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# Research paper

# Enhanced substrate degradation and methane yield with maleic acid pre-treatments in biomass crops and residues

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#### ABSTRACT

Organic acids are envisaged as alternative catalysts to strong mineral acids, in pre-treatment of lignocellulosic biomass for anaerobic digestion (AD). To evaluate this hypothesis, an untreated control and four pre-treatments (25 °C for 24 h) involving two levels of maleic acid (34.8 and 69.6 kg m<sup>-3</sup>), alone and combined with sulphuric acid (4 kg m<sup>-3</sup>), were studied in three agricultural substrates: Arundo (aka giant reed), Barley straw and B133 fibre sorghum. Methane production was assessed in a batch AD assay (35 °C for 51 days) with 4 g  $L^{-1}$  of volatile solid (VS) load. Fibre composition and structure were investigated through chemical analysis and Fourier transform infrared (FTIR) spectrometry. Arundo and B133 that were the most and least recalcitrant substrate, respectively, staged the highest and lowest increase in methane with high maleic acid: +62% over 218 cm<sup>3</sup> g<sup>-1</sup> of VS in untreated Arundo; +36% over 284 cm<sup>3</sup> g<sup>-1</sup> of VS in untreated B133. Barley straw showed an intermediate behaviour (+41% over 269 cm<sup>3</sup> g<sup>-1</sup> of VS). H<sub>2</sub>SO<sub>4</sub> addition to maleic acid did not improve CH<sub>4</sub> output. The large increase in methane yield determined by pre-treatments was reflected in the concurrent decrease of fibre (between 14 and 39% depending on fibrous component). Based on FTIR spectra, bands assigned to hemicellulose and cellulose displayed lower absorbance after pre-treatment, supporting the hypothesis of solubilisation of structural carbohydrates and change in fibre structure. Hence, maleic acid was shown a suitable catalyst to improve biodegradability of ligno-cellulosic biomass, especially in recalcitrant substrates as Arundo.

#### 1. Introduction

Ligno-cellulosic biomass of agricultural origin (i.e. energy crops and residues) is one of the key options for meeting the world's energy demand [1], while at the same time minimizing competition with food crops [2]. Anaerobic digestion (AD) is widely used at present for the energy conversion of agricultural biomasses.

These substrates are composed of the three fibre fractions (cellulose, hemicellulose and lignin) in a proportion and relationship that varies according to plant species. Among these fractions, lignin is fairly resistant to AD, retarding or preventing the

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hydrolysis of carbohydrates [3]. Therefore, during AD of lignocellulosic substrates, hydrolysis is considered the rate limiting step [4], influencing kinetics and, consequently, production of biogas. To overcome the recalcitrance of ligno-cellulosic substrates, pre-treatments are devised to loosen fibre structure, remove or rearrange lignin and hydrolyze cellulose and hemicellulose [5], resulting in faster hydrolysis and improved methane yield.

Pre-treatments rely on physical, chemical and biological means, sometimes combined [5]. Among them, chemical pre-treatments with the use of dilute sulphuric acid have been widely investigated [5–9], performing satisfactory results. The main reactions occurring during dilute sulphuric acid pre-treatment are the hydrolysis of hemicellulose [10], a partial hydrolysis of cellulose and a solubilisation of lignin, leading to changes in biomass structure [11]. The main drawbacks from the use of sulphuric acid are: i) corrosion of the equipment during AD, and ii) production of  $SO_4^2$ -boosting the activity of sulphate-reducing bacteria, which are

*Abbreviations:* AD, anaerobic digestion; AIL, acid insoluble lignin; Cell, cellulose; FTIR, Fourier transform infrared spectrometry; H-cell, hemicellulose; LSD, least significant difference; TS, total solids; VS, volatile solids.

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known to negative affect the incubation [12]. To avoid these constraints, the use of organic instead of mineral acids could be envisaged in the frame of a biomimetic approach, i.e. one that mimics natural enzymes. This is based on the fact that enzymes catalysing cellulose and hemicellulose hydrolysis act through a general acid-base mechanism by means of two carboxylic acids in the enzyme active site [13]. Accordingly, dicarboxylic acids having a similar catalytic structure as enzymes, were proposed as biomimetic catalysts [14], and they were shown to be actually more selective for  $\beta$ -(1,4)-glycolic bonds than sulphuric acid [13]. Maleic acid was shown the most favourable dicarboxylic acid as it concerns catalysis selectivity [15]. However, even if maleic acid is easier to handle than sulphuric acid due to its lower strength, this approach appears more expensive compared to sulphuric acid. For that reason Guo et al. [14] suggested a combination of mineral and organic acid as that fetching higher biomimetic catalysis while, at the same time, reducing the cost for organic acids with a cheaper mineral acid.

With this premise, a vast literature studied the biomimetic approach to enhance hemicellulose hydrolysis and cellulose access for enzymatic attack, in view of higher bioethanol output [8,14,16–18]. Conversely, only a study addressed methane potential [11]. Therefore, the objective of this work was to investigate bland pre-treatments with organic acid (maleic acid) and the combination of mineral and organic acid (sulphuric + maleic acid) on three ligno-cellulosic substrates of agricultural origin. The effects of pretreatments were evaluated on fibre composition, structure and methane yield compared to untreated substrates.

#### 2. Material and methods

#### 2.1. Investigated substrates

The three substrates, also used in a previous study on alkaline pre-treatments [19], were: Arundo donax L. (Arundo), a wild type sourced locally (44° 33' N, 11° 21' E; 32 m above sea level); B133 fibre sorghum [Sorghum bicolor (L.) Moench], a hybrid from Syngenta Seeds (Casalmorano, CR, Italy); Barley (Hordeum vulgare L.) straw, cv. Ketos from Limagrain (Busseto, PR, Italy). Biomass samples of the two crops and the crop residue were collected from fields at the experimental farm of the University of Bologna in Cadriano (same coordinates as above), Italy, during the year 2010. The soil is a deep alluvial loam, while climate falls in the Mediterranean North environmental zone [20]. Further detail on crop husbandry is given in the above referred work [19]. Arundo was harvested at initial senescence (October 5, 2010), while B 133 sorghum was harvested at hard dough stage (October 18, 2010). Barley straw was collected after combine harvesting of the dry grain, in summer 2010. Biomass samples of the three substrates were oven dried (60 °C) and ground at 2 mm for chemical characterization, pre-treatments and AD.

# 2.2. Acid pre-treatments

Pre-treatments were carried out with four combinations of organic and mineral acids: maleic acid at low and high concentration (34.8 and 69.6 kg m<sup>-3</sup>), alone and combined with sulphuric acid at 4 kg m<sup>-3</sup> (Table 1). Sulphuric acid concentration was based on a previous assay conducted on the same three substrates.

Pre-treatment procedure and subsequent assessments are summarized in Fig. 1.

Pre-treatment was carried out in the dark at 25 °C for 24 h, during which time substrates previously dried (60 °C) and ground (2 mm) were supplemented with the acid solutions and stirred (10,000 RCF). Substrate amounts were set to maintain a

#### Table 1

Pre-treatment conditions. O and M mean organic (maleic) and mineral (sulphuric) acid (Ac.), respectively; L and H indicate low and high concentration of maleic acid, respectively. Pre-treatments were conducted at 25  $^{\circ}$ C for 24 h.

Pre-treatment	Maleic acid	Sulphuric acid
	Kg m <sup>-3</sup>	
Untreated	-	_
OAc.L	34.8	_
OAc.H	69.6	_
M+OAc.L	34.8	4
M+OAc.H	69.6	4



**Fig. 1.** Scheme of pre-treatment procedure and subsequent assessments. OAc., organic (maleic) acid; M+OAc., mineral (4 kg of H<sub>2</sub>SO<sub>4</sub> m<sup>-3</sup>) and organic acid; L and H mean low (34.8 kg m<sup>-3</sup>) and high (69.6 kg m<sup>-3</sup>) level of maleic acid, respectively. Cell, cellulose; H-cell, hemicellulose; AIL, acid insoluble lignin; FTIR, Fourier transform infrared spectrometry; AD, anaerobic digestion; VS, volatile solids.

concentration of 100 mL of acid solution  $kg^{-1}$  of total solids (TS). At the end of the process, the liquid fraction was separated by vacuum pump equipped with a Whatman GF/C, Ø 47 mm filter, while the solid fraction was washed and dried (60 °C for 48 h) prior to chemical analysis (Fig. 1).

In parallel to this, the same pre-treatments were carried out on the three substrates in view of AD (Fig. 1).

#### 2.3. Anaerobic digestion

A batch AD assay was carried out using the same basic procedure described in Ref. [19]. Briefly, the inoculum was retrieved from the same source, and was subjected to the same period of adaptation. At the end, it showed the following data: TS, 32 mg g<sup>-1</sup>; volatile solids (VS), 26 mg g<sup>-1</sup>; total alkalinity, 29 g of CaCO<sub>3</sub> L<sup>-1</sup>; pH, 7.8.

The rest of the procedure was the same as in Refs. [19], namely as it concerns incubation temperature (35 °C), organic load (4 g of

Wavenumber ( $cm^{-1}$ )	Vavenumber (cm <sup>-1</sup> ) Group		Literary reference	
2900	C–H stretching	Cell	[23]	
1720	C=O stretching acetyl or carboxylic acid	H-cell and lignin	[24,25]	
1430	–CH <sub>2</sub> bending	Cell	[26]	
1375	C–H deformation	Cell	[27]	
1315	-CH <sub>2</sub> wagging vibrations	Cell and H-cell	[25]	
1158	C–O–C stretching	Cell and H-cell	[25]	
898	Glucose ring stretch, C–H deformation	Cell	[28]	

Cell, cellulose; H-cell, hemicellulose.

VS L<sup>-1</sup>), and the relatively high inoculum to substrate ratio (5.2:1, VS:VS) that was chosen to offset inhibiting factors [21,22]. The following six controls were added: blank (inoculum alone); blank plus the four combinations of maleic and sulphuric acid; glucose at the same organic load (4 g of VS L<sup>-1</sup>). This made up a total of 63 serum bottles under simultaneous incubation.

#### 2.4. Analytical methods

#### 2.4.1. Chemical analyses

Chemical determinations on untreated substrates were the same as in the cited work involving the same samples [19]: TS, VS, total Kjeldahl nitrogen, protein, lipids, starch, extractives, hemicellulose (H-cell) and cellulose (Cell), and acid insoluble lignin (AIL). The same data are, therefore, used in this work. Further detail on analytical methods is given in Ref. [19].

#### 2.4.2. Biogas measurement and analysis

Eleven times during the incubation (day 2, 4, 6, 9, 12, 16, 19, 25, 31, 41 and 51), biogas amount and composition were assessed with

the same procedure and equipment described in Ref. [19]. Methane production from the controls described in Section 2.3 were sub-tracted from methane produced from the respective substrates. Subsequent calculations [19] allowed us to express the amount of methane (cm<sup>3</sup> g<sup>-1</sup> of VS) produced during the incubation.

# 2.4.3. FTIR analysis

The structure of the solid fraction of untreated and pre-treated substrates was analysed with Fourier transform infrared (FTIR) spectrometry, using the same instrumentation, software and procedure as in Ref. [19]. Seven characteristic bands of absorbance (Table 2) were retained for subsequent discussion.

Thereafter, the total crystallinity index (TCI) and the lateral order index (LOI) were calculated as the respective 1375 to 2900 and 1430 to 898 cm<sup>-1</sup> peak ratio [26,27]. TCI and LOI express functional relationships in Cell fractions.

#### 2.5. Data analysis

Data of chemical analysis were subjected to one-way ANOVA for

compositional analysis of anticated substrates.
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Substrate	VS mg $g^{-1}$ of TS	C:N	Extr.	Protein	Lipids	Starch	Cell	H-cell	AIL		
			mg g <sup>-1</sup> of	$mg g^{-1}$ of VS							
Arundo	926 b	56.9 b	211 b	51.2	9.5 c	38.5 b	322 a	205 b	229 a		
Barley straw	924 c	64.1 a	221 b	44.4	12.2 b	16.2 c	317 a	214 a	215 ab		
B133 sorghum	949 a	54.2 b	349 a	52.2	16.3 a	53.3 a	228 b	126 c	198 b		

All traits except protein were significant at the ANOVA; a, b, c, significantly different means (LSD test at  $P \le 0.05$ ). VS, volatile solids; TS, total solids; Extr., extractives; Cell, cellulose; H-cell, hemicellulose; AlL, acid insoluble lignin.



Fig. 2. Cumulative CH<sub>4</sub> yield of untreated and pre-treated substrates during the anaerobic digestion assay. OAc., organic (maleic) acid; M+OAc., mineral (4 kg of H<sub>2</sub>SO<sub>4</sub> m<sup>-3</sup>) and organic acid; L and H mean low (34.8 kg m<sup>-3</sup>) and high (69.6 kg m<sup>-3</sup>) level of maleic acid, respectively. Vertical bars, ± standard deviation.



**Fig. 3.** Significant substrate × treatment interaction on cumulative CH<sub>4</sub> yield at the end of the incubation. U, L and H mean untreated and pre-treated with low (34.8 kg m<sup>-3</sup>) and high (69.6 kg m<sup>-3</sup>) level of maleic acid (OAc.) alone or in combination with sulphuric acid at 4 kg m<sup>-3</sup> (M+OAc.). LSD<sub>0.05</sub>, least significant difference at  $P \le 0.05$  (19.6 cm<sup>3</sup> of CH<sub>4</sub> g<sup>-1</sup> of VS). Vertical bars,  $\pm$  standard deviation.

the substrates factor. Data of cumulative CH<sub>4</sub> yield were subjected to two-way ANOVA addressing substrates, pre-treatments and their interaction. Fisher's least significant difference (LSD) at  $P \leq 0.05$  was used to separate data of significant ANOVA sources.

#### 3. Results and discussion

### 3.1. Compositional analysis of untreated substrates

Untreated substrates showed the characteristics reported in Table 3.

VS exhibited a limited variation (from 924 to 949 mg  $g^{-1}$  of TS). The C:N mass ratio ranged between 54 (B133 sorghum) and 64 (Barley straw), which is above the limit (30) indicated for optimum AD conditions [28]. The extractives were an approximate 200 mg  $g^{-1}$  of VS in Arundo and Barley straw; a much higher amount (~350 mg  $g^{-1}$  of VS) in B133 sorghum. Barley straw had a mildly lower protein content than the averaged Arundo and B133 sorghum (ca. 45 vs. 52 mg  $g^{-1}$  of VS), associated with a much lower starch content (16.2, 38.5 and 53.3 mg  $g^{-1}$  of VS in the three respective substrates). B133 sorghum evidenced a lower amount of Cell (228 mg  $g^{-1}$  of VS) and H-cell (126 mg  $g^{-1}$  of VS) than the averaged Arundo and Barley straw (319 and 210 mg  $g^{-1}$  of VS for the two respective carbohydrates). The sum of the two structural

carbohydrates amounted to an approximate 570, 580 and 380 mg g<sup>-1</sup> of VS in Arundo, Barley straw and B133 sorghum, respectively. AlL staged a narrow range between 198 and 229 mg g<sup>-1</sup> of VS in the respective B133 sorghum and Arundo. Similar structural carbohydrates and lignin content were found by Scordia et al. [17] in Arundo harvested in a wild area, while Sambusiti et al. [29] had found a higher content of lignin (+20%) in the same sorghum hybrid (B133).

#### 3.2. Methane yield during the incubation

The time trend of cumulative CH<sub>4</sub> yield is depicted in Fig. 2. In the first 4 days of incubation, CH<sub>4</sub> yield was similar in all substrates. Thereafter, pre-treated substrates exhibited a steep increase of CH<sub>4</sub> output, followed by a slowdown between ten and twenty days, and by subsequent resumption of the kinetics. Untreated substrates outlined a steady methanation rate (Fig. 2). At 10 days, ca. 50% of cumulative CH<sub>4</sub> yield was produced in the average of the three untreated substrates, vs. 68% in substrates pre-treated with OAc.L and M+OAc.L, and 76% with OAc.H and M+OAC.H (Fig. 2).

Enhanced CH<sub>4</sub> production in pre-treated substrates at the beginning of incubation might be due to conversion of easily degradable compounds, indicating an increase of the overall biodegradability after pre-treatment. This is in agreement with Lee

![](_page_4_Figure_12.jpeg)

**Fig. 4.** Fibre composition of untreated and pre-treated substrates. OAc., organic (maleic) acid; M+OAc., mineral (4 kg of  $H_2SO_4$  m<sup>-3</sup>) and organic acid; L and H mean low (34.8 kg m<sup>-3</sup>) and high (69.6 kg m<sup>-3</sup>) level of maleic acid, respectively. Cell, cellulose; H-cell, hemicellulose; AIL, lignin. Vertical bars, ± standard deviation.

![](_page_5_Figure_1.jpeg)

**Fig. 5.** Fingerprint range from 4000 to 600 cm<sup>-1</sup> of the FITR spectra of Untreated (black line), OAc.H (gray line) and M+OAc.H (dark gray) pre-treated Arundo (a), Barley straw (b) and B133 sorghum (c). OAc.H, organic (maleic) acid at high level (69.6 kg m<sup>-3</sup>); M+OAc., mineral (4 kg of H<sub>2</sub>SO<sub>4</sub> m<sup>-3</sup>) and organic acid at high level.

and Jeffries [8], who reported that pre-treatment with the same dicarboxylic acid of our experiment (maleic acid) released a remarkable amount of monomeric sugars derived from structural carbohydrates (H-cell and Cell). Conversely, the temporary slow-down of CH<sub>4</sub> production observed between 10 and 20 days was

likely due to a need of microbial adaptation before tackling the residual, more recalcitrant fraction.

In each substrate, cumulative CH<sub>4</sub> yield diverged progressively between untreated and pre-treated samples. Resulting from this, the substrate  $\times$  treatment interaction was statistically significant at the end of the incubation, meaning that pre-treatments exerted a different effect on final CH<sub>4</sub> output depending on each specific substrate (Fig. 3).

In untreated substrates, Arundo showed a lower CH<sub>4</sub> yield (218 cm<sup>3</sup> g<sup>-1</sup> of VS) than Barley straw and B133 sorghum (average, 276 cm<sup>3</sup> g<sup>-1</sup> of VS). In treated substrates, Arundo achieved a similar CH<sub>4</sub> yield with OAc.L and M+OAc.L (average, 272 cm<sup>3</sup> g<sup>-1</sup> of VS), and a further increase with M+OAc.H and OAc.H (330 and 354 cm<sup>3</sup> g<sup>-1</sup> of VS, respectively; Fig. 3). This corresponds to a 24, 51 and 62% respective gain in CH<sub>4</sub> yield vs. untreated Arundo. Pretreatments in Barley straw outlined a similar CH<sub>4</sub> yield in OAc.L and M+OAc.L (average, 323 cm<sup>3</sup> g<sup>-1</sup> of VS), as in OAc.H and M+OAc.H (average, 380 cm<sup>3</sup> g<sup>-1</sup> of VS), determining a respective 20 and 41% increase. Conversely, in B133 sorghum a variable CH<sub>4</sub> yield was obtained after pre-treatments: 307, 335, 373 and 398 cm<sup>3</sup> g<sup>-1</sup> of VS at M+OAc.L, OAc.L, M+OAc.H and OAc.H, respectively. Hence, CH<sub>4</sub> increase ranged between 8% (M+OAc.L) and 40% (OAc.H).

It is worth noting that B133 sorghum, a more biodegradable substrate, benefited less from pre-treatments than Barley straw and Arundo. Moreover, the addition of mineral acid to the organic acid determined a somewhat lower CH<sub>4</sub> production, especially in B133 sorghum (Fig. 3). This may be due to competition between sulphate-reducing bacteria and methane producing *Archea* [30].

Fernandes et al. [11] studied the effect of maleic acid (6 kg m<sup>-3</sup>) pre-treatment (150 °C for 30 min) on methane yield of three different substrates as hay, straw and bracken with different lignin content (25, 57 and 185 mg g<sup>-1</sup> of VS, respectively). They found that methane yield of hay and straw was not enhanced after pre-treatment, while a 57% increase (ca. 110 cm<sup>3</sup> g<sup>-1</sup> of VS) was shown in bracken. It appeared, therefore, that the effect of pre-treatment was more profound in ligno-cellulosic biomass with a higher lignin content. This is consistent with the trend observed in our experiment: methane yield after pre-treatment augmented in parallel with the content of lignin in untreated substrate: 25, 31 and 41% CH<sub>4</sub> yield increase with a lignin content of 198, 215 and 229 mg g<sup>-1</sup> of VS, in the respective B133 sorghum, Barley straw and Arundo (Table 3).

In the previous experiment addressing mild alkaline pretreatments (NaOH from 2 to 6 kg m<sup>-3</sup>) with the same three substrates [19], lower increases in CH<sub>4</sub> yield were observed: up to 10, 23 and 30% in B133 sorghum, Barley straw and Arundo, respectively. Hence, maleic acid was shown a catalyst with a remarkable potential, in light of the recent findings on AD of ligno-cellulosic biomass.

# 3.3. Changes in fibre composition and structure with pretreatments

During biodegradation, layers of lignin shield the two structural carbohydrates from enzymatic attack. Pre-treatments are devised to alter the ligno-cellulosic structure, even disrupt it, easing the bioconversion of ligno-cellulosic biomass. In this experiment, the reduction of the three structural components (Cell, H-cell and lignin) after pre-treatment varied in extent, depending on each specific substrate (Fig. 4). Conversely, the four acid combinations determined few, inconsistent variations between them. Arundo staged the strongest solubilisation of Cell and H-cell as the effect of pre-treatments (from 18 to 21% and from 32 to 39% in the two respective compounds), compared to B133 sorghum (from 5 to 19% and from 4 to 16%) and Barley straw (from 3 to 13% and from 25 to

#### Table 4

Absorbance related to bands assigned to H-cell and Cell (functional groups and linkages reported in Table 3). OAc. organic (maleic) acid; M+OAc. mineral (4 kg of H<sub>2</sub>SO<sub>4</sub> m<sup>-3</sup>) and organic acid; L and H mean low (34.8 kg m<sup>-3</sup>) and high (69.6 kg m<sup>-3</sup>) levels of maleic acid. LOI (Lateral Order Index) and TCI (Total Crystallinity Index) are based on specific peak ratios. In brackets, standard deviation.

Substrate	<b>Pre-treatments</b>	Wavenumbers (cm <sup>-1</sup> )							LOI	TCI
		2900	1720	1430	1375	1315	1158	898		
Arundo	Untreated	0.33	0.32	0.45	0.47	0.42	0.45	0.34	1.32 (0.01)	1.41 (0.01)
	OAc.L	0.12	0.15	0.23	0.26	0.24	0.29	0.15	1.57 (0.21)	2.08 (0.07)
	OAc.H	0.16	0.20	0.28	0.32	0.32	0.21	0.16	1.76 (0.33)	2.00 (0.52)
	M+OAc.L	0.14	0.17	0.25	0.27	0.25	0.26	0.14	1.69 (0.11)	1.89 (0.01)
	M+OAc.H	0.08	0.12	0.16	0.17	0.15	0.19	0.09	1.76 (0.01)	2.03 (0.01)
Barley straw	Untreated	0.23	0.14	0.32	0.33	0.31	0.40	0.21	1.49 (0.04)	1.40 (0.07)
	OAc.L	0.19	0.18	0.30	0.33	0.30	0.34	0.19	1.57 (0.05)	1.77 (0.35)
	OAc.H	0.25	0.23	0.32	0.37	0.32	0.40	0.24	1.37 (0.06)	1.68 (0.48)
	M+OAc.L	0.14	0.11	0.22	0.24	0.22	0.27	0.13	1.70 (0.03)	1.79 (0.12)
	M+OAc.H	0.16	0.16	0.24	0.28	0.25	0.30	0.17	1.47 (0.14)	1.77 (0.20)
B133 sorghum	Untreated	0.25	0.21	0.38	0.40	0.37	0.42	0.28	1.38 (0.02)	1.57 (0.06)
	OAc.L	0.17	0.20	0.28	0.33	0.31	0.38	0.20	1.41 (0.12)	2.05 (0.23)
	OAc.H	0.09	0.12	0.18	0.22	0.20	0.27	0.12	1.45 (0.07)	2.44 (0.32)
	M+OAc.L	0.14	0.19	0.28	0.33	0.31	0.42	0.19	1.50 (0.17)	2.42 (0.26)
	M+OAc.H	0.09	0.12	0.20	0.22	0.20	0.26	0.13	1.49 (0.07)	2.31 (0.25)

33%). Lastly, in the case of AIL the strongest reduction was observed in B133 sorghum (14%, as average of the four pre-treatments), compared to Arundo (12%) and Barley straw (9%) (Fig. 4).

In the literature, maleic acid has already performed satisfactorily in H-cell hydrolysis [13,31]. H-cell reduction up to ca. 80% was obtained in wheat straw pre-treated with low concentration of maleic acid (6 kg m<sup>-3</sup>) combined with microwave heating (170 °C for 30 min), while only a 12% Cell reduction was observed [32]. Likewise, Guo et al. [14] reported an 80% H-cell reduction in *Miscanthus* after pre-treatment (170 °C for 6 min) with maleic and sulphuric acid at 61.5 and 7.4 kg m<sup>-3</sup>, respectively.

In addition, these authors demonstrated that maleic acid is mainly active on the easily hydrolysable fraction of H-cell. In fact, H-cell in most ligno-cellulosic substrates may be present under two fractions, easy and hard to hydrolyze; the latter portion has been shown to account for 35% of the total [33]. Compared to the above referred works, our experiment exhibited up to 39% H-cell reduction, probably derived from the easily hydrolysable fraction. Beside its activity in the hydrolysis of H-cell, maleic acid proved also effective in reducing Cell and lignin (-19% and -15% in the two respective components, as average of all substrates and pretreatments vs. untreated) (Fig. 4).

Changes in fibre composition are also supported by FTIR analysis, which has already been used to study the structural characteristic of ligno-cellulosic material [34]. Seven bands of particular relevance were analysed. Their spectra referring to untreated, organic and mineral plus organic acid at high concentration (OAc.H and M+OAc.H, respectively) are displayed in Fig. 5; they are associated with the functional groups and compounds indicated in Table 2, according to the cited sources. In our experiment the bands assigned to H-cell and Cell exhibited a decrease in absorbance after pre-treatment (Table 4), supporting a solubilisation of the two structural carbohydrates and a change in fibre structure [33]. TCI and LOI, indicating the overall degree of order of Cell and the mount of crystalline vs. amorphous Cell, respectively, were augmented after pre-treatment. This means that only amorphous Cell was likely solubilised (Table 4), although these changes had positive effects on substrate biodegradability, speeding the hydrolytic phase and enhancing final methane production.

#### 4. Conclusions

Maleic acid, organic catalyst in the frame of a biomimetic approach to pre-treatments, was tested to improve methane yield from three ligno-cellulosic substrates. Remarkable physical and chemical changes of fibre structure occurred after pre-treatment, supporting the hypothesis of enhanced substrate degradation. This in turn determined increased methane yield, to an extent directly related to substrate recalcitrance: +41, +31 and +24% in the respective Arundo, Barley straw and B133 sorghum (averages of the four pre-treatments).

Although a considerable increase in methane output was evidenced, appraisal of costs and benefits is mandatory, before the implementation of such treatments may be envisaged in full scale biogas plants.

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