takes place. A comparison 588

589 with side effects reported in the bast fibre shows an induced morphological change on the 590 bast fibre on the pulling stem side compared to the opposite stem side and control plants, 591 and only if the tilt was at the stage of the G-layer deposition (middle and lower part of stem) (Ibragimova et al., 2017). We hypothesize that this effect is not observed in our 592 593 report because we may not have sampled the same stem section; for instance, 10 cm 594 long samples were taken 2 cm after the cotyledon versus sampling 1 cm after the 595 cotyledon (Ibragimova et al., 2017). In other words, we may have bypassed the area where this polarity occurred, probably by sampling slightly too far in the stem, even if the 596 597 stem sampling area was slightly curved.

598

599 4. Conclusion

600 The aim of this work was to gain new insights into the dynamics of the cell wall composition in the different cell types of the flax stem during plant development and after 601 602 a 90° tilt bending stress. The autofluorescence properties of proteins and phenolic 603 compounds when excited at a deep UV (DUV) wavelength accessible to the SOLEIL 604 synchrotron and multichannel fluorescence imaging have made it-possible to track 605 proteins and hydroxycinnamates. At contrary, whereas the quantification of the lignin fluorescence by the 480-550 nm filter could not be attained is our case. The specificity of 606 607 this filter was checked on extracted polymers. In our work, the higher amount of 608 fluorescence observed using this filter in both epidermis or cambium compared to xylem 609 cannot be assigned to lignin. We hypothesis that non-polymeric molecules, unidentified in 610 this work, overlap with the fluorescence of lignin. Further investigation are needs to 611 elucidate whose molecules are concerned. The main output is that there are higher 612 fluorescence signals of the hydroxycinnamate and both 'lignin' and proteins at the FG 613 stage for the five cell types of the stems, without exception. The fluorescence intensities 614 decrease at the M stage.

After a 90° tilt of the flax stem, generally, the fluorescence intensity was higher in all five cell types. This molecular investigation shows that full histological flax stem organization is involved in the gravitropic response following tilting. Our data support the fact that the gravitropic response is also dependent on the metabolic activity of the plant, and the variations in proteins, hydroxycinnamates and lignin are influenced by the developmental stage when tilt occurs.

621 For cambium as well as the secondary xylem, the active cells are strongly affected by 622 tilting, as demonstrated by a strong increase in the fluorescence intensity reflecting an 623 increase of the metabolic activity. No cambial activity in the sense of cell divisions was 624 measured in this work. So, in the sense of metabolic activity, if the gravitropic reaction 625 occurs at plant maturity, cambial overfluorescence intensity in tilted plants is still present 626 but is moderate compared to reactions occurring at vegetative or maturity stages. 627 The bast fibres were proven to also react after the stem tilt but moderately at the mature 628 plant stage. No evidence of stem polarity generated after tilt was demonstrated by

629 synchrotron DUV fluorescence imaging of the bast fibres, regardless of the

630 developmental stage examined.

631

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- 803

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805 Figures caption

- Figure 1. Scheme of the different steps of the experiment. The top part: a) Schematic
- 807 representation of the flax stems at different maturity stages. The pots were never inclined
- and so acted as control (_*C*) conditions, whereas other pots were tilted for 4 days. Tilting
- 809 (_90) was performed at three major stages of plant development (VS, FG and M), and
- tilted samples were collected 4 days after tilting, as depicted in the lower part b). About
- 811 10 cm stem portions taken approximately 2 cm above cotyledons were sampled from
- 812 whole stems and are illustrated by blue rectangles in b). Overall, 6 flax stem conditions
- 813 were explored (*VS_C*, *VS_90*, *FG_C*, *FG_90*, *M_C* and *M_90*).
- 814 **Figure 2.** Individual channel fluorescence and composite view of the VS_C flax stem
- 815 cross section. a) Fluorescence intensity of the channel of proteins, b) channel of
- 816 hydroxycinnamate in c) channel of lignin, d) the composite view with the merged a-c
- 817 channels, in e) a guide to interpret colours and in f) a phloroglucinol staining of the
- 818 quarter of the stem section to present the 5 cellular components of this study. The field of
- 819 view is 2.7 x 2.5 mm² for images a to d. In f, the inner stem part to the epidermis is (ep),

the bast fibre (bf), the cambium (ca), the xylem - secondary (xs) and the xylem - primary(xp).

Figure 3. Composite fluorescence images of the control plants _*C* or tilted plants _*90*, magnification 10x. Stem cross section from a control plant (a) and from a tilted plant (d) during the vegetative stage VS. Stem cross section from control (b) and tilted (e) plants during fast growth *FG*. Stem cross section of mature *M* plants and control (c) and tilted (f) plants. In b, a caudal pod is visible on the left side of the stem.

Figure 4. Fluorescence measurement in the cell types. A is the full stem section with the

4 sampling areas (labelled 1, 2, 3 and 4). The different cell types selected for

829 fluorescence intensity quantification are indicated. B focuses on area 1 to illustrate the

830 manual selection of pixels in the different cell types; the red lines go from the outer part to

the inner stem part to the epidermis (ep), the bast fibre (bf), the cambium (ca), the xylem -

secondary (xs) and the xylem - primary (xp). A zoom is provided for each vignette.

Figure 5. Average values of the fluorescence intensity (in arbitrary units) obtained from
the ANOVA development stages × conditions. The drop caps indicate whether the values
are significantly different from each other or not.

Figure 6. Cellular type average fluorescence intensity according to development and tilting
conditions. The drop caps indicate whether the values are significantly different from each
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839 **Figure 7.** FTIR spectrum of bast fibres.

In a), the blue dots indicate the location of the bast fibre local spectra collected around
the flax stem samples. In b), the absorption bands for the main components in bast fibre
are indicated.

Figure 8. Effect of the development stage. The bast fibre second-derivative micro-FTIR
spectral signals of control plants are superimposed. Red arrows indicate the bands of
interest for the study.

Figure 9. Effect of a 90° tilt on the second-derivative micro-FTIR spectra of the flax bast
fibre. Red arrows indicate the bands of interest for the study.

Figure 10. Investigation of the putative polarity in the stem by exploring the fluorescence

of control versus tilted stems in bast fibres. Fig 10a) shows the arbitrary positioning of the

850 fluorescence measurements around the stem section. Fig 10b) shows the location of the

pixel fluorescence measurement in the bast fibre for M_C (square) and M_90 (lozenge).

Fig 10c) shows the fluorescence intensity measured and expressed linearly for the

853 tilted/untreated plants.

854

855 Table caption

Table 1. Building colour images from the three emission filters. The same maximum

857 intensity values were set for all images.

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Proteins (Filter 327-353 nm)

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Channel of the	Filter /	Maximum	Expected	Expected
colour image	bandpass	intensity	compound	colour
Blue	327-353 nm	500	Protein	blue
Green	420-460 nm	400	Hydroxycinnamates	Green to yellow
Red	480-550 nm	300	Lignin	Red to yellow

