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Genotype, development and tissue-derived variation of cell-wall properties in the lignocellulosic energy crop Miscanthus

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PART OF A SPECIAL ISSUE ON PLANT CELL WALLS

⁵ Genotype, development and tissue-derived variation of cell-wall properties in the lignocellulosic energy crop *Miscanthus*

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• *Background and Aims* Species and hybrids of the genus *Miscanthus* contain attributes that make them front-runners among current selections of dedicated bioenergy crops. A key trait for plant biomass conversion to biofuels and biomaterials is cell-wall quality; however, knowledge of cell-wall composition and biology in *Miscanthus* species is limited. This study presents data on cell-wall compositional changes as a function of development and tissue type across selected genotypes, and considers implications for the development of miscanthus as a sustainable and renewable bioenergy feedstock.

• *Methods* Cell-wall biomass was analysed for 25 genotypes, considering different developmental stages and stem vs. leaf compositional variability, by Fourier transform mid-infrared spectroscopy and lignin determination. In addition, a *Clostridium phytofermentans* bioassay was used to assess cell-wall digestibility and conversion to ethanol.

• *Key Results* Important cell-wall compositional differences between miscanthus stem and leaf samples were found to be predominantly associated with structural carbohydrates. Lignin content increased as plants matured and was higher in stem tissues. Although stem lignin concentration correlated inversely with ethanol production, no such correlation was observed for leaves. Leaf tissue contributed significantly to total above-ground biomass at all stages, although the extent of this contribution was genotype-dependent.

• *Conclusions* It is hypothesized that divergent carbohydrate compositions and modifications in stem and leaf tissues are major determinants for observed differences in cell-wall quality. The findings indicate that improvement of lignocellulosic feedstocks should encompass tissue-dependent variation as it affects amenability to biological conversion. For gene–trait associations relating to cell-wall quality, the data support the separate examination of leaf and stem composition, as tissue-specific traits may be masked by considering only total above-ground biomass samples, and sample variability could be mostly due to varying tissue contributions to total biomass.

Key words: *Miscanthus*, biofuels, plant cell wall, Fourier transform mid-infrared spectroscopy, FTIR, lignin, fermentation, bioenergy, recalcitrance, lignocellulose, biomass, development, carbohydrates.

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INTRODUCTION

Plant biomass represents an abundant resource of renewable energy in the form of cell-wall polysaccharides. Dedicated energy crops as well as dual-purpose food and energy cultivars from the Panicoideae clade, which includes *Zea mays* (maize), *Miscanthus* spp. (miscanthus), *Sorghum bicolor* (sorghum), *Saccharum* spp. (sugarcane) and *Panicum virgatum* (switchgrass), are C₄ grasses that generate high yields of biomass (Feltus and Vandenbrink, 2012; van der Weijde *et al.*, 2013).

- 50 Among these, miscanthus represents one of the most promising dedicated second-generation bioenergy crops under development (Carroll and Somerville, 2009). Native to East Asia, members of the genus *Miscanthus* are perennial, rhizomatous plants, which remobilize nutrients to the rhizome during senes-
- 55 cence to ensure regrowth of the crop in the subsequent season (Robson *et al.*, 2012). Consequently, miscanthus is typically harvested during winter or early spring when nutrients have been translocated from above-ground tissues to rhizomes, thus providing a number of environmental advantages over annuals

as bioenergy crops, including lower requirements for fertilizer, reduced soil erosion and the potential for soil carbon sequestration (Clifton-Brown *et al.*, 2013). Furthermore, several varieties of miscanthus give high yields in cool climates, unusual within the C₄ grasses, making miscanthus a potentially viable and sustainable energy crop over a wide range of diverse geographical zones (Purdy *et al.*, 2013). Of the several identified miscanthus species, the most commonly investigated are *Miscanthus sinensis*, *Miscanthus sacchariflorus* and the vigorous but sterile triploid hybrids between the two, of which *Miscanthus* × *giganteus* is the most widely cultivated variety (Heaton *et al.*, 2008; Dwiyanti *et al.*, 2013; Liu *et al.*, 2013).

Most of the potential energy in lignocellulosic biomass is locked within secondary cell walls, a heterogeneous mix of predominantly cellulose, xylan and lignin polymers that interact to assemble a complex and dense matrix (McCann and Carpita, 2008; Chundawat *et al.*, 2011). The relative abundances and interactions among the polymers dictate biomass recalcitrance to saccharification (i.e. amenability to deconstruction to release fermentable sugars). Therefore, one of the key traits for

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the processing of plant biomass to produce biofuels and bioma-

¹²⁰ terials is secondary cell-wall quality (Himmel *et al.*, 2007; DeMartini *et al.*, 2013).

Lignin, one of the secondary cell-wall components, consists of a complex aromatic heteropolymer composed of varying percentages of three phenylpropanoid units: p-hydroxyphenyl (H),

- guaiacyl (G) and syringyl (S) (Fukushima and Dehority, 2000; Bonawitz and Chapple, 2010). The concentration of lignin, its composition and the manner in which it binds holocellulose within the cell-wall matrix is often seen as an exacerbating factor of cell-wall recalcitrance to enzymatic deconstruction
 (Vanholme *et al.*, 2010; Hodgson *et al.*, 2011; Ding *et al.*,
- 150 (Validonie et al., 2010, Hodgson et al., 2011, Dilg et al., 2012). However, the extent of this effect is not always consistent in literature reports and therefore recalcitrance should not be attributed solely to the presence of lignin (Grandis et al., 2014). In miscanthus, for instance, contrasting influences of
- lignin content on enzymatic hydrolysis have been reported (Lygin *et al.*, 2011; Zhang *et al.*, 2012). Other factors also influence enzymatic cell-wall hydrolysis, such as cellulose crystallinity (Hall *et al.*, 2010) and hemicellulosic xylan polysaccharides (DeMartini *et al.*, 2013), although the impact of these compositional and architectural features will vary depending on plant

species, developmental stage and tissue type. Despite the importance of optimizing miscanthus cell-wall

properties to improve its usefulness as a sustainable and economically viable bioenergy crop, there are surprisingly few reported

- studies on the cell-wall composition and biology of this genus (Slavov *et al.*, 2013*a*) and, to our knowledge, no reports addressing cell-wall changes as a function of tissue type and development.
- To investigate the chemical, structural and biological features of miscanthus biomass as a lignocellulosic feedstock, and to unveil how these characteristics vary among different genotypes, we embarked on an in-depth cell-wall analysis of 25 miscanthus genotypes from a larger replicated field trial comprising 244 genotypes. Several earlier studies on the entire field trial have focused on a diverse set of physiological and agronomical
- 155 focused on a diverse set of physiological and agronomical traits, including senescence (Robson *et al.*, 2012), flowering time (Jensen *et al.*, 2011), and canopy duration and leaf and stem morphology (Robson *et al.*, 2013). In addition, cell-wall composition of the full set of genotypes was previously
- 160 determined using gravimetric analytical methods in combination with near-infrared reflectance spectrophotometry (NIRS)-based calibration models (Allison *et al.*, 2011). Extending the level of detail of the latter study, we used a multidimensional approach, considering different developmental stages, stem vs. leaf com-
- positional variability, Fourier transform mid-infrared spectroscopy (FTIR) and acetyl bromide lignin determination on isolated cell-wall biomass. Furthermore, we employed a bioassay for the determination of cell-wall digestibility as a function of the ethanol yielded after fermentation with *Clostridium phyto-*
- 170 *fermentans*, an anaerobic soil bacterium that can convert a wide range of cell-wall components to ethanol, without the addition of exogenous cellulases and xylanases (Warnick *et al.*, 2002; Lee *et al.*, 2012*b*). Finally, we consider the possible implications of our findings in terms of future research strategies aimed at devel-
- 175 oping miscanthus into a sustainable energy crop by means of broadening our understanding of cell-wall compositional features, and impacts on biorefining.

MATERIALS AND METHODS

Plant material

Twenty-five genotypes of Miscanthus were selected from a spaced field trial of 244 accessions established in 2004 near Aberystwyth, UK (52·437848°N, 4·026688°W) described by Allison et al. (2011). Briefly, the trial field is on a WSW 7 % sloping field, relatively exposed to southerly and westerly 185 winds. The trial is organized into four randomized blocks, with longer block dimensions orientated perpendicularly to the main slope, and each surrounded by a dense guard perimeter of a commercially available variety of M. \times giganteus. The soil is characterized by a pH ranging from 5.1 to 6.3, and consists of a 190 stony seasonally waterlogged loam overlying shale, with the stone fraction estimated at 50 % of the soil mass in the 0-40 cm layer. Genotypes were selected to represent a wide range of compositional variability, estimated by gravimetric measurements of neutral detergent fibre, acid detergent fibre 195 and acid detergent lignin in bulked plant tissue samples (Allison et al., 2011). Priority was also given to genotypes included in genome-wide association studies (Slavov et al., 2013b). For each genotype, a single tiller of length equal or greater than three-quarters of the plant's total height (excluding 200 rhizome and inflorescence when present) was selected randomly and collected from three of the four replicate plots. Samples were collected at three time points during the 2012-2013 growing season. The time points corresponded to three developmental stages: 10 weeks after first shoot emergence, when the plants 205 were actively growing (AG); 18 weeks after emergence, a stage when the plants had mostly ceased their growth (peak biomass, PB); and at 42 weeks after emergence, when the plants had completely senesced (senesced stage, SS). Immediately after collection, the tillers were photographed, mea-210 sured and left at -20 °C overnight, before being freeze-dried. Once dry, stem and leaf tissue (including sheath) were separated and weighed, leaf contribution was recorded as percentage of total biomass dry weight, and individual tissues were ground to a particle size in the range 0.18 - 0.85 mm (mesh sizes 80 and 215 20). By the end of the growing season, 18 samples had been collected for each of the 25 selected lines (3 developmental stages \times 3 biological replicates \times 2 tissues).

Cell wall biomass preparation

All compositional analyses and the C. phytofermentans bioassay were carried out on purified cell wall, which was prepared following a procedure adapted from Foster et al. (2010). For each sample, approx. 1 g of ground plant biomass was extracted se-225 quentially as follows: with 30 mL ethanol, first for 12 h and then twice more for 30 min in a shaking incubator set at 40 $^{\circ}$ C/ 150 r.p.m.; three times with 20 mL chloroform/methanol **Q3** (1:1 v/v), for 30 min incubation at 25 °C and 150 r.p.m.; and finally, three times with 15 mL acetone, for 30 min, at $25 \,^{\circ}\text{C/}$ 230 150 r.p.m. Between each step of the extraction, the material was collected by centrifugation at 8000 r.p.m./10 min and the supernatants were discarded. Following the third acetone step, the samples were left to dry overnight in a fume hood. The 235 dried, solvent-extracted biomass was then re-suspended in 15 mL of 0.1 m sodium acetate buffer (pH 5.0) and heated to

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80 °C/20 min to induce starch gelatinization followed by cooling on ice. Subsequently, samples were centrifuged and supernatants were discarded, after which the resulting pellet was washed twice

with 30 mL deionized water, with resuspension, centrifugation 240 and supernatant removal being performed for each wash. Sodium azide was added at 0.0002 % (w/v) to inhibit microbial growth, and starch was removed by incubation with type-I porcine α -amylase (Sigma-Aldrich, St Louis, MO, USA; 47

245 units per 100 mg cell wall) in 0.1 m ammonium formate buffer (pH 6.0) at 25 °C/110 r.p.m. After 48 h, digestion was terminated by heating to 95 °C/15 min and samples were cooled on ice. The destarched cell-wall preparations were then washed three times in 30 mL deionized water and twice with 20 mL acetone, with centrifugation and supernatant removal, before being

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freeze-dried.

Fourier transform mid-infrared spectroscopy

- FTIR was performed on the prepared cell-wall biomass for all 255 miscanthus samples (25 lines \times 3 time points \times 2 tissues \times 3 plant replicates). Duplicate spectra were collected by attenuated total reflectance (ATR) in the range 4000–600 cm^{-1} using an Equinox 55 FTIR spectrometer (Bruker Optik, Ettlingen, Germany) equipped with a Golden Gate ATR accessory 260 (Specac, Slough, UK). Spectra were averaged over 32 scans at a
- resolution of 4 cm⁻¹ and corrected for background absorbance by subtraction of the spectrum of the empty ATR crystal. Absorbance spectra were converted to text files in Opus (v. 5.0; Bruker Optik), imported into MatLab (v. R2010b; MathWorks, 265
- Natick, MA, USA) and averaged. Full spectra, or fingerprint region spectra (1900-800 cm⁻¹), were transformed according to the Savitzky-Golay algorithm (order: 3; window: 15 pt), to improve peak resolution, and mean centre normalized (mean 0,
- s.d. 1) prior to principal components analysis (PCA) using the 270 Eigenvector PLS Toolbox (v. 7.0.3; Eigenvector Research, Wenatchee, WA, USA) to investigate the underlying relationships between the spectra.

275 Lignin measurement

Acetyl bromide lignin was determined in triplicate for all of the miscanthus samples (25 lines \times 3 time points \times 2 tissues \times 3 plant replicates) following the general procedures described by Foster et al. (2010) and Fukushima and Hatfield (2004), with some modifications, described as follows. Approximately 7.0 mg of the cell-wall samples was weighed into 10-mL Pyrex glass tubes fitted with polypropylene caps. For lignin solubilization, 500 µL of freshly prepared 25 % (v/v) acetyl bromide solution in glacial acetic acid was added to the samples, the tubes 285 were capped and placed in a heating block set at 50 °C for 2 h, after which the tubes were mixed using a vortex mixer every 15 min up to a total incubation time of 3 h. Following digestion, the tubes were cooled on ice and the contents of each were 290 diluted by the addition of 2000 μ L of 2 m NaOH. A further addition of 350 µL of 0.5 m hydroxylamine hydrochloride to each tube ensured the decomposition of polybromide ions (Monties, 1989). After vortex mixing, the final volume was adjusted to

10 mL with glacial acetic acid. The tubes were recapped, mixed by inversion and centrifuged to produce a particulate-free super-295 natant, and 200 µL of each sample was transferred to UV-transparent 96-well plates (UV-Star; Greiner Bio-One, Gloucestershire, UK). Absorbance at 280 nm was measured with a plate reader (µQuant; Bio-Tek Instruments, Winooski, VT, USA) using KC4 software (v. 3.3; Bio-Tek). An assay control sample of a standard cell-wall preparation was included 300 in all batches of the lignin assay as an internal standard. Additionally, negative controls containing no cell-wall material were included and their absorbance at 280 nm was set as absorbance baseline. A specific absorption coefficient (SAC) of 17.78 g^{-1} L cm⁻¹ has been reported for purified HCl-dioxane lignin from miscanthus samples (Lygin et al., 2011) and this was used to calculate the percentages of lignin in the cell-wall biomass samples as dry weight using the following equation: ABSL% = $(A_{280}/(SAC \times PL)) \times (V_R/W_S) \times 100\%$, where ABSL% is the acetyl bromide-soluble lignin percentage content, A_{280} is the ab- 310 sorption reading at 280 nm, PL is the pathlength determined for the 96-well microplates with a volume of 200 µL per well used during the analysis (0.556 cm), $V_{\rm R}$ is the reaction volume (litres) and $W_{\rm S}$ is the sample weight (g). Note that the ABSL method employed also measures ester-linked hydroxycinnamic acids 315 and it has been reported that these act as synthetic precursors and form an integral part of the lignin macromolecule (Ralph et al., 1994; Ralph, 2010; Tobimatsu et al., 2012).

C. phytofermentans bioassay of biomass digestibility

Ethanol yield analysis was performed as described previously (Lee et al., 2012a, b). C. phytofermentans strain ISDg (ATCC 700394) was cultured in a defined medium, MQM5.1, prepared as follows: $2.0 \text{ g L}^{-1} \text{ NaH}_2\text{PO}_4$, $10.0 \text{ g L}^{-1} \text{ K}_2\text{HPO}_4$, 1.0 g L^{-1} 325 $(NH_4)_2SO_4$, 1.0 g L⁻¹ L-cysteine hydrochloride monohydrate, 20 mL L⁻¹ XT solution (5.0 g L⁻¹ xanthine and 5.0 g L⁻¹ thymine in 0.06 m NaOH), 10 mL L^{-1} AA1 solution (5.0 g L^{-1} of each of the following amino acids: alanine, arginine, histidine, isoleucine, leucine, methinonine, proline and valine), and 10 mL 330 L^{-1} trace element solution (Balch *et al.*, 1979), resazurin (1 mg L⁻¹ trace element solution (Balch *et al.*, 1979), resizutin (Fing L⁻¹), which was added as an oxidation/reduction indicator. After autoclaving, 10 mL L⁻¹ CPV3 solution (20 mg L⁻¹ *p*-aminobenzoic acid, 1 mg L⁻¹ biotin, 30 mg L⁻¹ folinic acid, 80 mg L⁻¹ nicotinamide, 5 mg L⁻¹ pantethine, 2 mg L⁻¹ pyridox-335 al hydrochloride, 30 mg L⁻¹ riboflavin and 10 mg L⁻¹ thiamine) was added. The C. phytofermentans inoculum was initially grown in MQM5.1 with 3 g L^{-1} cellobiose as a carbon source using the anaerobic techniques described by Hungate (1969). Incubations were carried out in 10-mL volumes in 18×180 -mm tubes 340 sealed with neoprene caps.

For the biological conversion quality assay, the previously purified leaf and stem cell-wall biomass from the three replicates of the 25 miscanthus genotypes at PB developmental stage were analysed. Approximately 20-mg portions of each sample were 345 weighed in triplicate into autoclavable 2.2-mL polypropylene 96-well plates (Axygen Scientific, Union City, CA, USA), 0.92 mL of MQM5.1 media was added, and plates were sealed and autoclaved. Subsequently, 0.01 mL of the CPV3 solution and 0.01 mL of the prepared C. phytofermentans inoculum 350 was added to each well, and the samples were incubated without shaking at 37 °C/72 h. After incubation, the plates were centrifuged and a volume of 1.0 mL of each sample supernatant was collected and filtered through a 0.22-µm syringe filter unit (Millipore Corp., Billerica, MA, USA) and 5.0 µL of

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- ³⁵⁵ each sample was analysed by high-performance liquid chromatography (HPLC). The HPLC system (Waters Corp., Milford, MA, USA) was equipped with a carbohydrate analysis column (7.8×150 mm IC-Pak Ion Exclusion; Waters Corp.) and a refractive-index detector. The column was operated at
- 360 30 °C with 0.0025 M H₂SO₄ as the running buffer at a flow rate of 0.7 mL min⁻¹. The retention time for ethanol (17.84 \pm 0.02 min) was determined using a commercial mix (Fuel Ethanol Residual Saccharides Mix; catalogue number 48468-U; Sigma-Aldrich) containing glycerol, glucose, maltotriose,
- 365 maltose monohydrate, lactic acid, acetic acid, dextrin and ethanol. Standards were analysed at the beginning, middle and end of every distinct HPLC analysis-to ensure accuracy and precision of measurements.
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Statistical analysis

All calculations for descriptive statistics, analyses of variance and Tukey's range tests were performed using the statistical software Statistica (v. 8.0; StatSoft, Tulsa, OK, USA) at a 5 % significance level. For the leaf percentage dataset, analysis of variance (ANOVA) was used to test for the effect of genotype (25 levels) and developmental stage (three levels), and correlations were determined following natural logarithm transformation of the data, which showed a skewed distribution due to the exponential nature of tissue growth. With respect to lignin content and ethanol yield, the effect of tissue type (two levels) was also tested in addition to genotype and development

factors. Tukey's tests were used for multiple comparisons between factor levels. Effect sizes were calculated as eta-squared 4 statistics: $\eta^2 = SS_{effect}/SS_{total}$ (Cohen, 1973; Levine and Hullett, 2002), where SS is the sum of squares.

RESULTS

Morphological diversity within Miscanthus genotypes

The plant material used in this study included 25 miscanthus genotypes, with varying ploidy and represented two miscanthus species, M. sinensis and M. sacchariflorus, an inter-specific 425 hybrid M. \times giganteus, and other hybrids with different estimated admixture proportions of *M. sinensis* and *M. sacchariflorus* (Table 1). The mean contribution of leaf material (leaf blade and sheath) to total dry biomass among the genotypes was 63.8%(ranging from 42.2 to 80.4 %) at the AG stage, 55.6 % (36.3-430 78.8%) for PB stage and 36.1% (15.8-63.2%) for SS (Table 1). ANOVAs used to assess the effects of genotype, and developmental stage, on the percentage of leaf contribution to total biomass indicated that both factors were statistically significant, with large effect sizes: $\eta_{genotype}^2 = 0.35$ and $\eta_{developmental stage}^2 = 0.51$ (P < 0.0001). Variation in leaf percentage between the 435 three replicates of each genotype was not significant (P =0.4959), and Tukey's tests indicated that all three developmental stages were significantly different from each other. The interaction between genotype and development stage effects was not signifi-440 cant (P = 0.5858), indicating that although leaf percentage of

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TABLE I.	Description	of the 25	miscanthus	genotypes	used in	this study
				() · · · · / · · ·		

				Leaf percentage		445
Gen	notype Species	Ploidy	Active growth	Peak biomass	Senesced	
390 gig(01 $M. \times giganteus$	3 <i>n</i>	44.53 + 5.82	36.45 + 0.49	16.07 + 0.76	
gig($M. \times giganteus$	3 <i>n</i>	42.30 + 3.32	36.65 + 3.32	17.87 + 3.13	450
gig($M. \times giganteus$	4n	42.23 ± 1.00	36.25 ± 0.21	15.77 ± 5.20	
hyb	55 % M. sinensis; 45 % M. sacch	ariflorus* 2n	55.03 ± 3.35	48.60 ± 1.56	26.33 ± 2.91	
hyb	02 72 % M. sinensis; 28 % M. sacch	ariflorus 2n	67.67 ± 3.36	45.10 ± 6.93	36.63 ± 3.87	
hyb	64 % M. sinensis; 36 % M. sacch	ariflorus 3n	53.20 ± 6.92	36.55 ± 0.49	22.07 ± 1.45	
395 sac(01 M. sacchariflorus	2 <i>n</i>	53.20 ± 1.83	49.25 ± 1.91	32.07 ± 6.20	
sin0	01 M. sinensis	2n	68.17 ± 1.10	59.40 ± 7.07	38.57 ± 2.57	455
sin0	02 M. sinensis	2 <i>n</i>	61.80 ± 4.77	50.85 ± 6.72	31.40 ± 1.85	
sin0	03 M. sinensis	2 <i>n</i>	68.13 ± 22.13	60.55 ± 17.61	46.17 ± 21.27	
sin0	04 M. sinensis	2 <i>n</i>	74.27 ± 1.78	65.55 ± 4.88	34.23 ± 0.85	
sin0	05 M. sinensis	2 <i>n</i>	64.13 ± 4.43	56.85 ± 0.64	37.10 ± 3.30	
400 sin0	06 M. sinensis	2 <i>n</i>	71.57 ± 3.01	62.70 ± 0.14	52.67 ± 4.62	
sin0	07 M. sinensis	2 <i>n</i>	66.87 ± 2.01	56.60 ± 1.13	28.87 ± 2.51	460
sin0	08 M. sinensis	2 <i>n</i>	69.67 ± 10.21	63.35 ± 0.21	37.40 ± 1.28	100
sin0	09 M. sinensis	3 <i>n</i>	63.47 ± 1.01	59.70 ± 2.55	36.57 ± 1.52	
sin1	10 M. sinensis	2 <i>n</i>	66.97 ± 1.54	52.35 ± 2.90	36.43 ± 2.25	
sin1	11 M. sinensis	2 <i>n</i>	70.17 ± 0.76	60.95 ± 1.63	31.00 ± 12.91	
405 sin1	12 M. sinensis	2 <i>n</i>	76.03 ± 9.49	66.65 ± 11.81	59.87 ± 16.37	
sin1	13 M. sinensis	2n	80.43 ± 2.50	78.75 ± 0.78	63.17 ± 11.55	465
sin1	14 M. sinensis	2n	69.73 ± 2.65	64.50 ± 0.71	50.63 ± 2.75	405
sin1	15 M. sinensis	2n	60.50 ± 1.70	54.80 ± 6.08	31.70 ± 1.60	
sin1	16 M. sinensis	2n	66.63 ± 4.07	61.80 ± 4.81	39.47 ± 5.20	
sin1	17 M. sinensis	2n	67.37 ± 8.71	62.45 ± 16.48	38.27 ± 9.72	
410 sin1	18 M. sinensis	2n	70.90 ± 7.10	62.70 ± 0.99	41.93 ± 7.09	
Ove	erall		$63{\cdot}80 \pm 10{\cdot}14$	55.57 ± 10.93	36.09 ± 12.12	470

Values for leaf percentage of total biomass dry weight are expressed as mean \pm s.d. for the three replicated plants at the three developmental stages for each genotype.

*M. sinensis/M. sacchariflorus admixture proportions determined from single-nucleotide polymorphism data (Slavov et al., 2013b).

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total biomass varies throughout development, this variation is not significantly different between genotypes. The percentage of leaf

475 contribution to total biomass was negatively correlated with tiller length (r = -0.77, P < 0.0001) and total dry weight (r = -0.59; P < 0.0001; Fig. 1A, B). Box-and-whisker plots of the distribution of leaf percentage, tiller length and tiller weight (Fig. 1C) show that the relative contribution of leaf tissue to total biomass

- 480 decreased as plants matured. By contrast, tiller dry weight increased during plant growth to a maximum at peak biomass, and tiller length continuously increased until senescence, but in both cases the rate of change decreased as plants started to senesce. Mean leaf contribution to total biomass plotted against
- tiller length and weight, at the various developmental stages, shows that the genotypes used in this study differ in several allometric traits (Fig. 2). Throughout development hybrid genotypes ranked high in terms of tiller length and weight, but showed low leaf contributions to total biomass. This trend was predominant in $M. \times giganteus$ genotypes, whereas the other three hybrid

genotypes displayed less extreme traits. The *M. sinensis* genotypes included in this study showed a broad range in leaf contribution to total biomass, whereas the *M. sacchariflorus* genotype showed a tendency to fall between the *M. sinensis* genotypes and the other 535 hybrids.

Fourier transform mid-infrared spectroscopy

FTIR spectroscopy allowed us to investigate cell-wall composition in miscanthus stem and leaf samples, as well as to identify the major compositional shifts in each of these tissues during development. Comparison of the spectra for stem and leaf samples of the 25 genotypes at each developmental stage showed differences in the relative absorbance of the individual bands. However, they were too numerous and complex for detailed visual interpretation, and PCA was employed as an exploratory approach to identify the most distinctive features of the collected spectra. Following PCA, ten spectral bands



FIG. 1. Morphological characterization of 25 miscanthus genotypes at three developmental stages. Correlation of the natural logarithms of (A) tiller length and (B) total dry weight with percentage leaf contribution to total dry weight biomass for the 25 genotypes at three developmental stages (*r*, Pearson correlation coefficient). (C) Distribution of the transformed measurements as boxplots.



 ⁶³⁰ FIG. 2. Graphical presentation of the mean leaf percentage of total biomass plotted against mean tiller length and tiller dry weight for the 25 miscanthus genotypes at three developmental stages.
 ⁶⁹⁰

were detected as the main discriminant principal component (PC) loadings in the fingerprint region of the spectra (1900– 800 cm^{-1} ; Fig. 3A, B). The attribution of spectral areas to their corresponding cell-wall components was made according to the literature (Supplementary Data Table S1). Bands associated

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with cellulose were found at 1159 cm⁻¹ (d), 1061 cm⁻¹ (f), 1038 cm⁻¹ (g) and 993 cm⁻¹ (i) (Marry *et al.*, 2000; Wilson *et al.*, 2000; Oh *et al.*, 2005; McCann *et al.*, 2007; Schulz and Baranska, 2007; Adapa *et al.*, 2009; Gwon *et al.*, 2010; Matos *et al.*, 2013; Abidi *et al.*, 2014); pectin-associated loadings at 1746 cm⁻¹ (a), 1105 cm⁻¹ (e), 1017 cm⁻¹ (h) and 951 cm⁻¹

(j) (Séné *et al.*, 1994; Coimbra *et al.*, 1999; Kačuráková *et al.*, 2000; Wilson *et al.*, 2000; McCann *et al.*, 2001; Alonso-Simón *et al.*, 2004); and discriminant bands associated with S lignin at 1321 cm⁻¹ (b) and 1234 cm⁻¹ (c) (Labbé *et al.*, 2005; Gorzsás *et al.*, 2011; Zhou *et al.*, 2011).

Three PCA models were created, with the first one including all collected spectra (Fig. 3C). In this model, the first four PCs

accounted for nearly 84 % of the variance in the spectral data set, of which PC1 captured 41.25 %. No clustering was detected concerning the various miscanthus species; however, two clear clusters were observed along PC1 comprising spectra from 695 stem and from leaf tissue. The loadings of PC1 for this model (Fig. 3F) showed that differences in four regions (designated a, c, g and j; as described above) of the FTIR spectra were the main contributors to the differential clustering of stem and leaf samples. One prevalent positive loading, located at 1234 cm^{-1} 700 (c), coincides with a band frequently associated with S units in core lignin. However, the other three main loadings of PC1 overlapped spectral regions typically associated with structural carbohydrate in lignocellulosic samples: positive peaks at 1746 cm^{-1} (a) and 1038 cm^{-1} (g), and a negative peak at 951 cm^{-1} 705 (i). This indicates that PC1 is mostly correlated with portions of the spectra associated with carbohydrates. Further PCA models were created after the spectral data had been split into separate subsets comprising each tissue type: leaf (Fig. 3D)



⁶⁰ F1G. 3. Mean FTIR spectra of (A) leaf and (B) stem samples of 25 miscanthus genotypes at three developmental stages in the range 1900–800 cm⁻¹. Plot of principal component one (PC1) and principal component two (PC2) scores for (C) all samples, (D) stem samples and (E) leaf samples. PC1 loading plot for (F) all samples and (G) stem samples. Spectral bands: a, 1745 cm⁻¹; b, 1325 cm⁻¹; c, 1230 cm⁻¹; d, 1159 cm⁻¹; e, 1105 cm⁻¹; f, 1060 cm⁻¹; g, 1037 cm⁻¹; h, 1017 cm⁻¹; i, 993 cm⁻¹; j, 950 cm⁻¹. Abbreviations: AG, active growth; PB, peak biomass; SS, senesced stage.

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and stem (Fig. 3E). For leaf samples, the first five PCs captured slightly more than 83 % of the total variance, but no clear clusters could be discerned along any of the PC axes (shown for PC1 and

- PC2 in Fig. 3D). In contrast, analysis of the stem spectral data (the 830 first four principal components accounted for almost 85 % of the variance) detected two distinctive clusters correlating to developmental stage along PC1 (Fig. 3E): one consisting of stem cell-wall samples from actively growing plants, and 835 another of overlapping stem samples collected at peak biomass and after senescence. For this PCA model, six spectral regions featured prominently in the loadings for PC1 (Fig. 3G). These loadings, which captured 49.31 % of the variance in stem spectral data, showed that this principal component is mostly correlated with spectral regions attributed to cell-wall polysacchar-840
- ide components: positively at 1746 cm⁻¹ (a), 1017 cm⁻¹ (h) and 993 cm⁻¹ (i); and negatively at 1159 cm⁻¹ (d), 1105 cm⁻¹ (e) and 1061 cm⁻¹ (f). In addition, it was also evident that the bands at 1321 cm⁻¹ (b) and 1234 cm⁻¹ (c), associated with S lignin mono-845
- mers, were perceptible negative loadings, thus suggesting higher amounts of S lignin in mature stem tissues.

Acetyl bromide lignin content

850 Lignin content was measured as ABSL% of cell-wall biomass dry weight (Table 2). The mean lignin content of the 25 selected genotypes increased in both tissues as plants matured, and ranged from 18.3 % in leaf tissue at AG stage to 23.7 % in stem tissue at SS stage. Additionally, lignin content was typically higher in stem samples than in leaf samples of the 855 same genotype.

The statistical significance of development, tissue and genotype effects was confirmed by ANOVA (P < 0.0001 for all three factors), and Tukey's tests showed distinction between AG, PB and SS, and between stem and leaf. Furthermore, the following effect sizes were determined: $\eta^2_{developmental stage} = 0.50$, $\eta^2_{tissue} = 0.10$ and $\eta^2_{genotype} = 0.06$. The lignin content variation 890 between the three plant replicates of each genotype (P =0.5553), the interaction between genotype and developmental stage (P = 0.88558) and the interaction genotype \times developmental stage \times tissue (P = 0.2056) were all not significant. By 895 contrast, the interactions of genotype \times tissue and developmental stage \times tissue were both significant (P < 0.0001). In light of these results, the importance of genotype and tissue on lignin content was assessed at each developmental stage individually. The resulting ANOVA showed that tissue was the only factor 900 that had a significant effect at the developmental stages considered (P < 0.0001 at each of the three developmental stages). Genotype had a significant effect on ABSL% in samples collected during AG (P = 0.0002) and during PB (P = 0.0052), but not in senesced samples (P = 0.6215). The interaction 905 between genotype and tissue was not significant at the actively growing stage (P = 0.2214), but was significant during peak biomass (P = 0.0005) and senescence (P = 0.0334). All these results suggest that although genotype has a significant effect on lignin content, its influence decreases over development, 910 until it has no significant effect on lignin concentration in samples collected during senescence. This decrease in the relevance of genotype is supported by a reduction of its effect size throughout development: $\eta^2_{AG} = 0.30$, $\eta^2_{PB} = 0.13$ and $\eta_{SS}^2 = 0.12.$ 915

TABLE 2. Acetyl bromide lignin percentage of cell-wall biomass dry weight

860		Active	growth	Peak b	iomass	Sene	esced
G	Genotype	Leaf	Stem	Leaf	Stem	Leaf	Stem
g	rig01	17.27 ± 1.38	18.72 ± 0.67	19.61 ± 0.86	22.75 ± 0.73	20.48 ± 0.84	24.45 ± 1.22
g g	tig02	19.48 ± 0.85	19.87 ± 1.21	18.71 ± 0.15	23.39 ± 0.98	20.80 ± 0.70	24.85 ± 0.78
865 g	tig03	18.19 ± 0.40	19.74 ± 1.48	18.55 ± 0.59	24.05 ± 1.11	21.36 ± 2.88	24.44 ± 0.93
ĥ	yb01	17.08 ± 0.23	18.03 ± 1.09	19.03 ± 0.71	21.72 ± 1.80	22.18 ± 1.57	23.55 ± 0.66
h	yb02	17.43 ± 0.33	19.76 ± 1.22	19.17 ± 0.15	23.15 ± 1.39	22.51 ± 0.45	23.77 ± 0.62
h	iyb03	17.91 ± 1.72	20.07 ± 0.89	18.52 ± 0.45	23.64 ± 0.46	20.05 ± 0.51	24.26 ± 0.62
sa	ac01	17.38 ± 0.78	17.18 ± 1.23	19.36 ± 1.38	20.92 ± 0.13	22.02 ± 2.31	23.25 ± 1.16
si	in01	19.70 ± 1.50	19.76 ± 0.69	21.47 ± 1.00	21.72 ± 0.28	23.22 ± 1.92	22.98 ± 0.04
⁸⁷⁰ si	in02	18.63 ± 1.88	20.41 ± 1.73	20.69 ± 0.65	21.54 ± 1.30	23.59 ± 3.03	23.88 ± 0.81
si	in03	18.07 ± 1.22	18.18 ± 3.11	19.24 ± 1.05	19.90 ± 2.45	23.80 ± 0.87	22.56 ± 2.19
si	in04	18.68 ± 0.53	20.00 ± 0.37	19.45 ± 1.27	20.50 ± 1.30	24.14 ± 2.33	22.32 ± 0.39
si	in05	18.88 ± 1.60	19.45 ± 0.81	19.31 ± 1.37	22.55 ± 0.58	23.82 ± 2.48	23.76 ± 0.87
si	in06	18.95 ± 0.98	17.68 ± 1.17	20.50 ± 1.32	21.62 ± 1.16	22.22 ± 2.70	22.86 ± 0.43
si	in07	18.63 ± 1.40	19.35 ± 2.22	19.85 ± 1.54	24.33 ± 1.28	24.01 ± 2.19	22.93 ± 0.75
875 si	in08	17.95 ± 0.66	18.09 ± 2.15	19.43 ± 1.43	21.34 ± 0.35	22.81 ± 2.34	23.01 ± 1.10
si	in09	18.46 ± 1.38	21.51 ± 0.65	19.24 ± 0.99	22.76 ± 0.32	24.41 ± 2.97	24.05 ± 0.96
si	in10	19.73 + 0.68	19.74 + 1.63	20.83 + 1.46	22.85 + 0.70	23.11 + 1.22	24.10 + 0.54
si	in11	18.30 + 0.53	19.13 + 2.61	19.33 + 1.12	22.13 + 0.87	22.22 + 2.05	24.06 + 1.27
si	in12	18.03 + 1.50	17.79 + 2.21	18.74 + 1.12	21.82 + 2.67	21.68 + 2.39	23.70 + 1.22
si	in13	17.23 + 0.35	16.44 + 0.14	19.41 + 0.83	18.62 + 3.08	21.12 + 1.39	22.93 + 0.53
880 si	in14	17.28 + 1.43	19.39 + 2.10	19.10 + 0.16	21.99 + 1.66	22.12 + 0.22	24.54 + 0.78
si	in15	19.96 + 0.14	21.26 + 1.56	20.77 + 1.03	22.49 + 1.60	23.39 + 0.97	24.56 + 0.94
si	in16	17.59 + 1.13	19.93 + 0.76	18.47 + 0.67	22.88 + 1.66	21.89 + 1.38	23.92 + 0.03
si	in17	17.16 + 2.02	21.22 + 1.13	20.85 + 1.00	22.52 + 0.23	22.03 + 0.47	22.68 + 1.16
si	in18	19.20 + 0.60	20.34 + 1.24	20.24 + 1.21	22.67 + 0.76	23.68 + 2.15	24.06 + 0.55
0	Overall	18.29 ± 0.88	19.32 + 1.29	19.59 + 0.84	22.15 + 1.28	22.51 + 1.2	23.66 + 0.71
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Values are mean \pm s.d. calculated for a total of 18 samples per each of the 25 miscanthus genotypes (3 plants \times 3 developmental stages \times 2 tissues).

945 Lignin content and ethanol yield

The digestibility of stem and leaf cell-wall samples from the 25 genotypes at peak biomass was evaluated based on the ethanol concentration in the supernatant after 72 h of incubation with

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 TABLE 3. Supernatant ethanol concentrations as mg of ethanol yielded per g of dry cell-wall biomass after 72 h of incubation with Clostridium phytofermentans

955	Genotype	Leaf	Stem
	gig01	50.90 ± 4.61	42.21 ± 7.10
	gig02	51.14 ± 3.79	44.32 ± 3.63
	gig03	51.30 ± 1.76	45.05 ± 1.92
060	hyb01	48.65 ± 6.31	42.16 ± 2.63
900	hyb02	47.01 ± 7.23	45.67 ± 2.69
	hyb03	49.00 ± 0.82	44.36 ± 1.75
	sac01	47.54 ± 2.75	48.03 ± 4.04
	sin01	50.44 ± 3.30	46.04 ± 5.66
	sin02	47.13 ± 2.95	47.55 ± 7.43
0.65	sin03	50.44 ± 3.91	48.62 ± 4.30
965	sin04	48.19 ± 6.32	48.56 ± 6.62
	sin05	47.19 ± 6.14	43.73 ± 1.96
	sin06	47.14 ± 1.90	48.17 ± 5.00
	sin07	46.77 ± 2.12	46.09 ± 3.33
	sin08	46.24 ± 2.74	47.07 ± 6.14
970	sin09	50.96 ± 5.25	46.88 ± 3.55
	sin10	44.83 ± 4.87	48.83 ± 2.79
	sin11	47.25 ± 2.08	45.73 ± 2.07
	sin12	52.46 ± 3.41	45.67 ± 2.89
	sin13	44.75 ± 1.33	52.02 ± 3.27
	sin14	51.34 ± 2.59	47.07 ± 1.19
	sin15	43.49 ± 1.60	47.04 ± 0.90
975	sin16	45.56 ± 3.55	43.33 ± 4.37
	sin17	46.99 ± 2.66	45.51 ± 3.54
	sin18	52.19 ± 5.66	44.03 ± 1.24
	Overall	48.36 ± 2.53	46.15 ± 2.28

000	Values are mean \pm s.d. for six samples per each of the 25 miscanthus
980	genotypes (3 plant replicates \times 2 tissues).

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C. phytofermentans. Ethanol yields expressed as milligrams of ethanol yielded per gram of the cell-wall biomass dry weight 1005 $(mg_{ethanol} g_{biomass}^{-1})$ ranged from a maximum of 52.5 mg g⁻¹ to a minimum of 42.2 mg g^{-1} , in a leaf and a stem sample, respectively (Table 3). ANOVA detected that the differences in ethanol vielded by the three plant replicates of each genotype were not significant (P = 0.1994). By contrast, a significant difference was detected in 1010 the ethanol yields of the various genotypes (P < 0.0001) and between the two tissues (P < 0.0001), with leaf tissue typically generating higher ethanol concentrations than the stem tissue of a given genotype (Table 3), contrasting with the observed lignin contents, which were higher in stem tissues (Fig. 4A). There 1015 was a significant correlation between lignin content and ethanol yield for stem and leaf samples collected at PB, with a Pearson correlation coefficient of r = -0.61 (P < 0.0001) indicating a negative association between lignin content and amenability to C. phytofermentans-mediated cell-wall deconstruction 1020 (Fig. 4B). Although this is a meaningful correlation, the data indicate that other factors besides lignin concentration have an exacerbating effect on recalcitrance, as supported by the individual analysis of stem and leaf data. In stem samples the interaction between ethanol vield and lignin concentration showed a coeffi-1025 cient of r = -0.65 (P = 0.0005). However, for PB leaf samples the interaction was not significant (r = -0.31, P = 0.1326).

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DISCUSSION

Biomass accumulation

Most studies on cell-wall composition in energy crops use total above-ground biomass for their analysis, as this is the most relevant material for downstream applications. However, several studies focusing on the use of forage grasses and cereal straw for animal nutrition have shown that the leaf fraction is different in terms of cell-wall composition and ruminant digestibility



FIG. 4. Lignin percentage of biomass dry weight and ethanol yield ($m_{ethanol} g_{biomass}^{-1}$) for 25 miscanthus genotypes during the peak biomass developmental stage (A) Distribution of lignin and ethanol yield measurements. (B) Pearson correlation (r) of ethanol yield with lignin content.

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when compared with the stem fraction (Love et al., 1998; McCartney et al., 2006).

- The analysis of biomass accumulation on the 25 genotypes 1065 included in our study has shown that leaf material (blade and sheath) contributed on average to more than half of the total dry biomass during the first two harvest time points. At the senesced stage, the leaf contribution was reduced to an average
- 1070 of 36.1 %, mainly due to leaf abscission during senescence. In addition to the significant contribution of leaf biomass to total above-ground biomass, it is important to emphasize the variation of leaf contributions, ranging from 42.2 to 80.4 % at actively growing, 36.3 to 78.8 % at peak biomass and 15.8 to 63.2 % at
- senescent stages (Table 1). These varying tissue contributions 1075 can have a substantial performance and economic impact on downstream biorefining processes, as compositional differences between stem and leaf biomass will lead to tissue-specific amenability for biological conversion to ethanol (discussed below).
- Our data also indicate that caution is required when interpreting 1080 correlations of cell-wall phenotyping data obtained from pooled total above-ground biomass with genetic/genomic data, as part of the observed variation might actually be due to differences in the tissue contributions to total biomass.
- 1085 The negative associations of leaf biomass contribution with tiller weight and length indicate that low-yield genotypes are more likely to have a higher leaf biomass percentage. However, significant differences were observed in leaf percentages across the genotypes, but not between the independent plant replicates of a given genotype. This observation suggests 1090 that the leaf to stem ratio could at least in part be a heritable genotype-specific trait, providing opportunities for the breeding of miscanthus cultivars that are simultaneously high yielders and high leaf biomass producers.
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Biomass composition

FTIR spectroscopy has become a powerful fingerprinting method to monitor modifications in plant cell-wall composition 1100 as it provides information about the main polysaccharides and lignin present in the cell wall (Kačuráková et al., 2000; Mouille et al., 2003; Derkacheva and Sukhov, 2008). Multivariate analysis of the FTIR data across the three developmental stages showed a distinct clustering of the spectra obtained from stem and leaf samples (Fig. 3C). This spectral segregation suggests significant 1105 compositional differences between stems and leaves, which is in accordance with reports in other species, such as maize, sorghum and rice (Oryza sativa) (Krakowsky et al., 2006; Murray et al., 2008; Jahn et al., 2011). These studies also showed that there is little genetic correlation of cell-wall 1110 polymer composition between leaf and stem tissues, suggesting that cell-wall composition is under separate genetic control in these tissues. Of the four spectral areas detected as prevalent loadings of PC1 (Fig. 3F), one is associated with lignin (c), while all 1115 others correlate with structural carbohydrates (a, g and j). As has

- been reported for sorghum (Petti et al., 2013), the prominence of band c in our data suggests higher amounts of S lignin in stem when compared with leaf tissues. However, given that the remaining three major PC1 loadings coincide with carbohydrate bands,
- 1120 it is likely that overall compositional shifts between leaf and stem cell-wall samples are more significant in their polysaccharide fractions.

Analysis of the FTIR data from different developmental stages showed that the cell-wall composition of stems from actively growing samples differs significantly from those at peak biomass or after senescence, as indicated by the discrete clusters 1125 formed during PCA (Fig. 3E). This finding most likely relates to the smaller proportion of secondary walls in actively growing stems when compared with samples at peak biomass and senesced stage. Bands associated with S lignin are noticeable negative loadings (Fig. 3G), and suggest a higher occurrence in 1130 stem samples collected during PB and SS. This is in agreement with other reports showing that more S lignin is deposited in stems when plants mature and cease to elongate, leading to a concomitant increase in the S/G ratio (Chen et al., 2002; Jung and Engels, 2002; Grabber et al., 2004). However, as above, PC1 1135 (which is responsible for the separation between elongating and mature/senesced stem samples) is predominantly correlated with carbohydrate regions of the spectra.

Grass cell walls typically contain less pectin than their dicot counterparts (Caffall and Mohnen, 2009; Vogel, 2008). It was 1140 therefore unexpected that, in addition to cellulose, variation was detected in spectral regions attributed to pectin while there was no such variation for hemicellulose. However, the masking of bands associated with hemicellulose in the spectral region defined between 1200 and 800 cm⁻¹ (Ridley et al., 2001) 1145 remains a possibility until further investigation reveals the precise nature of the structural polysaccharides involved. The presence of negative and positive PC1 loadings associated with pectins (a/j in Fig. 3F; and a,h/e in Fig. 3G) might indicate extensive differences in the structure and substitution of pectic polysaccharides 1150 between leaf and stem tissue and also as stems mature. This is in agreement with the fact that grasses display a marked developmental preference for accumulating differently modified pectins in specific cell types (Carpita, 1996). Furthermore, in the dicot *Linum usitatissimum* (flax), it has been shown that pectin synthesis 1155 and modification is different in stems and in leaves and that stem pectin incurs greater modifications during plant elongation (Bédouet et al., 2006). As for cellulose, the dominant positive band g (Fig. 3F) could indicate higher cellulose contents in stem samples. By contrast, the observed opposition of bands d, f and i 1160 in the PC1 loading plot (Fig. 3G) suggests modifications in cellulose structure as more advanced stages of maturity are reached. In effect, it has been reported that cellulose crystallinity differs between primary and secondary plant cell walls (Kataoka and Kondo, 1998; Park et al., 2013). 1165

With leaf tissue samples (Fig. 3D), the compositional differences detected by FTIR were not sufficient to create PCA clusters. This possibly reflects the fact that leaf material is less changeable, and undergoes less secondary cell-wall thickening as it matures.

The ABSL values obtained for lignin content at the senesced stage were in close agreement with other values reported for several miscanthus genotypes (Lygin et al., 2011; Domon et al., 2013), although no data for actively growing and peak biomass lignin content in miscanthus are available for compari-1175 son. As expected, there was a significant developmental and tissue effect for lignin content with (1) an overall increase in lignin as the plants mature, and (2) a higher content of lignin in stem tissues than in leaf tissues. Higher stem versus leaf lignin content has been reported for a wide range of grasses, including switchgrass (Mann et al., 2009; Shen et al., 2009)

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and M. \times giganteus (Hodgson et al., 2010; Le Ngoc Huyen et al., 2010). Our data also highlighted the limited predictive power of tissue lignin content when measured for a certain developmental stage for a specific genotype. For instance, none of the five lowest ranking genotypes for leaf lignin content at AG stage ranks

- 1185 among the five lowest for PB stage and only two of the highest ranking genotypes for stem lignin content at PB rank among the five highest at SS stage. The data also support the concept of distinct genetic control of cell-wall composition in leaf and stem tissue. As an example, stem tissue lignin content of the 1190
- three $M. \times giganteus$ genotypes included in our study ranked among the highest five at SS, while the corresponding leaf content values ranked among the lowest five. While the overall variation in lignin content across the different genotypes
- remained fairly consistent for leaf tissue with increasing maturity 1195 (AG 16.9 %, PB 16.2 %, SS 21.7 %), the variation for stem lignin content is larger for AG and PB (30 and 30.7 %, respectively), but decreases at SS (11.3 %). The decrease in variation of stem lignin content may reflect a convergence in developmental variability as plants senesce, and most likely accounts for the observed

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absence of the genotype effect in senesced samples.

We have employed a biological assay, which uses C. phytofermentans, as a means to convert isolated cell-wall biomass to ethanol. The data obtained were subsequently correlated with 1205 the lignin concentration of the same samples, thus providing a measure of the interaction of lignin content with biomass amenability to conversion. Significant variation was observed

- between the two tissues and across the genotypes in terms of ethanol yields. Supernatant ethanol concentrations showed a vari-1210 ation of 20.6 % in leaf samples and 23.4 % in stem samples across
- the genotypes (Table 3). By contrast, the average ethanol yield in leaf samples was only 4.8 % higher than in stem samples, compared with the 13.1 % difference observed in the lignin concentration between the two tissues at PB. Our data suggest that the degree
- at which tissues are lignified does not completely account for the 1215 convertibility of lignocellulosic biomass. This is supported by the fact that drastically different coefficients were found when correlating lignin contents with ethanol yields from stem or leaf samples $(r_{\text{stem}} = -0.65 \text{ and } r_{\text{leaf}} = -0.31)$. These results indicate that 1220
- lignin content has a higher relevance for the recalcitrance of stem tissue than it does for leaf tissue sampled during PB. At this stage, leaf tissue amenability to conversion may be far more influenced by other factors than it is by lignin concentration. Similar results have been reported by Le Ngoc Huyen et al.
- (2010), who found that foliar tissues show less recalcitrance 1225 than stem tissues, despite also containing appreciable amounts of lignin. Moreover, the fact that stem and leaf tissues display distinct behaviours during conversion indicates the divergent compositional arrangement of these tissues. PCA of the FTIR
- spectra obtained for stem and leaf samples collected at PB revealed 1230 discriminant loadings and clustering patterns along PC1 similar to those seen for the data across the three developmental stages (compare Fig. 3F with Supplementary Data Fig. S1). It is very likely that the divergent compositional features at the
- 1235 polysaccharide level and the lignin monomer content may be factors affecting cell-wall recalcitrance in addition to mere lignin concentration, thus making it difficult if not impossible to use the extent of tissue lignification solely as a predictor of cellwall recalcitrance.

CONCLUDING REMARKS

Our studies provide evidence that structural polysaccharides are main contributors to the compositional variability during stem development and between stem and leaf tissue. Hence, we hypothesize that the observed differences in recalcitrance 1245 between stem and leaf tissues are mainly attributed to divergent carbohydrate composition and cross-linking patterns between these two tissues. Variation in the relative contributions of leaf and stem tissues to total above-ground biomass, together with reports indicating that their composition is under separate 1250 genetic control, emphasize that improvement of cell-wall quality traits for the processing of miscanthus lignocellulosic biomass to biofuels and biomaterials must consider these observations. For gene-trait associations relating to cell-wall quality it is best practice to obtain leaf and stem compositional data separately as 1255 tissue-specific traits may be masked by total above-ground biomass and variability between samples could be largely due to varying tissue contributions to the total biomass.

SUPPLEMENTARY DATA

Supplementary data are available online at www.aob.oxfordjournals.org and consist of the following. Table S1: assignment of relevant FTIR absorption bands characteristic of miscanthus cell-wall biomass. Figure S1: principal components analysis of 1265 FTIR spectra of all samples from 25 miscanthus genotypes at peak biomass stage.

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LITERATURE CITED

- Abidi N, Cabrales L, Haigler C. 2014. Changes in the cell wall and cellulose content of developing cotton fibers investigated by FTIR spectroscopy. Carbohydrate Polymers 100: 9-16.
- Adapa P, Karunakaran C, Tabil L, Schoenau G. 2009. Potential applications of infrared and Raman spectromicroscopy for agricultural biomass. 1285 Agricultural Engineering International: CIGR Ejournal 11: 1-25.
- Allison GG, Morris C, Clifton-Brown J, Lister SJ, Donnison IS. 2011. Genotypic variation in cell wall composition in a diverse set of 244 accessions of Miscanthus. Biomass & Bioenergy 35: 4740-4747.
- Alonso-Simón A, Encina AE, García-Angulo P, Álvarez JM, Acebes JL. 2004. FTIR spectroscopy monitoring of cell wall modifications during the 1290 habituation of bean (Phaseolus vulgaris L.) callus cultures to dichlobenil. Plant Science 167: 1273-1281.
- Balch WE, Fox GE, Magrum LJ, Woese CR, Wolfe RS. 1979. Methanogens: reevaluation of a unique biological group. Microbiological Reviews 43: 260 - 296
- Bédouet L, Denys E, Courtois B, Courtois J. 2006. Changes in esterified pectins 1295 during development in the flax stems and leaves. Carbohydrate Polymers **65**: 165-173.
- Bonawitz N, Chapple C. 2010. The genetics of lignin biosynthesis: connecting genotype to phenotype. Annual Review of Genetics 44: 337-363.

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1270

1260

1325

1345

Caffall KH, Mohnen D. 2009. The structure, function, and biosynthesis of plant cell wall pectic polysaccharides. *Carbohydrate Research* 344: 1879–1900.

- 1300 cell wall pectic polysaccharides. Carbohydrate Research 344: 1879–1900.
 Carpita NC. 1996. Structure and biogenesis of the cell walls of grasses. Annual Review of Plant Physiology and Plant Molecular Biology 47: 445–476.
 - Carroll A, Somerville C. 2009. Cellulosic biofuels. Annual Review of Plant Biology 60: 165–182.

Chen L, Auh C, Chen F, et al. 2002. Lignin deposition and associated changes in anatomy, enzyme activity, gene expression, and ruminal degradability in

stems of tall fescue at different developmental stages. *Journal of Agricultural and Food Chemistry* **50**: 5558–5565.

Chundawat SPS, Beckham GT, Himmel ME, Dale BE. 2011. Deconstruction of lignocellulosic biomass to fuels and chemicals. Annual Review of Chemical and Biomolecular Engineering 2: 121–145.

1310 Clifton-Brown J, Robson P, Davey C, et al. 2013. Breeding Miscanthus for bioenergy. In: Saha M, Bhandari H, Bouton J, eds. Bioenergy feedstocks. New York: John Wiley & Sons, Inc., 67–81.

Cohen J. 1973. Eta-squared and partial eta-squared in fixed factor ANOVA designs. *Educational and Psychological Measurement* **33**: 107–112.

 Coimbra MA, Barros A, Rutledge DN, Delgadillo I. 1999. FTIR spectroscopy
 as a tool for the analysis of olive pulp cell-wall polysaccharide extracts. Carbohydrate Research 317: 145–154.

DeMartini JD, Pattathil S, Miller JS, Li H, Hahn MG, Wyman CE. 2013. Investigating plant cell wall components that affect biomass recalcitrance in poplar and switchgrass. *Energy & Environmental Science* **6**: 898–909.

Derkacheva O, Sukhov D. 2008. Investigation of lignins by FTIR spectroscopy.
 Macromolecular Symposia 265: 61–68.

Ding S-Y, Liu Y-S, Zeng Y, Himmel ME, Baker JO, Bayer EA. 2012. How does plant cell wall nanoscale architecture correlate with enzymatic digestibility? *Science* 338: 1055–1060.

Domon J-M, Baldwin L, Acket S, et al. 2013. Cell wall compositional modifications of *Miscanthus* ecotypes in response to cold acclimation. *Phytochemistry* 85: 51–61.

- Dwiyanti MS, Stewart JR, Yamada T. 2013. Germplasm resources of *Miscanthus* and their application in breeding. In: Saha M, Bhandari H, Bouton J, eds. *Bioenergy feedstocks*. New York: John Wiley & Sons, Inc., 49–66.
- Feltus FA, Vandenbrink JP. 2012. Bioenergy grass feedstock: current options and prospects for trait improvement using emerging genetic, genomic, and systems biology toolkits. *Biotechnology for Biofuels* 5: 80.
 - Foster CE, Martin TM, Pauly M. 2010. Comprehensive compositional analysis of plant cell walls (lignocellulosic biomass) Part I: Lignin. *Journal of Visualized Experiments* 37: e1745.
- Fukushima RS, Dehority BA. 2000. Feasibility of using lignin isolated from forages by solubilization in acetyl bromide as a standard for lignin analyses. *Journal of Animal Science* 78: 3135–3143.

Fukushima RS, Hatfield RD. 2004. Comparison of the acetyl bromide spectrophotometric method with other analytical lignin methods for determining lignin concentration in forage samples. *Journal of Agricultural and Food Chemistry* 52: 3713–3720.

1340 Grabber JH, Ralph J, Lapierre C, Barrière Y. 2004. Genetic and molecular basis of grass cell-wall degradability. I. Lignin-cell wall matrix interactions. Comptes Rendus Biologies 327: 455–465.

Grandis A, Souza AP, Tavares EQP, Buckeridge MS. 2014. Using natural plant cell wall degradation mechanisms to improve second generation bioethanol. In: McCann M, Buckeridge M, Carpita N, eds. *Plants and bioenergy*. New York: Springer, 211–230.

- Gorzsás A, Stenlund H, Persson P, Trygg J, Sundberg B. 2011. Cell-specific chemotyping and multivariate imaging by combined FT-IR microspectroscopy and orthogonal projections to latent structures (OPLS) analysis reveals the chemical landscape of secondary xylem. *The Plant Journal* 66: 903–914.
- Gwon JG, Lee SY, Doh GH, Kim JH. 2010. Characterization of chemically modified wood fibers using FTIR spectroscopy for biocomposites. *Journal of Applied Polymer Science* 116: 3212–3219.
 - Hall M, Bansal P, Lee JH, Realff MJ, Bommarius AS. 2010. Cellulose crystallinity – a key predictor of the enzymatic hydrolysis rate. *FEBS Journal* 277: 1571–1582.

 Heaton EA, Dohleman FG, Long SP. 2008. Meeting US biofuel goals with less land: the potential of *Miscanthus*. *Global Change Biology* 14: 2000–2014.
 Himmel ME, Ding S, Y. Johnson DK, et al. 2007. Biomosc proclaitmency and in

Himmel ME, Ding S-Y, Johnson DK, et al. 2007. Biomass recalcitrance: engineering plants and enzymes for biofuels production. Science 315: 804–807. Hodgson EM, Fahmi R, Yates N, et al. 2010. Miscanthus as a feedstock for fast-pyrolysis: does agronomic treatment affect quality? *Bioresource Technology* 101: 6185–6191.

Hodgson EM, Nowakowski DJ, Shield I, et al. 2011. Variation in Miscanthus chemical composition and implications for conversion by pyrolysis and thermo-chemical bio-refining for fuels and chemicals. Bioresource Technology 102: 3411–3418.

Hungate R. 1969. A roll tube method for cultivation of strict anaerobes. In: Norris J, Ribbons DW, eds. *Methods in microbiology*. New York: Academic Press, 117–132.

Jahn CE, Mckay JK, Mauleon R, et al. 2011. Genetic variation in biomass traits among 20 diverse rice varieties. Plant Physiology 155: 157–168.

Jensen E, Farrar K, Thomas-Jones S, Hastings A, Donnison I, Clifton-Brown J. 2011. Characterization of flowering time diversity in *Miscanthus* species. *GCB Bioenergy* 3: 387–400.

Jung HG, Engels FM. 2002. Alfalfa stem tissues: cell wall deposition, composition, and degradability. Crop Science 42: 524–534.

- Kačuráková M, Capek P, Sasinková V, Wellner N, Ebringerová A. 2000. FT-IR study of plant cell wall model compounds: pectic polysaccharides and hemicelluloses. *Carbohydrate Polymers* 43: 195–203.
- Kataoka Y, Kondo T. 1998. FT-IR microscopic analysis of changing cellulose crystalline structure during wood cell wall formation. *Macromolecules* 31: 760–764.

Krakowsky MD, Lee M, Coors JG. 2006. Quantitative trait loci for cell wall components in recombinant inbred lines of maize (*Zea mays* L.) II: Leaf sheath tissue. *Theoretical and Applied Genetics* **112**: 717–726.

Labbé N, Rials TG, Kelley SS, Cheng Z-M, Kim J-Y, Li Y. 2005. FT-IR imaging and pyrolysis-molecular beam mass spectrometry: new tools to investigate wood tissues. *Wood Science and Technology* **39**: 61–76.

Le Ngoc Huyen T, Rémond C, Dheilly RM, Chabbert B. 2010. Effect of harvesting date on the composition and saccharification of *Miscanthus* × *giganteus*. *Bioresource Technology* 101: 8224–8231.

- Lee SJ, Warnick TA, Leschine SB, Hazen SP. 2012a. A high-throughput biological conversion assay for determining lignocellulosic quality. In: Normanly J, ed. *High-throughput phenotyping in plants*. New York: Humana Press, 341–349.
- Lee SJ, Warnick TA, Pattathil S, et al. 2012b. Biological conversion assay using *Clostridium phytofermentans* to estimate plant feedstock quality. *Biotechnology for Biofuels* 5: 5.
- Levine TR, Hullett CR. 2002. Eta squared, partial eta squared, and misreporting of effect size in communication research. *Human Communication Research* 28: 612–625.
- Liu C, Xiao L, Jiang J, et al. 2013. Biomass properties from different Miscanthus species. Food and Energy Security 2: 12–19.

Love G, Snape C, Jarvis M. 1998. Comparison of leaf and stem cell-wall components in barley straw by solid-state ¹³C NMR. *Phytochemistry* **49**: 1191–1194.

Lygin AV, Upton J, Dohleman FG, *et al.* 2011. Composition of cell wall phenolics and polysaccharides of the potential bioenergy crop – *Miscanthus*. *Global Change Biology Bioenergy* **3**: 333–345.

Mann DGJ, Labbé N, Sykes RW, et al. 2009. Rapid assessment of lignin content and structure in switchgrass (*Panicum virgatum* L.) grown under different environmental conditions. *BioEnergy Research* 2: 246–256.

Marry M, McCann MC, Kolpak F, White AR, Stacey NJ, Roberts K. 2000. Extraction of pectic polysaccharides from sugar-beet cell walls. *Journal* of the Science of Food and Agriculture 80: 17–28.

Matos DA, Whitney IP, Harrington MJ, Hazen SP. 2013. Cell walls and the developmental anatomy of the *Brachypodium distachyon* stem internode. 1405 *PLoS ONE* 8: e80640.

McCann MC, Carpita NC. 2008. Designing the deconstruction of plant cell walls. *Current Opinion in Plant Biology* 11: 314–320.

McCann MC, Bush M, Milioni D, *et al.* 2001. Approaches to understanding the functional architecture of the plant cell wall. *Phytochemistry* 57: 811–821.

McCann MC, Defernez M, Urbanowicz BR, et al. 2007. Neural network analysis of infrared spectra for classifying cell wall architectures. Plant Physiology 143: 1314–1326.

- McCartney DH, Block HC, Dubeski PL, Ohama AJ. 2006. Review: The composition and availability of straw and chaff from small grain cereals for beef cattle in Western Canada. *Canadian Journal of Animal Science* 86: 443–455.
- Monties B. 1989. Lignins. In: Dey PM, Harborne JB eds. *Methods in plant biochemistry*. New York: Academic Press, 113–158.

1490

- Mouille G, Robin S, Lecomte M, Pagant S, Höfte H. 2003. Classification and identification of arabidopsis cell wall mutants using Fourier-transform infrared (FT-IR) microspectroscopy. *Plant Journal* 35: 393–404.
- Murray SC, Rooney WL, Mitchell SE, et al. 2008. Genetic improvement of sorghum as a biofuel feedstock: II. QTL for stem and leaf structural carbohydrates. Crop Science 48: 2180–2193.
 - **Oh SY, Yoo DI, Shin Y, Seo G. 2005.** FTIR analysis of cellulose treated with sodium hydroxide and carbon dioxide. *Carbohydrate Research* **340**: 417–428.
- Park YB, Lee CM, Koo BW, Park S, Cosgrove DJ, Kim SH. 2013. Monitoring meso-scale ordering of cellulose in intact plant cell walls using sum frequency generation spectroscopy. *Plant Physiology* 163: 907–913.
 - Petti C, Harman-Ware AE, Tateno M, et al. 2013. Sorghum mutant RG displays antithetic leaf shoot lignin accumulation resulting in improved stem saccharification properties. *Biotechnology for Biofuels* 6: 146.
- Purdy SJ, Maddison AL, Jones LE, et al. 2013. Characterization of chillingshock responses in four genotypes of Miscanthus reveals the superior tolerance of M. × giganteus compared with M. sinensis and M. sacchariflorus. Annals of Botany 111: 999–1013.
 - Ralph J. 2010. Hydroxycinnamates in lignification. *Phytochemistry Reviews* 9: 65–83.
- Ralph J, Hatfield RD, Quideau S, Helm RF, Grabber JH, Jung H-JG. 1994.
 Pathway of *P*-coumaric acid incorporation into maize lignin as revealed by

NMR. Journal of the American Chemical Society 116: 9448–9456.
Ridley BL, O'Neill MA, Mohnen D. 2001. Pectins: structure, biosynthesis, and oligogalacturonide-related signaling. *Phytochemistry* 57: 929–967.

- Robson P, Mos M, Clifton-Brown J, Donnison I. 2012. Phenotypic variation in senescence in *Miscanthus*: towards optimising biomass quality and quantity. *BioEnergy Research* 5: 95–105.
- Robson P, Jensen E, Hawkins S, et al. 2013. Accelerating the domestication of a bioenergy crop: identifying and modelling morphological targets for sustainable yield increase in *Miscanthus*. Journal of Experimental Botany 64: 4143–55.
- Schulz H, Baranska M. 2007. Identification and quantification of valuable plant substances by IR and Raman spectroscopy. *Vibrational Spectroscopy* 43:
- ⁴⁴⁵ 13–25.

- Séné CFB, McCann MC, Wilson RH, Grinter R. 1994. FT-Raman and FT-infrared spectroscopy: an investigation of five higher plant cell walls and their components. *Plant Physiology* 106: 1623–1633.
- Shen H, Fu C, Xiao X, et al. 2009. Developmental control of lignification in stems of lowland switchgrass variety Alamo and the effects on saccharification efficiency. *BioEnergy Research* 2: 233–245.
- Slavov G, Allison G, Bosch M. 2013a. Advances in the genetic dissection of plant cell walls: tools and resources available in *Miscanthus*. Frontiers in Plant Science 4: 217.
- Slavov G, Robson P, Jensen E, et al. 2013b. Contrasting geographic patterns of genetic variation for molecular markers vs. phenotypic traits in the energy grass Miscanthus sinensis. Global Change Biology Bioenergy 5: 562–571. 1485
- **Tobimatsu Y, Elumalai S, Grabber JH, Davidson CL, Pan X, Ralph J. 2012.** Hydroxycinnamate conjugates as potential monolignol replacements: *in vitro* lignification and cell wall studies with rosmarinic acid. *ChemSusChem* **5**: 676–686.
- van der Weijde T, Alvim Kamei CL, Torres AF, et al. 2013. The potential of C4 grasses for cellulosic biofuel production. Frontiers in Plant Science 4: 107.
- Vanholme R, Demedts B, Morreel K, Ralph J, Boerjan W. 2010. Lignin biosynthesis and structure. *Plant Physiology* **153**: 895–905.
- Vogel J. 2008. Unique aspects of the grass cell wall. *Current Opinion in Plant Biology* 11: 301–307.
- Warnick TA, Methé BA, Leschine SB. 2002. Clostridium phytofermentans sp. nov., a cellulolytic mesophile from forest soil. International Journal of Systematic and Evolutotionary Microbiology 52: 1155–1160.
- Wilson RH, Smith AC, Kačuráková M, Saunders PK, Wellner N, Waldron KW. 2000. The mechanical properties and molecular dynamics of plant cell wall polysaccharides studied by Fourier-transform infrared spectroscopy. *Plant Physiology* 124: 397–406.
- Zhang T, Wyman CE, Jakob K, Yang B. 2012. Rapid selection and identification of *Miscanthus* genotypes with enhanced glucan and xylan yields from hydrothermal pretreatment followed by enzymatic hydrolysis. *Biotechnology for Biofuels* 5: 56.
- **Zhou G, Taylor G, Polle A. 2011.** FTIR-ATR-based prediction and modelling of lignin and energy contents reveals independent intra-specific variation of these traits in bioenergy poplars. *Plant Methods* **7**: 1–10.

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