

Revisiting the contribution of ATR-FTIR spectroscopy to characterize plant cell wall polysaccharides

Xuwei Liu, Catherine M.G.C. Renard, Sylvie Bureau, Carine Le Bourvellec

▶ To cite this version:

Xuwei Liu, Catherine M.G.C. Renard, Sylvie Bureau, Carine Le Bourvellec. Revisiting the contribution of ATR-FTIR spectroscopy to characterize plant cell wall polysaccharides. Carbohydrate Polymers, 2021, 262, pp.117935. 10.1016/j.carbpol.2021.117935 . hal-03182808

HAL Id: hal-03182808 https://hal.inrae.fr/hal-03182808

Submitted on 22 Mar 2023

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



Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License

Version of Record: https://www.sciencedirect.com/science/article/pii/S0144861721003222 Manuscript_e2efc31663fd0ecc28253bdb110de926

Revisiting the contribution of ATR-FTIR spectroscopy to

characterize plant cell wall polysaccharides

Xuwei Liu^a, Catherine M.G.C. Renard^{a, b}, Sylvie Bureau^a, Carine Le Bourvellec^{a, *} ^aINRAE, Avignon University, UMR SQPOV, F-84000 Avignon, France ^bINRAE, TRANSFORM, F-44000 Nantes, France

Corresponding author*

Carine Le Bourvellec (carine.le-bourvellec@inrae.fr)

INRAE, UMR408 SQPOV « Sécurité et Qualité des Produits d'Origine Végétale »

228 route de l'Aérodrome

CS 40509

F-84914 Avignon cedex 9

Tél: +33 (0)4 32 72 25 35

Other authors

Xuwei Liu: xuwei.liu@inrae.fr

Catherine M.G.C Renard: catherine.renard@inrae.fr

Sylvie Bureau: sylvie.bureau@inrae.fr





3 Abstract

The contribution of ATR-FTIR spectroscopy to study cell wall polysaccharides 4 (CWPs) was carefully investigated. The region 1800-800 cm⁻¹ was exploited using 5 principal component analysis and hierarchical clustering on a large range of different 6 powders of CWPs based on their precise chemical characterization. Relevant 7 wavenumbers were highlighted for each CWP: 1035 cm⁻¹ was attributed to 8 xylose-containing hemicelluloses, 1065 and 807 cm^{-1} to mannose-containing 9 hemicelluloses, 988 cm⁻¹ to cellulose, 1740 and 1600 cm⁻¹ to homogalacturonans 10 according to the degree of methylation. Some band positions were affected by 11 12 macromolecular arrangements (especially hemicellulose-cellulose interactions). 13 However, as arabinan and galactan did not reveal distinctive absorption bands, 14 ATR-FTIR spectroscopy did not allow the discrimination of cell walls differing by the abundance of these polysaccharides, e.g., those extracted from apple and beet. 15 Therefore, the application of ATR-FTIR could remain sometimes limited due to the 16 complexity of overlapping spectra bands and vibrational coupling from the large 17 diversity of CWP chemical bonds. 18

19 Keywords: ATR-FTIR; Polysaccharides; Cell walls; Cellulose; Pectins;
20 Hemicelluloses

21 Abbreviations:

AIS, alcohol insoluble solids; ATR-FTIR, Attenuated Total Reflectance Fourier
Transform Infrared Spectroscopy; DW, dry weight; PCA, Principal Component

24 Analysis; HCA: Hierarchical Cluster Analysis.

25 **1. Introduction**

26 Plant cell walls of the primary walls of dicots and non-grass monocots are 27 dynamic and ordered networks of natural carbohydrate polymers, constituted by an amorphous matrix mainly composed of pectins embedded in a network of cellulose 28 29 and hemicelluloses, as well as minor amounts of structural glycoproteins, phenolic 30 compounds and enzymes (Carpita, Sabularse, Montezinos, & Delmer, 1979). Cell 31 walls are highly variable according to species, developmental and maturity stages, 32 plant organs and environmental conditions (Anderson & Kieber, 2020; Burton, Gidley, 33 & Fincher, 2010). Therefore, this makes it difficult to easily identify and quantify cell 34 wall components.

35 In general, the structure and composition of plant cell walls are characterized 36 after sample extraction, pretreatment (e.g., acid hydrolysis) and diverse specific 37 biochemical analyses (e.g., chromatography, mass spectrometry, spectrophotometry), which are expensive and time-consuming. An advanced tool based on mid-infrared 38 39 spectroscopy would provide the advantages of rapid and easy analysis of the prepared 40 samples. Attenuated Total Reflectance Fourier Transform Infrared Spectroscopy 41 (ATR-FTIR) has been increasingly used for the rapid characterization of cell walls of 42 fruits and vegetables (Chylinska, Szymanska-Chargot, & Zdunek, 2016; Coimbra, Barros, Barros, Rutledge, & Delgadillo, 1998; Coimbra, Barros, Rutledge, & 43 44 Delgadillo, 1999; Ferreira, Barros, Coimbra, & Delgadillo, 2001; Kacurakova, Capek, Sasinkova, Wellner, & Ebringerova, 2000; Szymanska-Chargot, Chylinska, Kruk, & 45 Zdunek, 2015). Especially in recent years, it has become a powerful research 46

technology to clarify the composition of dry carbohydrate samples (Canteri, Renard,
Le Bourvellec, & Bureau, 2019). This method appears really convenient insofar as it
avoids undesirable structural changes that may occur during sample analysis, e.g.,
extraction and preparation. Moreover, ATR-FTIR can detect changes in the fruit and
vegetables during processing at the cell wall level (Lan, Renard, Jaillais, Leca, &
Bureau, 2020).

53 The identification of cell wall polysaccharides by infrared spectroscopy is 54 generally carried out on the different polysaccharide fractions obtained by sequential 55 extractions (the extraction of pure polysaccharides is imperfect), followed by ethanol precipitation and anion exchange chromatography. These extracted polysaccharides 56 are then characterized using chemical and biochemical methods (Brahem, Renard, 57 58 Gouble, Bureau, & Le Bourvellec, 2017; Coimbra et al., 1999; Renard, 2005; Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013). However, 59 these studies do not use purified polysaccharides to confirm the absorption bands 60 61 identified by comparison with the literature. Moreover, some studies have also performed polysaccharide analysis using spectral data but obtained from KBr pellets 62 63 or aqueous solutions, the classical way before the development of the ATR method (Kacurakova et al., 2000). Fruits and vegetables are highly hydrated and susceptible 64 65 to environmental conditions. Drying, not only prevents samples from oxidation and hydrolysis under the action of endogenous enzymes, but also concentrates samples by 66 67 water elimination, so it significantly improves the reflectance spectra of some specific components present in lower content than water (Lan et al., 2020). Therefore, the 68

69 systematic analysis of purified solid materials from cell wall polysaccharides by70 ATR-FTIR may improve their identification.

71 Moreover, some challenges exist due to the ATR-FTIR response of the different 72 cell wall polysaccharides. For example, according to our previous research, in spite of 73 very different structures and compositions, apple and beet cell walls were poorly 74 discriminated by Principal Component Analysis (PCA) based on ATR-FTIR spectra 75 (Liu, Renard, Rolland-Sabaté, Bureau, & Le Bourvellec, 2021). Therefore, we need to reconsider these results, knowing that the interactions between the internal 76 77 components of the cell walls (Le Bourvellec & Renard, 2012; Liu, Le Bourvellec, & Renard, 2020) may affect the absorption of these very complex mixtures. This study 78 combined ATR-FTIR and stoichiometry to characterize the abundance and 79 80 composition of cell wall polysaccharides, taking into account the heterogeneity and interactions between different cell wall components. To track the characteristic peaks 81 of each cell wall component, spectral data and conventional chemical methods are 82 83 used to study the composition of cell walls and internal structures. In order to evaluate the available information based on extracted samples, powders of cellulose, 84 85 hemicelluloses, and pectins were also scanned in ATR-FTIR. To identify the typology of the cell walls, both PCA and Hierarchical Cluster Analysis (HCA) were performed. 86 This study provided new explanations and experimental ideas for studying complex 87 natural polymer systems, and guidance for using ATR-FTIR data to clarify 88 89 carbohydrate structures, physical properties and interactions.

90 2. Materials and methods

91

2.1. Monosaccharide and polysaccharide samples

92 Monosaccharides (D-(+)-Arabinose, D-(-)-Fucose, D-(+)-Xylose, D-(+)-Mannose, 93 L-Rhamnose, D-(+)-Glucose, and D-(+)-galactose) and D-(+)-Galacturonic acid monohydrate were obtained from Fluka (Buchs, Switzerland). Arabinan (sugar beet), 94 linear 1,5-α-L-arabinan (sugar beet), debranched arabinan, galactan (potato), 95 96 rhamnogalacturonan I (from potato pectic fibre), and rhamnogalacturonan (from soybean pectic fibre), xylan, arabinoxylan, glucomannan, and xyloglucan were 97 98 provided from Megazyme (Bray, Ireland). Commercial apple and citrus peel pectins 99 (degrees of methylation \sim 75%), microcrystalline cellulose and poly-galacturonic acid 100 were provided by Sigma-Aldrich (Deisenhofen, Germany). Homogalacturonan DM 101 70 was supplied by Watrelot et al. (2013). The common names of cell wall 102 components and their abbreviations used in this study are presented in Table 1.

103 Native and modified cell walls and pectins from apple, beet and kiwifruit were supplied and characterized by Liu et al. (2021). The native cell wall samples were 104 105 named as apple cell wall (ACN), beet cell wall (BCN), ripe kiwifruit cell wall (KCRN) and overripe kiwifruit cell wall (KCON), and samples after boiling at pH 2.0, 3.5, and 106 6.0 are designated (AC, BC, KCR or KCO) - 2, (AC, BC, KCR or KCO) - 3, and (AC, 107 108 BC, KCR or KCO) - 6, respectively. Extracted pectins at pH 2.0, 3.5 and 6.0 from apple, beet and two kiwifruit cell walls at pH 2.0, 3.5, and 6.0 are designated (AP, BP, 109 110 KPR or KPO) - 2, (AP, BP, KPR or KPO) - 3, and (AP, BP, KPR or KPO) - 6, 111 respectively.

112 Table 1. The common names of cell wall components, their abbreviations and their ATR-FTIR frequencies (cm⁻¹) determined with our spectrometer of the studied plant cell wall

113 polysaccharides.

	Sample names	Abbreviations	Linkable peaks or regions (cm ⁻¹)
Monosaccharides	D-(-)-Arabinose	Ara	1312, 1128s, 1088s, 1050vs, 991vs, 940, 890s, 841s
	D-(+)-Xylose	Xyl	1146, 1123s, 1034vs, 1016s, 930s, 902s
	D-(+)-Mannose	Man	1110s, 1064s, 1034vs, 1016vs, 966s, 949s, 912, 879
	D-(+)-Galactose	Gal	1154s, 1142, 1100s, 1056vs, 1039vs, 990, 971, 953s, 827s
	L-Rhamnose	Rha	1375, 1290, 1226, 1145, 1116s, 1074s, 1026vs, 976s, 907, 874, 82
	D-(+)-Glucose	Glc	1228, 1206, 1150s, 1100s, 1052s, 1016vs, 991vs, 912s, 840s
	D-(-)-Fucose	Fuc	1334s, 1140s, 1095s, 1083vs, 1050vs, 976vs, 921, 868, 814
	D(+)-Galacturonic acid monohydrate	Gal A	1756s, 1708s, 1275, 1218, 1155, 1095vs, 1062vs, 1025vs, 823s
β-glucans	Microcrystalline cellulose	MCCE	1640, 1428, 1367, 1320, 1308, 1200, 1160, 1052 <i>s</i> , 1030 <i>vs</i> , 988 <i>s</i> , 8
	Yeast β-glucan	YGLU	1640, 1428, 1367, 1308, 1200, 1160, 1068s, 1030vs, 988s, 886
	Curdlan (1,3-β-o-glucan)	CGLU	1640, 1428, 1367, 1308, 1200, 1160, 1068s, 1030vs, 988s, 886
Hemicelluloses	Rye Arabinoxylan (59% xylose)	ARHV	1164, 1035vs, 983s, 890
	Wheet Arabinoxylan (64% xylose)	AXMB	1164, 1035vs, 983s, 890
	Wheet Arabinoxylan (77% xylose)	AXLB	1164, 1035vs, 983s, 890
	Xylan (Beechwood)	XYBW	1164, 1035vs, 983s, 890
	1,4-β-D-Mannan	MANB	1367, 1065vs, 1035s, 1013vs, 938s, 890, 870s, 807s
	Galactomannan (Carob)	GAMA	1065vs, 1027vs, 870s, 807s
	Xyloglucan (from tamarind seed)	XYGT	(1040-1010) <i>vs</i> , 939, 890
	Xyloglucan Oligosaccharides	XYGO	(1040-1010) <i>vs</i> , 939, 890
	Xyloglucan (Hepta-, +Octa, +Nona-saccharides)	XYGH	(1040-1010)vs, 939, 890
Pectins	Citrus peel pectin	CPPC	1740s, 1600, 1440, 1230, 1141, 1097s, 1014vs, 954, 914, 831

Table 1. (Continues)

Linkable peaks or regions (cm⁻¹) Sample names Abbreviations Commercial apple pectin APPC 1740s, 1600, 1440, 1230, 1141, 1097s, 1014vs, 954, 914, 831 ARSB Arabinan (sugar beet) 1600, the region of (1100 - 950) Linear 1,5-α-L-arabinan (sugar beet) LNAR 1208, 1115, 1086s, 1071s, 1043vs, 1022vs, 1004vs, 982vs, 948s Debranched arabinan (sugar beet) DBAR 1600, 1208, 1115, 1086s, 1071s, 1043vs, 1022vs, 1004vs, 982vs, 948s Galactan (Potato) GTAN 1600, 1405, 1039vs, 884s Rhamnogalacturonan I (from potato pectic fibre) RGPP 1740, 1600s, 1410, 1238, 1141s, 1097s, 1074s, 1014vs, 954 Rhamnogalacturonan (from soybean pectic fibre) RGSP 1600s, 1410, 1141s, 1097s, 1074s, 1014vs, 954 Homogalacturonan DM 70 HGTN 1740vs, 1440, 1230, 1140, 1097s, 1014vs, 970s, 914 Poly-galacturonic acid GALN 1590s, 1410s, 1330, 1141s, 1097s, 1014vs, 954s

114 * IR band intensity: *vs*, very strong; *s*, strong.

115 2.2. Characterization of carbohydrate composition

Sugar analysis was performed as previously described by Liu et al. (2021). For 116 117 neutral sugars analysis, 10 mg of cell walls or cellulose were submitted to a Saeman acid hydrolysis (Saeman, Moore, Mitchell, & Millett, 1954) and then to simple 118 hydrolysis (dissolved in 1 mol/L sulfuric acid) whereas soluble polysaccharides (10 119 120 mg) were only submitted to simple hydrolysis. The derivatization to alditol acetates (Englyst, Wiggins, & Cummings, 1982) allows the detection of sugars by gas 121 chromatography with a flame ionization detector (Agilent, Inc., Palo Alto, USA). 122 123 Galacturonic acid was measured by a meta-hydroxyl-diphenyl assay (Blumenkrantz & 124 Asboe-Hansen, 1973). The methanol was measured by a stable isotope dilution assay 125 using headspace-GC-MS (QP2010 Shimadzu Kyoto, Japan) as described by Renard & 126 Ginies (2009). The degree of methylation (DM) was then calculated as the molar ratio 127 of methanol to galacturonic acid.

128 **2.3. ATR-FTIR spectra**

All cell wall polysaccharide samples, in the form of dry powder, were stored in P₂O₅ atmosphere before analysis to remove residual water. ATR-FTIR spectra data (4000 to 600 cm⁻¹) were acquired at room temperature in a Tensor 27 FTIR spectrometer (Bruker Optics®, Wissembourg, France), using a single-reflectance horizontal ATR cell (Golden Gate with a diamond crystal, Bruker Optics®) equipped with a system to press the dried homogenized samples on the crystal surface (Bureau et al. 2012). Each sample was analyzed three times (using after homogenization three different aliquots of the powders) to consider its heterogeneity, and each spectrum was the average of 16 scans. Spectral pre-processing and data treatment using multivariate analyses were performed with MATLAB 7.5 (Mathworks Inc. Natick, MA) software using the SAISIR package (Cordella & Bertrand, 2014). The spectral data were pretreated with baseline correction and standard normal variate (SNV) to correct multiplicative interferences and variations in baseline shift before any multivariate analysis.

143 2.4. Statistical analysis

All biochemical analyses were presented as mean values of analytical triplicates 144 and the reproducibility of the results was expressed as pooled standard deviations 145 146 (Pooled SD). Pooled SD was calculated per series of replicates using the sum of individual variances weighted by the individual degrees of freedom (Box, Hunter, & 147 148 Hunter, 1978). A PCA was applied on the ATR-FTIR spectra in the range between 1800 and 800 cm^{-1} in order to study the repartition of the cellulose, hemicelluloses 149 150 and pectins in a space according to their composition and absorption bands. Spectral data pre-processing and PCA were performed using MATLAB 7.5 (Mathworks Inc. 151 152 Natick, MA) software using the SAISIR package (Cordella & Bertrand, 2014). HCA was performed using R software using FactoMineR (for computing) and factoextra 153 (for visualizing the results) (R Core Team., 2014). 154

155 **3. Results and discussion**

156 **3.1.** Characteristic bands of cell wall polysaccharides in the ATR-FTIR spectra

The compositions of the 58 cell wall polysaccharides from extracted and 157 commercial origin were determined in this study by both, the classical methods (Table 158 2, see Liu et al., 2021 for cell walls and extracted pectins) and ATR-FTIR 159 spectroscopy (Table 1, Figure 1). Detailed peak positions and assignments of each 160 pure cell wall polysaccharide were limited to the specific bands in the range of 161 1800-800 cm^{-1} (detected in solid or liquid form) in agreement with the previous 162 works (Canteri et al., 2019; Coimbra et al., 1998, 1999; Ferreira et al., 2001; Filippov 163 & Kohn, 1975; Gnanasambandam, R., Proctor, 2000; Kacurakova et al., 2000; 164 Kyomugasho, Christiaens, Shpigelman, Van Loey, & Hendrickx, 2015; McCann, 165 166 Hammouri, Wilson, Belton, & Roberts, 1992; Monsoor, Kalapathy, & Proctor, 2001; Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013) and 167 168 summarized in Table 3. Strong absorption bands in this region corresponding to the specific wavenumbers assigned to pectins (e.g., rhamnogalacturonan 169 and homogalacturonan), hemicelluloses (e.g., xyloglucan, mannan, galactomannan, 170 arabinoxylan and xylan) and cellulose (Figure 1), are detailed below. 171

172 **Table 2.** Composition of extracted cell walls and pectins from fruits and vegetables and commercial purified

173 cellulose, hemicelluloses and pectin components (mg/g dry weight, except for degree of methylation expressed

174 in %).

Codes	Rha	Fuc	Ara	Xyl	Man	Gal	Glc	Gal A	MeOH	DM
β-glucans										
MCCE	3	0	0	17	13	0	861	-	-	-
CGLU	0	0	0	0	0	0	705	-	-	-
YGLU	0	0	0	10	6	0	650	-	-	-
Hemicelluloses										
GAMA	0	0	5	0	776	190	7	-	-	-
ARHV	0	0	321	486	0	17	0	-	-	-
AXMB	1	0	288	693	0	7	5	-	-	-
AXLB	1	0	209	733	0	6	5	-	-	-
XYBW	12	0	6	716	0	10	8	-	-	-
MANB	0	0	0	0	960	18	8	-	-	-
XYGT	1	0	12	282	0	140	399	-	-	-
XYGO	3	0	37	287	0	123	515	-	-	-
XYGH	1	0	3	281	0	106	506	-	-	-
Commercial										
pectins										
CPPC	20	1	21	4	0	138	33	535	74	73
APPC	12	0	20	8	1	88	63	564	77	79
ARSB	52	0	584	0	0	114	0	83	-	-
LNAR	25	0	841	0	0	78	0	51	-	-
DBAR	68	0	504	0	0	193	48	98	-	-
GTAN	51	0	23	4	0	628	9	79		
RGPP	54	0	10	3	0	105	0	473	-	-
RGSP	81	64	21	120	0	96	0	403	-	-
HGTN	-	-	-	-	-	-	-	814	100	68
GALN	-	-	-	-	-	-	-	850	0	0
Pooled SD	1.0	0.3	8.3	7.0	5.8	2.3	8.6	4.0	2.5	3.2

175 Pooled SD: pooled standard deviation. Rha: rhamnose, Fuc: fucose, Ara: arabinose, Xyl: xylose, Man: mannose, 176 Gal: galactose, Glc: glucose, Gal A: galacturonic acid, MeOH: methanol, DM: degree of methylation. 177 Hemicelluloses (ARHV: Rye Arabinoxylan (59% Xylose), AXMB: Wheet Arabinoxylan (64% Xylose), AXLB: 178 Wheet Arabinoxylan (77% Xylose), XYBW: Xylan (Beechwood), MANB: 1,4-β-D-Mannan, XYGT: Xyloglucan 179 (from tamarind seed), XYGO: Xyloglucan Oligosaccharides, XYGH: Xyloglucan Oligosaccharides (Hepta-, +Octa, 180 +Nona-saccharides), GAMA: Galactomannan (Carob)); Pectins (APPC: Apple pectin, CPPC: Citrus peel pectin, 181 HGTN: Homogalacturonan DM 70, GALN: Polygalacturonic acid, DBAR: Debranched arabinan, ARSB: 182 Arabinan, LNAR: Linear arabinan, RGPP: Rhamnogalacturonan I, RGSP: Rhamnogalacturonan, GTAN: Galactan 183 (Potato)); β-glucans (MCCE: Microcrystalline cellulose, CGLU: 1,3-beta-o-glucan and YGLU: Yeast beta-glucan).





Wavenumbers (cm⁻¹)

186 Figure 1 ATR-FTIR spectra (pre-processed with Standard Normal Variate) of commercial purified and extracted 187 cell wall polysaccharides in solid form: A. Monosaccharides (Rha: rhamnose, Fuc: fucose, Ara: arabinose, Xyl: 188 xylose, Man: mannose, Gal: galactose, Glc: glucose, Gal A: galacturonic acid); B. Hemicelluloses (ARHV: Rye 189 Arabinoxylan (59% Xylose), AXMB: Wheet Arabinoxylan (64% Xylose), AXLB: Wheet Arabinoxylan (77% 190 Xylose), XYBW: Xylan (Beechwood), MANB: 1,4-β-D-Mannan, XYGT: Xyloglucan (from tamarind seed), 191 XYGO: Xyloglucan Oligosaccharides, XYGH: Xyloglucan Oligosaccharides (Hepta-, +Octa, +Nona-saccharides), 192 GAMA: Galactomannan (Carob)); C. Pectins (APPC: Apple pectin, CPPC: Citrus peel pectin, HGTN: 193 Homogalacturonan DM 70, GALN: Polygalacturonic acid, DBAR: Debranched arabinan, ARSB: Arabinan, LNAR:

- 194 Linear arabinan, RGPP: Rhamnogalacturonan I, RGSP: Rhamnogalacturonan, GTAN: Galactan (Potato)); D.
- 195 β-glucans (MCCE: Microcrystalline cellulose; CGLU: curdlan 1,3-beta-o-glucan; YGLU: Yeast beta-glucan).

Wavenumber range	r range Mone, or polysoceboride		Corresponding		
(cm ⁻¹)	in which it was detected	Band assignments	Wavenumber	References	
(detected)	III which it was detected		range (cm ⁻¹)*		
1756 & 1708	Gal A	Galacturonic acid (absence of glycosidic bond)	-	-	
				(Filippov & Kohn, 1975; Gnanasambandam, R.,	
1740	APPC, CPPC and HGTN	C=O stretching vibration of alkyl ester (pectin)	1745-1730	Proctor, 2000; McCann et al., 1992; Monsoor et	
				al., 2001; Szymanska-Chargot & Zdunek, 2013)	
1640	MCCE, CGLU and YGLU	H–O–H bending vibration absorbed water	1640	(Szymanska-Chargot et al., 2015)	
	CALN CDDC ADOD			(Filippov & Kohn, 1975; Gnanasambandam, R.,	
1605 - 1595	GALN, CPPC, ARSB,	COO antisymmetric stretching polygalacturonic acid,	1630-1600	Proctor, 2000; McCann et al., 1992; Monsoor et	
	DBAR, RGPP and RGSP	free carboxyl group		al., 2001; Szymanska-Chargot & Zdunek, 2013)	
1525		Amid II N-H deformation (proteins); lignin and	1550	(MaConn et al. 1002)	
1525	KCR/OS	phenolic back bone	1550	(MCCallifiet al., 1992)	
1474	ARHV, XYBW and XYGT	Xylose-containing hemicellulose	-	-	
1440	ADDC CDDC and UCTN	Asymmetric stretching modes vibration of methyl	1440	(Canteri et al., 2019; Szymanska-Chargot et al.,	
1440	APPC, CPPC and HOTN	esters (pectin)	1440	2015)	
1428	MCCE, CGLU and YGLU	CH ₂ symmetric bending (cellulose)	1428	(Szymanska-Chargot & Zdunek, 2013)	
1410	CALN DOD and DOOD	COO ⁻ symmetric stretching, free carboxyl group	1410	(Compared a Charact & Zharach 2012)	
1410	GALN, KGPP and KGSP	(rhamnogalacturonan and homogalacturonan)	1410	(Szymanska-Chargot & Zdunek, 2013)	
	MCCE CCLU and VCLU	C. Hailanting and CH. has dies (callulate		(Symmetry Charget et al. 2015)	
1367	57 MCCE, CGLU and YGLU, XYGT, GAMA and MANB	C-H vibrations and CH_2 bending (cellulose,	1370, 1362	(Szymańska-Chargot et al., 2013;	
		hemicelluloses)		Szymanska-Chargot & Zdunek, 2013)	

Table 3. The main ATR-FTIR absorption bands, the polysaccharides in which they were detected and their tentative assignment. For polysaccharide identification (detailed in Table 1).

Table	3. ((Continues)
		(/

Wavenumber range (cm ⁻¹) (detected)	Mono- or polysaccharide in which it was detected	Band assignments	Corresponding Wavenumber range (cm ⁻¹)*	References
1331 1330	Fuc HGTN, GALN, RGPP and	Fucose (absence of glycosidic bond) Bending of O–H groups in pyranose ring of pectins	- 1331-1320	(Szymanska-Chargot et al., 2015;
1312	RGSP Ara	Arabinose (absence of glycosidic bond)	-	Szymanska-Chargot & Zdunek, 2013)
1308	MCCE, CGLU and YGLU	CH ₂ symmetric bending or CH ₂ rocking vibration (cellulose)	1317-1313	(Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013)
1290	Rha	Rhamnose (absence of glycosidic bond)	-	
1238	RGPP	Rhamnogalacturonan	-	
1230	APPC, CPPC and HGTN	C–O stretching (pectins)	1240, 1230	(Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013)
1218	Gal A	Galacturonic acid (absence of glycosidic bond)	-	-
1164	ARHV, XYBW and XYGT	Glycosidic bond vibrations (O–C–O) (xylose-containing hemicellulose)	1173, 1153, 1147	(Coimbra et al., 1999; Kacurakova et al., 2000)
1160	KCR/Os	Glycosidic bond vibrations (O–C–O) (cellulose in cell walls)	1160	(Canteri et al., 2019; Szymanska-Chargot & Zdunek, 2013)
1152	MANB and GAMA	Glycosidic bond vibrations (O–C–O) (mannose-containing hemicellulose)	1150	(Szymanska-Chargot et al., 2015)
1141	APPC, CPPC, GALN, HGTN, RGPP and RGSP	Glycosidic bond vibrations (O–C–O) (pectin)	1150-1143	(Coimbra et al., 1998, 1999)
1126	Ara	Arabinose (absence of glycosidic bond)	-	

Table 3. ((Continues))
	(001111100)	

Wavenumber range (cm ⁻¹) (detected)	Mono- or polysaccharide in which it was detected	Band assignments	Corresponding Wavenumber range (cm ⁻¹)*	References
1097	APPC, CPPC, HGTN, GALN, RGPP and RGSP	C–O stretching, C–C stretching ring pectin	1100-1090	(Coimbra et al., 1998; Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013)
1074	GTAN, RGPP and RGSP	C-C stretching ring (galactan and rhamnogalacturonan)	1072, 1070	(Kacurakova et al., 2000)
1068	MCCE, CGLU and YGLU	C–O stretching, C–C stretching, C6–H2–O6 (cellulose)	1059, 1047	(Szymanska-Chargot et al., 2015)
1065	MANB and GAMA	C–O stretching, C–C stretching (mannose-containing hemicellulose)	1064	(Kacurakova et al., 2000)
1039	GTAN	C–C stretching ring (galactan)	1038	(Kacurakova et al., 2000)
1035	ARHV, XYBW and XYGT	C–O stretching, C–C stretching (xylose-containing hemicellulose)	1042, 1041, 1038	(Canteri et al., 2019; Coimbra et al., 1999; Kacurakova et al., 2000)
1030	MCCE, CGLU and YGLU	C–O stretching, C–C stretching, C6–H2–O6 (cellulose)	1034, 1030	(Kacurakova et al., 2000; Szymanska-Chargot & Zdunek, 2013)
1027	GAMA	C–O stretching, C–C stretching, C6–H2–O6 (galactomannan)	1034	(Kacurakova et al., 2000)
1014	APPC, CPPC, GALN, HGTN, DBAR, ARSB, GTAN, RGPP and RGSP	C–O stretching, C–C stretching pectin (C2–C3, C2–O2, C1–O1) backbone vibrations (pectin)	1020, 1015, 1014	(Coimbra et al., 1998, 1999; Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013)
1013	MANB	C–O stretching, C–C stretching (mannan)	-	
991	Glc and Ara	Glucose and arabinose (absence of glycosidic bond)	-	
988	MCCE, CGLU and YGLU	C–O stretching, C–C stretching cellulose (C6–H2–O6)	1000, 985	(Canteri et al., 2019; Szymanska-Chargot & Zdunek, 2013)

	(a)
Toble 4	(Continues)
I ADIC J.	COMUNUEST
	(, , , , , , , ,

Wavenumber range (cm ⁻¹) (detected)	Mono or polysaccharide in which it was detected	Band assignments	Corresponding Wavenumber range (cm ⁻¹)*	References
983	ARHV and XYBW	Xylan and arabinoxylan	-	
982	DBAR, ARSB and LNAR	Arabinan	-	
970	APPC, CPPC and HGTN	Pectins	972	(Kacurakova et al., 2000)
954	RGSP	CO bending (pectins)	952	(Coimbra et al., 1999; Szymanska-Chargot & Zdunek, 2013)
939	XYGT, GAMA and MANB	Ring vibration (hemicellulose and arabinan)	941	(Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013)
925 and 908	Xyl	Xylose (absence of glycosidic bond)	-	
914	APPC, CPPC, HGTN and GALN	Ring vibration (pectin)	-	
890	XYGT, ARHV and XYBW	C1–H bending (xylose-containing hemicellulose)	893	(Szymanska-Chargot & Zdunek, 2013)
886	KCR/Os	C1–H bending (cellulose)	899, 895	(Canteri et al., 2019; Szymanska-Chargot & Zdunek, 2013)
884	GTAN	C1-H bending (galactan)	883	
870	GAMA and MANB	C1-H bending (mannose-containing polysaccharide)	-	
840	Ara	Arabinose (absence of glycosidic bond)	-	
831	APPC, CPPC, GALN and HGTN	Ring vibration (pectin)	833-830	(Szymanska-Chargot et al., 2015; Szymanska-Chargot & Zdunek, 2013)
807	GAMA and MANB	Ring vibration (mannose-containing hemicellulose)	-	

197 * Reference from the literatures. Monosaccharides (Rha: rhamnose, Fuc: fucose, Ara: arabinose, Xyl: xylose, Man: mannose, Gal: galactose, Glc: glucose, Gal A: galacturonic acid);



203 3.1.1. Monosaccharides

204 Monosaccharides, without any glycosidic linkage to other units, are the simplest 205 component units of the cell wall polysaccharides, and have been shown to be important in polymer analysis and structure elucidation (Wiercigroch et al., 2017). It 206 was necessary to consider here hexopyranoses (glucose, mannose and galactose), 207 208 dehydro-hexopyranoses (rhamnose and fucose), pentopyranose (xylose), and 209 pentofuranose (arabinose), standards with different positions of hydroxyl groups on C-2, C-3, and C-4 (Figure 1A). The spectra of pentoses (arabinose and xylose) were 210 dominated by a band at 991 and 1034 cm⁻¹, respectively, mainly due to the v(C-C), 211 212 v(C-O) and $\beta(C-CH)$ vibrations (Edwards, 1976). For hexoses, D-(+)-glucose was observed by the main band at 991 cm⁻¹. D-(+)-galactose is almost identical in structure 213 214 to D-(+)-glucose, with a different orientation in the C-4 OH group, but with a distinct spectral difference due to their free crystalline structure (Figure 1A). Similarly, the 215 spectrum of other hexopyranoses with different hydroxyl group orientations on C-2, 216 C-3, and C-4 was also significantly different. Therefore, the relative positions of 217 (C-OH) essentially affected the spectrum through variations in their spatial 218 219 arrangement and interactions. The detailed ATR-FTIR bands of monosaccharides with 220 modes assignments are listed in Table 1.

221 **3.1.2. Pectic components**

Pectins are acidic hetero-polysaccharides mainly composed of homogalacturonan,rhamnogalacturonan I and II, and different neutral sugar side chains (e.g., arabinan)

and galactan). The spectra of apple pectins (APPC) and citrus peel pectins (CPPC) 224 had very similar characteristic bands (Figure 1C) centering at 1740, 1600, 1440, 1230, 225 1141, 1097, 1014, 954, 914 and 831 cm⁻¹. They have also similar levels of neutral 226 sugars, galacturonic acid and degree of methylation (Table 2). The band at 1740 cm^{-1} 227 is assigned to the esterified galacturonic acids. This band is characteristic of pectins 228 229 with a high degree of methylation (DM) such as high methylated homogalacturonan 230 (DM=70) (HGTN). Inversely, the band of poly-galacturonic acid (GALN), which is non-esterified, was at 1600 cm^{-1} due to the COO⁻ carboxylate ion stretching. These 231 two main characteristic bands of pectins are fixed and similar to those previously 232 233 reported (Coimbra et al., 1999; Filippov & Kohn, 1975; Gnanasambandam, R., Proctor, 2000; Kacurakova et al., 2000; Reintjes, Musco, & Joseph, 1962; 234 Szymanska-Chargot et al., 2015; Wojdyło, Figiel, Lech, Nowicka, & Oszmiański, 235 2014). 236

237 For a further interpretation of these spectra, pure pectic components were characterized in the same conditions. Notably, although these parts of the 238 polysaccharide components were not a complete pectin structure found in the plant, 239 they are representative of different subunits of the pectins. For rhamnogalacturonan 240 (RGPP and RGSP), the band at 1014 cm⁻¹ was the strongest (Figure 1C). However, 241 this peak overlapped with the main peak of commercial pectins (APPC and CPPC). 242 243 Some other specific bands, assigned to rhamnogalacturonan (RGPP and RGSP) can be used such as 1410, 1238 and 1074 cm⁻¹. In fact, RGSP contained more xylose than 244 245 RGPP (120 mg/g vs 3 mg/g respectively) (Table 2). Based on the maximum peak of 24

xylose at 1035 cm⁻¹ for RGSP, but not for RGPP (Figure 1C), 1035 cm⁻¹ may be used
to differentiate the abundance of xylose in cell wall polysaccharides.

248 In the case of the main neutral sugar side chains of pectins (Figure 1C): (i) galactan (GTAN) was characterized by bands at 1039 cm⁻¹, 1014 and 884 cm⁻¹; and 249 (ii) arabinan, such as present in sugar beet arabinan (ARSB), linear arabinan (LNAR) 250 251 and debranched arabinan (DBAR), were more visible in the region between 1100 and 950 cm⁻¹. The arabinans with different linearity and branching degrees had some 252 minor peak intensity differences in this region. For example, the main band of LNAR 253 (with the highest arabinose content, Table 2) was close to 980 cm^{-1} , which may be 254 255 used to assess arabinan. However, the characteristic peaks of arabinan (sugar beet) were in the region and also within the range of the main peaks of other pectins (e.g., 256 commercial pectins, rhamnogalacturonan and galactan) not allowing a clear 257 assignment of bands between 1100 and 950 cm⁻¹. In aqueous solutions, Kacurakova et 258 al. (2000) observed bands at 1070 and 1043 cm⁻¹ for rhamnogalacturonan, 1072 cm⁻¹ 259 for galactan and 1039 cm⁻¹ arabinan, which were too close for reliable discrimination. 260 These differences may be due to the different states (solid or solution) of the cell wall 261 262 compounds and probably also to the specificity of the used spectrometers.

For pectic homogalacturonans with more or less methylation, the main characteristic peaks (especially 1740, 1600, 1097 and 1014 cm⁻¹) were stable regardless of whether they are in a solid crystalline state or in an aqueous solution. However, the identification of spectral characteristic peaks of arabinan, galactan, and 267 rhamnogalacturonan required caution as their peaks were dependent on the sample268 state, which could lead to their overlapping with those of hemicelluloses and pectins.

269

3.1.3. Hemicelluloses and cellulose

Bands at 1474, 1367, 1164, 1152, 1065, 1035, 1027, 1013, 983, 939, 890, 870 and 270 807 cm⁻¹ were identified for hemicelluloses (Figure 1B). Xylan (XYBW) had 271 characteristic absorption bands at 1035 cm⁻¹ and 983 cm⁻¹. The arabinoxylans with 272 different xylose contents (Table 2), e.g., ARHV (59%), AXMB (64%) and AXLB 273 (77%), showed a main band at 1035 cm⁻¹ for which the intensity varied with the 274 xylose content. Xyloglucan (XYGT, XYGO and XYGH) absorbed in the region of 275 1040-1010 cm⁻¹, like hemicelluloses and cellulose making it difficult to be identified. 276 Galactomannan (GAMA) and mannan (MANB) had bands at 1027 and 1013 cm⁻¹, 277 respectively. In addition, they had a common secondary band at 1065 cm⁻¹, and two 278 specific peaks at 870 and 807 cm⁻¹ (also found in softwood sample, Simonović, 279 Stevanic, Djikanović, Salmén, & Radotić, 2011), which allowed them to be 280 281 distinguished from others (Figure 1B). Therefore, the bands which could be assigned to xylose- and mannose- containing hemicelluloses were 1474, 1164, 1035, 983 and 282 890 cm⁻¹ for xylose-, and 1152, 1065, 1027, 1013, 939, 870 and 807 cm⁻¹ for 283 mannose-, respectively (Table 3). This was confirmed for xylose- containing 284 hemicelluloses including XYBW, ARHV, AXMB, AXLB, XYGT, XYGO and XYGT, 285 and mannose- containing hemicelluloses including MANB and GAMA. 286

287

The bands characteristic of cellulose, β -(1 \rightarrow 4)-linked glucan, for MCCE, were at

288 1640, 1428, 1367, 1320, 1308, 1200, 1160, 1052, 1030, 988 and 893 cm⁻¹ (Figure 1D), 289 whereas bands at 1068 and 886 cm⁻¹ (instead of 1052 and 893 cm⁻¹) were identified 290 for β-(1→3)-glucans (YGLU and CGLU). These slight differences for two peaks may 291 contribute to distinguish 1,3-β- and 1,4-β- bonds.

292 **3.1.4.** Glycosidic linkage

293 The glycosidic linkages are an important characteristic structure in polysaccharides and influence the spectral changes in aqueous solutions (Jockusch et 294 295 al., 2004; Kačuráková & Mathlouthi, 1996; Kanou, Nakanishi, Hashimoto, & Kameokaj, 2005; Nikonenko, Buslov, Sushko, & Zhbankov, 2000; Wiercigroch et al., 296 2017). For example, the spectra of the glycosidic bonds with different positions and 297 configurations of oligo- and poly- saccharides in aqueous solution differ markedly 298 299 from monosaccharides, with stretching vibrations [v(CO)] of the C-O-C glycosidic linkage being the marker of the polysaccharide configuration. These vibrations appear 300 in the two spectral ranges 1160-1130 and 1000-960 cm⁻¹ (Kacurakova et al., 2000; 301 302 Kačuráková & Mathlouthi, 1996; Wiercigroch et al., 2017). Similarly, these bands appear in cell wall polysaccharides in the solid states when compared with 303 monosaccharides (Figure 1). For xylose units with β -(1 \rightarrow 4) glycosidic linkage 304 (XYBW) bands were found at about 1164 cm⁻¹, for glucose units with β -(1 \rightarrow 4) 305 glycosidic linkage (YGLU, CGLU and MCCE) bands were found at 1160 cm⁻¹ and 306 for mannose units with β -(1 \rightarrow 4) glycosidic linkage (MANB) bands were found at 307 1152 cm⁻¹. Bands at 1141 cm⁻¹ for pectins originate from the stretching motion of the 308

309 CO bond within the glycosidic linkage (Coimbra et al., 1999). Therefore, the 310 involvement of glycosidic bonds influenced the spectrum through changes in their 311 spatial arrangement, type and position.

However, some monosaccharides also showed spectral bands in the region between 1100 to 1000 cm⁻¹ presenting overlapping with those of polysaccharides (Figure 1, Table 1).



316 Figure 2 PCA scores scatter plots of 1) commercial and extracted pectins, 2) rhamnogalacturonan, 3) galactan, 4) arabinan, 5) mannose- containing hemicelluloses, 6) xylose- containing 317 hemicelluloses and 7) cellulose (A), and all cell wall materials (excluding monosaccharides): 8) kiwifruit cell walls, 9) apple and beet cell walls (C). ATR-FTIR spectra in the range 1800 to 800 318 cm⁻¹ with their PCA loading profile of components PC1 and PC2 (B) and (D). Monosaccharides (Rha: rhamnose, Fuc: fucose, Ara: arabinose, Xyl: xylose, Man: mannose, Gal: galactose, Glc: 319 glucose, Gal A: galacturonic acid); Hemicelluloses (ARHV: Rye Arabinoxylan (59% Xylose), AXMB: Wheet Arabinoxylan (64% Xylose), AXLB: Wheet Arabinoxylan (77% Xylose), XYBW: 320 Xylan (Beechwood), MANB: 1,4-β-D-Mannan, XYGT: Xyloglucan (from tamarind seed), XYGO: Xyloglucan Oligosaccharides, XYGH: Xyloglucan Oligosaccharides (Hepta-, +Octa, 321 +Nona-saccharides), GAMA: Galactomannan (Carob)); Pectins (APPC: Apple pectin, CPPC: Citrus peel pectin, HGTN: Homogalacturonan DM 70, GALN: Polygalacturonic acid, DBAR: 322 Debranched arabinan, ARSB: Arabinan, LNAR: Linear arabinan, RGPP: Rhamnogalacturonan I, RGSP: Rhamnogalacturonan, GTAN: Galactan (Potato)); β-glucans: (MCCE: Microcrystalline 323 cellulose; CGLU: curdlan, 1,3-beta-o-glucan; YGLU: Yeast beta-glucan).

324 **3.2. Discrimination of cell walls**

325 Two multivariate analyses were used to study the discrimination of complex fruit 326 cell walls in comparison with that obtained on standard compounds. A Principal component analysis (PCA) was carried out using the spectral data (Figure 2A) to 327 identify the mapping of cell wall polysaccharides (excluding extracted cell walls and 328 329 monosaccharides). PC1 and PC2 explained respectively 50% and 16% of the total 330 variance. Along the PC1, the samples were well discriminated due to the different types of cell wall polysaccharides. Positive loadings of PC1 covered wavenumbers 331 characteristic of xvlose-containing (1035 cm^{-1}) hemicellulose and cellulose (988 cm^{-1}) 332 333 (Figure 2B). The negative high values of PC1 were obtained for wavenumbers at 1740 and 1600 cm⁻¹ characteristic of esterified and non-esterified carboxyl groups in 334 335 pectins (Figure 2B), respectively and of 1440, 1320, 1230, 1141, 1097, 914 and 831 cm⁻¹ bands characteristic of pectins (Figure 2B). In addition, negative PC1 loading 336 appeared for wavenumbers at 1410 cm^{-1} characteristic for rhamnogalacturonan. The 337 338 PC2 separated the esterified pectins and arabinan on the negative side (at the bottom) from the less esterified pectins, galactan and mannose-containing hemicelluloses (at 339 the top). The bands at (1600 and 1141 cm⁻¹), 1410 cm⁻¹, and (1065 and 807 cm⁻¹) 340 341 were attributed to respectively free carbonyl group of pectins, rhamnogalacturonan and mannose-containing hemicelluloses with a positive correlation to PC2. The region 342 closes to 1740 and 970 cm⁻¹ corresponded to the esterified pectins with a negative 343 correlation to PC2. 344

345 The same approach was used on the extracted cell walls from apple, beet and kiwifruit (Figure 2C). The cell wall polysaccharides were divided into nine groups: 1) 346 347 commercial and extracted pectins; 2) rhamnogalacturonans; 3) galactans; 4) arabinans; 5) mannose- containing hemicelluloses; 6) xylose- containing hemicelluloses; 7) 348 349 cellulose; 8) kiwifruit cell walls; 9) apple and beet cell walls. The first seven groups 350 are similar to Figure 2A, with the later addition of the extracted cell walls in the 351 center of the Figure 2C. This is consistent with the expected results as they contain 352 intact cell wall polysaccharide components. However, the extracted cell walls issued 353 from different species and extraction conditions were not well separated, especially for apple and beet cell walls. How can we explain the lack of discrimination of cell 354 walls between apple and beet, whereas their composition differ significantly (Liu et 355 356 al., 2021)? As shown on the PCA, the kiwifruit cell walls were at the upper left in the middle of the Figure 2C, while the apple and beet cell walls were at the bottom right 357 in the middle of the Figure 2C. This was probably linked to the high cellulose and 358 359 relatively low galacturonic acid for kiwifruit, and the high galacturonic acid for both apple and beet (140 to 206 mg/g for apple, 141 to 225 mg/g for beet) (Liu et al., 2021). 360 361 However, the cell walls of apple and beet were not distinguished, in spite of marked chemical and structural differences (Liu et al., 2021). Even a PCA performed on the 362 ATR-FTIR spectra of the cell walls of apple and beet alone did not separate them well 363 (data not shown). Beet cell walls are rich in arabinan (111 to 189 mg/g), contain 364 365 ferulic acid, and have only minor amounts of rhamnose, fucose, xylose and mannose. Apple cell walls contain as much or more xylose (65 to 75 mg/g) and galactose (63 366

-67 mg/g) than arabinose (27 to 66 mg/g). However, the expected separation of apple
and beet cell walls spectra by the characteristic peaks of arabinan (982 cm⁻¹) or
galactan (1039 cm⁻¹) was not observed. Probably the characteristic peaks were not
detected or were overlapped with those of other cell wall components, e.g.,
hemicelluloses or cellulose (Figure 1). This may be a limitation of ATR-FTIR in cell
wall characterization, because arabinan and galactan are well known to change during
ripening or processing.

374 PC loadings (Figure 2D) were very close to those obtained on the standard compounds (Figure 2B), probably reflecting the fact that the cell walls are by 375 376 themselves combinations of pectins, hemicelluloses and cellulose. PC2 loadings had an extra shoulder peak at 1526 cm⁻¹, which may be attributed to lignin and ferulic acid, 377 378 but occurring in combination with hemicelluloses (e.g., xylans and xyloglucans). The non-polysaccharide compounds in the cell wall, such as proteins (e.g., C = O of 379 amides at 1655 cm⁻¹ and N-H of amides at 1540, and 1234 cm⁻¹), lignins (e.g., 1520, 380 1410 and 921 cm⁻¹) and other phenolic compounds (e.g., 1520, 1440, 1284, 1196 and 381 1075 cm⁻¹) have absorption bands which mare interfere with those of polysaccharides. 382 383 This needs to be taken into account when characterizing cell walls using ATR-FTIR. 384 Moreover, the structure of xylans varies with the source of the plant, such as wheat or beechwood. As an example, some phenolic compounds can couple beechwood xylan 385 386 chains via ferulate dimerization (dehydrodiferulate cross-links) and/or incorporation into lignin, thus affecting their spectral bands (Kačuráková et al., 1999). 387



389 Figure 3 Hierarchical cluster analysis dendrogram of 58 cell wall and cell wall polysaccharide and monosaccharide samples based on average ATR-FTIR spectra in the range 4000 to 600 cm⁻¹ 390 using Ward's clustering algorithm with Euclidian distance. From left to right, the groups are (i) commercial and extracted pectins; (ii) kiwifruit cell walls and RG; (iii) monosaccharides; (iv) 391 cellulose and hemicelluloses and (v) apple and beet cell walls. Monosaccharides (Rha: rhamnose, Fuc: fucose, Ara: arabinose, Xyl: xylose, Man: mannose, Gal: galactose, Glc: glucose, Gal A: 392 galacturonic acid); Hemicelluloses (ARHV: Rye Arabinoxylan (59% Xylose), AXMB: Wheet Arabinoxylan (64% Xylose), AXLB: Wheet Arabinoxylan (77% Xylose), XYBW: Xylan 393 (Beechwood), MANB: 1,4-β-D-Mannan, XYGT: Xyloglucan (from tamarind seed), XYGO: Xyloglucan Oligosaccharides, XYGH: Xyloglucan Oligosaccharides (Hepta-, +Octa, 394 +Nona-saccharides), GAMA: Galactomannan (Carob)); Pectins (APPC: Apple pectin, CPPC: Citrus peel pectin, HGTN: Homogalacturonan DM 70, GALN: Polygalacturonic acid, DBAR: 395 Debranched arabinan, ARSB: Arabinan, LNAR: Linear arabinan, RGPP: Rhamnogalacturonan I, RGSP: Rhamnogalacturonan, GTAN: Galactan (Potato)); β-glucans: (MCCE: Microcrystalline 396 cellulose; CGLU: curdlan, 1,3-beta-o-glucan; YGLU: Yeast beta-glucan).

Hierarchical cluster analysis (HCA) is widely applied as an unsupervised 397 classification method to calculate distances between samples and cluster them 398 according to this distance, based here on their spectra (Granato, Santos, Escher, 399 Ferreira, & Maggio, 2018). HCA highlighted five groups (Figure 3). Group 1 400 401 clustered samples with high galacturonic acid contents from apple, beet, citrus and 402 kiwifruit pectins. Group 2 contained samples with linear pectins and less side chains 403 from kiwifruit cell walls, polygalacturonic acid and rhamnogalacturonan. Group 3 associated monosaccharide samples with absence of glycosidic bonds. Group 4 404 405 clustered samples including hemicelluloses and cellulose. Group 5 clustered samples with high methylated pectins and rich in arabinose and galactose, extracted from 406 apple, beet cell walls, galactan and arabinan. Therefore, ATR-FTIR spectroscopy 407 408 coupled with chemometrics allowed a good discrimination of cell walls related to their compositions and structures, giving some classes according to the different kinds 409 of pectins, hemicelluloses and cellulose. 410

The cell walls represented a polymer system in a complex mixture with a 411 diversity of both compositions and structures. PCA and HCA performed on the 412 413 spectral data allowed to discriminate samples according to their cell wall 414 polysaccharides. ATR-FTIR could be considered as a fast and easy way to distinguish 415 different types of cell walls. Due to their complexity and the numerous spectral bands 416 for each component, with the addition of overlapping, it was therefore difficult to 417 assign each band to a compound chemical structure. However, the observed changes in intensity or presence/absence of some bands reflected the differences in 418 36

419 composition between the cell walls. And, slight changes in the strength and position
420 of individual bands could also be due to the different conformations of cell wall
421 polymers and the interactions between individual components.

422 **4.** Conclusions

ATR-FTIR spectra in the region between 1800 and 800 cm⁻¹ combined with PCA has been widely applied to study the main polysaccharides present in the complex cell walls. However, it is not always possible to analyze the structural changes in cell wall polysaccharides at the molecular level and not all absorption bands allow differentiation. Their complex structures, compositions, glycosidic linkage patterns and the interactions between polysaccharides (or even with polyphenols and proteins) make the application of ATR-FTIR to plant cell walls still challenging.

430 What can be determined is that: 1) xylan, arabinoxylan and xyloglucan all had the same characteristic band at 1035 cm⁻¹; 2) cellulose showed a characteristic band at 431 988 cm⁻¹; 3) the mannan and galactomannan were identified by the bands at 1065 and 432 807 cm⁻¹; 4) the degree of methylation of pectin homogalacturonans was easily 433 determined by the relative height of the two bands at 1740 and 1600 cm⁻¹. According 434 to these results, the analysis of purified cell wall polysaccharides could be easily and 435 436 successfully performed directly with these bands giving information on the main cell wall compounds: pectins, hemicelluloses and cellulose. 437

However, some difficulties remain to identify intact cell wall components and inparticular to discriminate cell walls of apple and beet in relation to the bands of

arabinan and galactan. The main chain made of arabinan did not give available
characteristic peaks. The specific band of galactan at 1039 cm⁻¹ was overlapped with
the bands of hemicelluloses. Therefore, the application of ATR-FTIR spectroscopy for
the characterization of cell wall polysaccharides requires more in-depth research and
should be used in combination with other analytical techniques.

445 Acknowledgements

LIU Xuwei would like to acknowledge China Scholarship Council (CSC) and
Institut National de Recherche pour l'Agriculture, l'Alimentation, et l'Environnement

448 (INRAE) for financial support to his PhD study.

449 **References**

- Anderson, C. T., & Kieber, J. J. (2020). Dynamic Construction, Perception, and
 Remodeling of Plant Cell Walls. *Annual Review of Plant Biology*, 71(1), 39–69.
 https://doi.org/10.1146/annurev-arplant-081519-035846
- Blumenkrantz, N., & Asboe-Hansen, G. (1973). New method for quantitative
 determination of uronic acids. *Analytical Biochemistry*, 54(2), 484–489.
 https://doi.org/10.1016/0003-2697(73)90377-1
- Brahem, M., Renard, C. M. G. C., Gouble, B., Bureau, S., & Le Bourvellec, C. (2017).
 Characterization of tissue specific differences in cell wall polysaccharides of ripe
 and overripe pear fruit. *Carbohydrate Polymers*, *156*, 152–164.
- 459 https://doi.org/10.1016/j.carbpol.2016.09.019
- Bureau, S., Ścibisz, I., Le Bourvellec, C., & Renard, C. M. G. C. (2012). Effect of
 sample preparation on the measurement of sugars, organic acids, and
 polyphenols in apple fruit by mid-infrared spectroscopy. *Journal of Agricultural and Food Chemistry*, 60(14), 3551–3563. https://doi.org/10.1021/jf204785w
- Burton, R. A., Gidley, M. J., & Fincher, G. B. (2010). Heterogeneity in the chemistry,
 structure and function of plant cell walls. *Nature Chemical Biology*, 6(10), 724–
 732. https://doi.org/10.1038/nchembio.439
- 467 Canteri, M. H. G., Renard, C. M. G. C., Le Bourvellec, C., & Bureau, S. (2019).
 468 ATR-FTIR spectroscopy to determine cell wall composition: Application on a
 469 large diversity of fruits and vegetables. *Carbohydrate Polymers*, 212, 186–196.
 470 https://doi.org/10.1016/j.carbpol.2019.02.021
- 471 Carpita, N., Sabularse, D., Montezinos, D., & Delmer, D. P. (1979). Determination of
 472 the pore size of cell walls of living plant cells. *Science*, 205(4411), 1144–1147.
 473 https://doi.org/10.1126/science.205.4411.1144
- 474 Chylinska, M., Szymanska-Chargot, M., & Zdunek, A. (2016). FT-IR and FT-Raman
 475 characterization of non-cellulosic polysaccharides fractions isolated from plant
- 476 cell wall. *Carbohydrate Polymers*, *154*, 48–54.
- 477 https://doi.org/10.1016/j.carbpol.2016.07.121

- 478 Coimbra, M. A., Barros, A., Barros, M., Rutledge, D., & Delgadillo, I. (1998).
- 479 Multivariate analysis of uronic acid and neutral sugars in whole pectic samples
 480 by FT-IR spectroscopy. *Carbohydrate Polymers*, *37*, 241–248.
- 481 https://doi.org/10.1016/S0144-8617(98)00066-6
- 482 Coimbra, M. A., Barros, A., Rutledge, D. N., & Delgadillo, I. (1999). FTIR
 483 spectroscopy as a tool for the analysis of olive pulp cell-wall polysaccharide
 484 extracts. *Carbohydrate Research*, *317*(1–4), 145–154.
- 485 https://doi.org/10.1016/S0008-6215(99)00071-3
- 486 Cordella, C. B. Y., & Bertrand, D. (2014). SAISIR: A new general chemometric
 487 toolbox. *TrAC Trends in Analytical Chemistry*, *54*, 75–82.
- 488 https://doi.org/10.1016/j.trac.2013.10.009
- 489 Edwards, S. L. (1976). An Investigation of the Vibrational Spectra of the Pentose
 490 Sugars. Lawrence University.
- 491 Englyst, H., Wiggins, H. S., & Cummings, J. H. (1982). Determination of the
 492 non-starch polysaccharides in plant foods by gas-liquid chromatography of
 493 constituent sugars as alditol acetates. *The Analyst*, *107*(1272), 307–318.
- 494 https://doi.org/10.1039/an9820700307
- Ferreira, D., Barros, A., Coimbra, M. A., & Delgadillo, I. (2001). Use of FT-IR
 spectroscopy to follow the effect of processing in cell wall polysaccharide
- 497 extracts of a sun-dried pear. *Carbohydrate Polymers*, 45(2), 175–182.
- 498 https://doi.org/10.1016/S0144-8617(00)00320-9
- Filippov, M. P., & Kohn, R. (1975). Determination of the esterification degree of
 carboxyl groups of pectin with methanol by means of infrared spectroscopy. *Chem. Zvesti*, 29(1), 88–91. Retrieved from
- 502 https://www.chempap.org/?id=7&paper=5409
- 503 Gnanasambandam, R., Proctor, A. (2000). Determination of pectin degree of
 504 esterification by diffuse reflectance. *Food Chemistry*, 68, 327–332.
 505 https://doi.org/10.1016/s0308-8146(99)00191-0
- 506 Granato, D., Santos, J. S., Escher, G. B., Ferreira, B. L., & Maggio, R. M. (2018). Use

- of principal component analysis (PCA) and hierarchical cluster analysis (HCA)
 for multivariate association between bioactive compounds and functional
 properties in foods: A critical perspective. *Trends in Food Science and Technology*, 72, 83–90. https://doi.org/10.1016/j.tifs.2017.12.006
- Jockusch, R. A., Kroemer, R. T., Talbot, F. O., Snoek, L. C., Çarçabal, P., Simons, J.
 P., ... Von Helden, G. (2004). Probing the Glycosidic Linkage: UV and IR
 Ion-Dip Spectroscopy of a Lactoside. *Journal of the American Chemical Society*, *126*(18), 5709–5714. https://doi.org/10.1021/ja031679k
- 515 Kacurakova, M., Capek, P., Sasinkova, V., Wellner, N., & Ebringerova, A. (2000).
 516 FT-IR study of plant cell wall model compounds: pectic polysaccharides and
- 517hemicelluloses. Carbohydrate Polymers, 43(2), 195–203.

518 https://doi.org/10.1016/S0144-8617(00)00151-X

- 519 Kačuráková, M., & Mathlouthi, M. (1996). FTIR and laser-Raman spectra of
 520 oligosaccharides in water: Characterization of the glycosidic bond. *Carbohydrate*521 *Research*, 284(2), 145–157. https://doi.org/10.1016/0008-6215(95)00412-2
- Kačuráková, M., Wellner, N., Ebringerová, A., Hromádková, Z., Wilson, R. H., &
 Belton, P. S. (1999). Characterisation of xylan-type polysaccharides and
 associated cell wall components by FT-IR and FT-Raman spectroscopies. *Food Hydrocolloids*, *13*(1), 35–41. https://doi.org/10.1016/S0268-005X(98)00067-8
- Kanou, M., Nakanishi, K., Hashimoto, A., & Kameokaj, T. (2005). Influences of
 monosaccharides and its glycosidic linkage on infrared spectral characteristics of
 disaccharides in aqueous solutions. *Applied Spectroscopy*, 59(7), 885–892.
 https://doi.org/10.1366/0003702054411760

530 Kyomugasho, C., Christiaens, S., Shpigelman, A., Van Loey, A. M., & Hendrickx, M.

- E. (2015). FT-IR spectroscopy, a reliable method for routine analysis of the degree of methylesterification of pectin in different fruit- and vegetable-based matrices. *Food Chemistry*, 176, 82–90.
- 534 https://doi.org/10.1016/j.foodchem.2014.12.033
- 535 Lan, W., Renard, C. M. G. C., Jaillais, B., Leca, A., & Bureau, S. (2020). Fresh,

- freeze-dried or cell wall samples: Which is the most appropriate to determine
 chemical, structural and rheological variations during apple processing using
 ATR-FTIR spectroscopy? *Food Chemistry*, *330*, 127357.
- 539 https://doi.org/10.1016/j.foodchem.2020.127357
- Le Bourvellec, C., & Renard, C. M. G. C. (2012). Interactions between polyphenols
 and macromolecules: Quantification methods and mechanisms. *Critical Reviews in Food Science and Nutrition*, 52(3), 213–248.
- 543 https://doi.org/10.1080/10408398.2010.499808
- Liu, X., Le Bourvellec, C., & Renard, C. M. G. C. (2020). Interactions between cell
 wall polysaccharides and polyphenols: Effect of molecular internal structure. *Comprehensive Reviews in Food Science and Food Safety*, 19(6), 3574–3617.
 https://doi.org/10.1111/1541-4337.12632
- Liu, X., Renard, C. M. G. C., Rolland-Sabaté, A., Bureau, S., & Le Bourvellec, C.
 (2021). Modification of apple, beet and kiwifruit cell walls by boiling in acid
 conditions: Common and specific responses. *Food Hydrocolloids*, *112*.
 https://doi.org/10.1016/j.foodhyd.2020.106266
- McCann, M. C., Hammouri, M., Wilson, R., Belton, P., & Roberts, K. (1992). Fourier
 transform infrared microspectroscopy is a new way to look at plant cell walls. *Plant Physiology*, *100*(4), 1940–1947. https://doi.org/10.1104/pp.100.4.1940
- Monsoor, M. A., Kalapathy, U., & Proctor, A. (2001). Determination of
 polygalacturonic acid content in pectin extracts by diffuse reflectance Fourier
 transform infrared spectroscopy. *Food Chemistry*, 74(2), 233–238.
 https://doi.org/10.1016/S0308-8146(01)00100-5
- Nikonenko, N. A., Buslov, D. K., Sushko, N. I., & Zhbankov, R. G. (2000).
 Investigation of stretching vibrations of glycosidic linkages in disaccharides and
 polysaccarides with use of IR spectra deconvolution. *Biopolymers (Biospectroscopy)*, 57(4), 257–262.
- 563 https://doi.org/10.1002/1097-0282(2000)57:4<257::AID-BIP7>3.0.CO;2-3
- 564 R Core Team. (2014). A Language and Environment for Statistical Computing. R

- 565 Foundation for Statistical Computing, 2. Retrieved from
 566 http://www.r-project.org
- 567 Reintjes, M., Musco, D. D., & Joseph, G. H. (1962). Infrared Spectra of Some Pectic
 568 Substances. *Journal of Food Science*, 27(5), 441–445.
- 569 https://doi.org/10.1111/j.1365-2621.1962.tb00124.x
- 570 Renard, C. M. G. C. (2005). Variability in cell wall preparations: Quantification and
 571 comparison of common methods. *Carbohydrate Polymers*, 60(4), 515–522.
 572 https://doi.org/10.1016/j.carbpol.2005.03.002
- 573 Renard, C. M. G. C., & Ginies, C. (2009). Comparison of the cell wall composition
 574 for flesh and skin from five different plums. *Food Chemistry*, *114*(3), 1042–1049.
 575 https://doi.org/10.1016/j.foodchem.2008.10.073
- Saeman, J. F., Moore, W. E., Mitchell, R. L., & Millett, M. A. (1954). Techniques for
 the determination of pulp constituents by quantitiative paper chromatography. *Tappi Journal*, *37*(8), 336–343.
- 579 Simonović, J., Stevanic, J., Djikanović, D., Salmén, L., & Radotić, K. (2011).
 580 Anisotropy of cell wall polymers in branches of hardwood and softwood: A
 581 polarized FTIR study. *Cellulose*, *18*(6), 1433–1440.
- 582 https://doi.org/10.1007/s10570-011-9584-1
- 583 Szymanska-Chargot, M., Chylinska, M., Kruk, B., & Zdunek, A. (2015). Combining
 584 FT-IR spectroscopy and multivariate analysis for qualitative and quantitative
 585 analysis of the cell wall composition changes during apples development.
- 586 *Carbohydrate Polymers*, *115*, 93–103.
- 587 https://doi.org/10.1016/j.carbpol.2014.08.039
- Szymanska-Chargot, M., & Zdunek, A. (2013). Use of FT-IR Spectra and PCA to the
 Bulk Characterization of Cell Wall Residues of Fruits and Vegetables Along a
 Fraction Process. *Food Biophysics*, 8(1), 29–42.
 https://doi.org/10.1007/s11483-012-9279-7
- 592 Watrelot, A. A., Le Bourvellec, C., Imberty, A., & Renard, C. M. G. C. (2013).
 593 Interactions between pectic compounds and procyanidins are influenced by

- methylation degree and chain length. *Biomacromolecules*, 14(3), 709–718.
 https://doi.org/10.1021/bm301796y
- Wiercigroch, E., Szafraniec, E., Czamara, K., Pacia, M. Z., Majzner, K., Kochan,
 K., ... Malek, K. (2017). Raman and infrared spectroscopy of carbohydrates: A
 review. Spectrochimica Acta Part A: Molecular and Biomolecular
 Spectroscopy, 185, 317–335. https://doi.org/10.1016/j.saa.2017.05.045
- 600 Wojdyło, A., Figiel, A., Lech, K., Nowicka, P., & Oszmiański, J. (2014). Effect of
- 601 Convective and Vacuum-Microwave Drying on the Bioactive Compounds, Color,
- and Antioxidant Capacity of Sour Cherries. *Food and Bioprocess Technology*.
- 603 https://doi.org/10.1007/s11947-013-1130-8