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Application of Enzymatic and Bacterial Biodelignification Systems for Enhanced Breakdown of Model Lignocellulosic Wastes

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8

9 ABSTRACT

10 This paper explores the extent to which enzymatic and bacterial biodelignification

11 systems can breakdown lignocellulose in model wastes to potentially enhance biogas

12 generation. Two representative lignocellulosic wastes (newspaper and softwood)

13 commonly found largely undegraded in old landfills were used. A fungal peroxidase (lignin

14 peroxidase) enzyme and a recently isolated lignin-degrading bacterial strain

15 (Agrobacterium sp.) were used. Tests were conducted in stirred bioreactors with

16 methanogens from sewage sludge added to produce biogas from breakdown products.

17 Addition of lignin peroxidase resulted in ~20% enhancement in cumulative methane

18 produced in newspaper reactors. It had a negative effect on wood. Agrobacterium sp. strain

19 enhanced biodegradation of both wood (~20% higher release of soluble organic carbon and

20 enhanced breakdown) and newspaper (~2-fold biogas yield). The findings of this paper

21 have important implications for enhanced breakdown in old landfills that are rich in these

22 wastes, and anaerobic operations utilising lignocellulosic wastes for higher degradation

23 efficiencies and biogas production.

24 KEYWORDS: Lignocellulose, peroxidase enzyme, bacterial biodelignification

25 1 INTRODUCTION

26 The rate of landfill biogas production is initially high as the easily degradable fraction is 27 broken down, the rate then decreases with a subsequent long 'tail' of emissions typically 28 observed. This happens due to the accumulation of waste fractions that are difficult for 29 landfill microbiota to break down, i.e. lignocellulose-containing wastes (Barlaz et al., 30 1989). Moreover, these slow rates of gas generation after the easily-degradable matter has 31 depleted are insufficient for energy generation or flaring, and so the biogas typically 32 escapes into the atmosphere. As landfill biogas mainly comprises of methane and carbon 33 dioxide, which are two major contributors to climate change (O'Dwyer et al., 2017), this is 34 not a sustainable situation. 35 The recalcitrance of lignocellulosic wastes results from the presence of lignin, which is a 36 complex heteropolymer and the second most abundant biopolymer in nature. Due to its 37 very complex structure, including ether and C-C double bonds, phenolic monomers etc, 38 most microorganisms found in landfills find it extremely difficult to breakdown this 39 polymer (Bugg et al., 2011a; Cragg et al., 2015). Lignin also forms a 'glue'-like structure 40 around other easily degradable matter (e.g. cellulose, hemicellulose) in lignocellulosic 41 wastes to, in effect, protect against microbial attack, hence further decreasing the 42 bioavailability of the biomass (Brandt et al., 2013). 43 Accelerating breakdown of lignin and subsequent methanogenesis in the waste body 44 would help to confine methane emissions to a shorter period of higher methane 45 concentration in biogas, allowing more of the landfill gas to be used for energy 46 generation/stopping escape into the atmosphere and reducing the long 'tail' of low 47 emissions. This would also allow more rapid stabilization of sites and so reduce long-term

management costs for operators. Lignocellulosic wastes are also diverted to anaerobic
digestion (AD) plants for production of biogas (Wyman et al., 2017). However, the same
problem of recalcitrance of these wastes to biodegradation makes it difficult to utilise their
full potential in AD operations (Hassan et al., 2018). Again, accelerating the breakdown of
lignocellulose could also offer better biogas recovery resulting in improved biogas yields
from lignocellulosic waste.

54 This acceleration may be achieved by biotechnological methods. For instance, some microorganisms have evolved specifically to degrade lignin, with white-rot fungi being one 55 56 of the few well-studied organisms (Bugg et al., 2011a). White-rot fungi produce 57 extracellular peroxidase enzymes to break down lignin. However, the process is extremely 58 slow due to slow fungal growth over the biomass (Bugg et al., 2011a; Bugg et al., 2011b; 59 Cragg et al., 2015) and is limited in typical waste environments such as landfill due, for 60 example, to a lack of oxygen (Geoffrey, 2003; Leonowicz et al., 1999). Some bacteria are 61 also able to metabolise lignocellulose (Mathews et al., 2015; Rahmanpour et al., 2016; Xu 62 et al., 2018) and have been shown to be more flexible in terms of the conditions under 63 which they can operate (Bugg et al., 2011a; Bugg et al., 2011b; Cragg et al., 2015; Rashid 64 et al., 2017).

Acceleration of breakdown of lignin-rich materials will require more rapid enzymatic activity, which can be addressed in two ways: (i) extracellular enzymes similar to those produced by white-rot fungi, which are available commercially, can be added to the system thereby eliminating the need to use the organism itself in biodelignification (Hettiaratchi et al., 2014; Hettiaratchi et al., 2015; Jayasinghe et al., 2011; Jayasinghe et al., 2014; Jayasinghe et al., 2013). Researchers have applied this technique of enzymatic

71 biodelignification with the prospect of enhanced recovery of chemicals/biogas with varying 72 degrees of success (Schroyen et al., 2017; Schroyen et al., 2014; Schroyen et al., 2015) 73 although not to realistic, complex waste materials such as wood or newspaper. (ii) a second 74 approach is the application of bacteria for delignification, this has received only very 75 limited attention but is gaining popularity and shows promise in enhancing the breakdown 76 of lignocellulosic wastes for various biotechnological applications (e.g. biogas generation, 77 bioethanol production, renewable chemicals) (Bugg et al., 2011a; Bugg et al., 2011b; 78 Mathews et al., 2015; Rashid et al., 2017). However, the application of this technique to 79 complex waste materials likely to be found in significant quantities in landfills is yet to 80 attract significant attention. 81 Many of the aforementioned studies have been carried out under highly-controlled

conditions (e.g. highly buffered systems) and with idealised materials (e.g. model lignin
molecules, dissolved pure kraft lignin). The question of how well these enzymes and
bacteria can breakdown lignocellulose in realistic waste materials found typically
undegraded in landfills, with the goal of enhancing gas production and hence stabilising the
waste mass is considered in this study.

The objective of this paper is to test the hypothesis that certain enzymes and bacteria can break down lignocellulose in complex and realistic waste materials, found to be largely undegraded in landfills (Ximenes et al., 2015; Ximenes et al., 2017; Ximenes et al., 2008), and so enhance rates of biogas production. Conversion of breakdown products to biogas through methanogenic activity (provided through addition of methanogen-rich sewage sludge) is recorded as volume of gas produced, alongside key parameters of the system (chemical oxygen demand, organic carbon, pH etc.). As part of the bacterial

94	biodelignification experiments, the hypothesis that the solid:liquid ratio might also play an
95	important role in the ability of the Agrobacterium sp. strain to enhance lignocellulosic
96	breakdown and subsequent methanogenesis is also tested.

2 MATERIALS & METHODS

Two waste materials are considered: newspaper (52 g/ m^2 standard recycled paper) and 98 99 softwood (kiln-dried Nordic redwood pine timber, high heartwood proportion). For 100 particle-size reduction, prior to starting an experiment, approx. 200g of waste were fed to a 101 Fritsch 55743 rotary knife mill (approx. feed size: 2 cm long squares for newspaper and 102 cubes for wood) with a 2mm screen. For experiments requiring powdered wood, the knife-103 milled material that passed through the 2mm screen was collected and passed through a 104 0.15 mm sieve. These wastes were chosen to represent the range of lignocellulosic wastes 105 found in MSW landfills, with newspaper containing a smaller amount of lignin in 106 comparison to softwood which is very lignin-rich. 107 Sewage sludge from an anaerobic digester at a wastewater facility in Cardiff, was used as 108 a source of methanogens $(5.52\% \pm 0.002 \text{ dry solids of which } 59.46\% \pm 0.023 \text{ volatile})$ 109 solids) in order to model the presence of methanogens in the landfill environment. Sewage 110 sludge was used due to its reported relatively more homogeneous behaviour as an inoculum 111 across the literature for anaerobic biodegradation in comparsion to landfill leachate, which 112 varies much more widely around the world (Pearse et al., 2018). The sludge was sampled 113 from the digester in three 5-litre high-density polyethylene jerrycans, immediately 114 transferred to the lab and used to start the bioreactor experiments. For experiments where 115 the bacteria was acclimatisation at 30°C for three days prior to starting an experiment, the 116 jerrycans were immediately taken to the lab from the site, transferred to an incubator

117 maintained at 30°C and shaken manually twice a day to ensure homogenisation. For every 118 experimental run, fresh sludge was sampled from the same site and sampling point. 119 Commercially available lignin peroxidase (Merck product code: 42603-10MG-F) was 120 used in the enzyme delignification experiments. Agrobacterium sp. (GenBank accession 121 JX872342, bacterial phylum α-Proteobacteria), supplied by T. Bugg (Warwick University, 122 UK), which is a facultative anaerobic lignin degrader isolated from landfill soil that grows 123 optimally at 30 °C (Rashid et al., 2017) was used in the bacterial biodelignification 124 experiments. 125 2.1 **Bacterial Culture** 126 The Agrobacterium sp. cultures were maintained on Luria-Bertani (LB) agar. The

127 bacterium was cultured in LB broth at 30 °C and the cultures were harvested (centrifugation

128 for 10 min at 3394 rcf – relative centrifugal force) in the exponential phase according to its

129 growth curve. The cultures were then washed with M9 mineral media $(6.78g/l Na_2HPO_4,$

130 3 g/l KH_2PO_4 , 1 g/l NH_4CL , 0.5 g/l NaCL) to remove any carbon from the LB broth,

131 followed by centrifugation again and resuspension in M9 before addition to the bioreactors.

132 2.2 Experimental Design

Preliminary Small-scale Bacterial Biodelignification: To identify optimal conditions for later bioreactor experiments on wood a series of preliminary experiments were undertaken. This is particularly critical for wood due to the typically lengthy degradation time-scales (experiments lasting years in some studies) (Wang et al., 2013) and the lignin content of wood and its various sizes found undegraded in landfills (small cm-scale chips to m-scale blocks) (De la Cruz et al., 2013; Wang & Barlaz, 2016; Ximenes et al., 2017; Ximenes et al., 2018). Since wood contains roughly 1.5 times more lignin than newspaper and is unprocessed lignocellulose, where the structure is intact, it represents some of the most
difficult to degrade wastes in landfills and should prove to be a more challenging substrate
for biodegradation. To get an understanding of the impact of the physical state of the wood
on biodegradation by *Agrobacterium sp.*, small-scale tests in 50 ml sterile tubes were
conducted.

145 Four wood sample types were tested in these experiments exploring the impact of waste 146 form/size and accessibility of biodegradable materials. Each consisted of one particle size 147 range (<2 mm or <0.15 mm) and was either tested as-is or autoclaved prior to the 148 experiment. In the latter case wood was autoclaved at 120°C for 15 mins to test its effect on 149 deconstructing the lignocellulosic matrix for easier microbial access to key polymers, i.e. 150 lignin, cellulose and hemicellulose (Pecorini et al., 2016). Each type was then tested with 151 and without Agrobacterium sp. in duplicate specimens. Each specimen comprised 1 g of 152 wood added to 20 ml of M9 mineral medium in 50 ml tubes with either 1 ml of 153 Agrobacterium sp. starter culture or 1 ml sterile M9 medium (control) (Rashid et al., 2017). 154 Flasks were placed in a shaking incubator at 200 rpm and 30°C for seven days. For these 155 small-scale tests, sludge was absent since the purpose was to test the ability of this strain to 156 breakdown lignocellulose, whilst the presence of other microbial communities from the 157 sludge could interfere with the monitoring of the activity of only this strain. Total carbon 158 released into the liquid phase during the test and the organic carbon content of the solid 159 residue after bacterial treatment were analysed. Biogas production, pH and chemical 160 oxygen demand were not monitored in these preliminary small-scale tests. 161 Bioreactor System: The experimental apparatus (supplied by Anaero Technology UK) 162 consisted of 15 one-litre reactors submerged in a water bath maintained at the required

163 temperature. Gas flow meters based on the water displacement method (Wickham et al., 164 2016) combined with an Arduino (for data logging) were used to monitor biogas 165 production. Biogas was collected in 5 litre Tedlar® bags attached to the outlets of the gas 166 flow meter for each reactor, and all biogas data are reported at STP. The reactors were 167 continuously stirred at 45 rpm during the experimental work and the duration of the 168 experiments was between 30-35 days. 169 Bioreactor-scale Bacterial Biodelignification (Experiment 1): Mechanistic insights from 170 the preliminary small-scale tests were used to inform the design of larger-scale 1 L (total 171 volume) tests. The rationale behind larger-scale experiments was to be able to study 172 whether enhanced biodegradation could be carried out using the Agrobacterium sp. whilst 173 monitoring for key variables such as biogas production, release of organic carbon etc, all of 174 this done in the presence of methanogenic microbial communities from sewage sludge. 175 These bacterial experiments contained 4g of waste, and employed a mixture of sludge and 176 M9 minimal media (for the agrobacterium sp. (Rashid et al., 2017)). 177 Each bioreactor comprised 600 ml M9 medium (650 ml in controls), 50 ml suspension 178 of Agrobacterium sp. (0 ml in controls), 50 ml methanogen-rich sewage sludge 179 (acclimatised at 30 °C for three days prior to introduction) and 4g of lignocellulosic 180 material (wood or newspaper), together with 300 ml of headspace. The headspace 181 comprised of air due to Agrobacterium sp. being a facultative anaerobe, i.e. able to work 182 under anaerobic conditions but preferring aerobic conditions (Rashid et al., 2017; Taylor et 183 al., 2012). This reflects typical conditions in bioreactor landfills, where the waste mass is 184 initially aerobic and the system slowly goes anaerobic with the passage of time (Benson et 185 al., 2007). Five cases were tested (Table 1), each in triplicate, with Agrobacterium sp.

186	applied to vessels containing wood (<0.15 mm, non-autoclaved), newspaper and no waste,
187	whilst control vessels with only methanogens (no Agrobacterium) were supplied with wood
188	or newspaper. All bioreactors were incubated in the water bath apparatus at 30 °C for 35
189	days. Liquid samples (5 ml) were taken periodically for analysis (section 2.3) from the
190	sampling port using sterile pipettes and transferred to 15 ml sterile containers. Solid residue
191	was obtained from all bioreactors by collection on Whatman No. 42 $\ensuremath{\mathbb{R}}$ filter paper (2.5 μm)
192	via vacuum filtration then drying at 105 °C. Biogas collected during the experiment in 5
193	litre Tedlar gas bags was analysed for methane content.
194	Progressively diminishing rates of biogas and organic carbon release for newspaper
195	reactors prompted a 'restarting' of the experiment, where the contents of the bioreactors
196	were centrifuged at 3394 rcf for 10 min under aseptic conditions to remove the supernatant
197	and fresh autoclaved M9 was added in the same amounts as the beginning of the test (see
198	above). This allowed for the hypothesis that towards the end of the experiment (whilst there
199	is still carbon present in the liquid phase but gas production has almost stopped),
200	accumulation of recalcitrant or toxic substances in the liquid phase inhibits further
201	conversion/breakdown to biogas to be tested. 'Restarting'/removing the liquid and
202	introducing a fresh medium could be beneficial and allow for biogas production to continue
203	again from the left-over solid residue. 'Restarted' bioreactors were incubated under the
204	same conditions for 20 additional days, with liquid and gas sampling.
205	Impact of Solid:Liquid Ratio on Bacterial Biodelignification (Experiment 2): Previous
206	work has shown that microbial activity is affected by the waste to inoculum ratio (with
207	particularly low ratios shown to inhibit methanogenesis) (Moset et al., 2015). As mentioned
208	previously, to test the hypothesis that the solid:liquid ratio might also play an important role

in the ability of the *Agrobacterium sp.* to breakdown lignocellulose, the impact of the
Solid:Liquid (S:L) ratio on bacterial biodelignification of these wastes was also studied by
increasing the mass of waste added to 12 g. The experiments (Table 1) are otherwise the
same as the earlier bioreactor-scale tests, apart from there not being a 'waste-free' control
triplicate. A 4 g test (for wood and newspaper respectively) was also carried out as a repeat

of that in the previous experiment to allow comparability and to serve as controls to test theabove hypothesis.

216 <u>Bioreactor-scale Enzymatic Biodelignification (Experiment 3)</u>:. 1 L (total volume) tests

to study whether enhanced biodegradation could be achieved using lignin peroxidase.

218 Treatments are summarised in Table 1. 4 g of waste ≤ 2 mm, i.e. newspaper or wood (non-

autoclaved), and 2.22 mg of lignin peroxidase (LiP) were added to 600 ml of sludge (source

of methanogens) for a volatile solids ratio of sludge to waste of 4:1 (Labatut et al., 2011).

221 The reason for a much larger sludge volume here pertains to the fact that inoculum to

substrate ratio is a major factor in the determination of the biomethane potential of a

substrate (Raposo et al., 2006), hence the enzyme experiments were conducted at the ideal

volatile solids ratio of the sludge to the waste based on previous studies (Moset et al., 2015;

225 Peña Contreras et al., 2018). The chosen waste:LiP mass ratio was based on previous work

226 (Hettiaratchi et al., 2014; Hettiaratchi et al., 2015; Jayasinghe et al., 2011; Jayasinghe et al.,

227 2014; Jayasinghe et al., 2013) where the volatile solids (VS):LiP ratio was optimised for

maximum biogas production. LiP was activated with H_2O_2 prior to addition at the optimal

LiP: H_2O_2 ratio (Hettiaratchi et al., 2014; Jayasinghe et al., 2011; Jayasinghe et al., 2013).

230 The controls comprised of exactly the same experimental setup but lacked the peroxidase

enzyme in 'waste-containing' controls. 'Waste-free' controls were also carried out with 0

- and 2.22 mg of LiP. All bioreactors were incubated at 38 °C, as this is the optimal
- 233 operational temperature employed by the anaerobic digester used as the source of
- 234 methanogens.

	Experiment 1		Experiment 2		Experiment 3	
	With Agrobacterium	Without Agrobacterium	With Agrobacterium	Without Agrobacterium	With enzyme*	Without enzyme*
No waste	3				3	2
Newspaper (4 g)	3	3	3		3	2
Newspaper (12 g)			3			
Wood (4 g)	3	3	3		3	2
Wood (12 g)			3			

235 Table 1. Conditions tested in the three main experiments. Numbers refer to number of replicates for each treatment.

Note: Wood sample size < 0.15 mm, apart from samples denoted with * which had size < 2mm. Other experimental conditions are as described in the text.

236 **Table 2.** Waste Characterisation.

Sample	VS (%)	OC (%)	Lignin (%)
Wood	99.62 ± 0.00	50.60 ± 0.40	38.45 ± 0.13
Newspaper	84.26 ± 0.00	36.72 ± 8.09	25.73 ± 0.89

237 *Note: Data presented as percentage of dry mass.*

238 2.3 Analytical Methods

239 Liquids: All 5 ml liquid samples taken during bioreactor experiments were filtered 240 through sterile 0.2 μ m filters prior to analysis. The pH was measured on the entire sample 241 post-filtration according to standard methods (APHA, 2012). The Folin-Ciocalteau method 242 was used for measuring phenol release (Meda et al., 2005; Rashid et al., 2017). 0.8 ml of 243 deionized water and 0.5 ml Folin-Ciocalteau's reagent (Merck F9252) were added to 0.2 ml 244 filtered liquid sample. 2.5 ml of 20% Na₂CO₃ was then added and the samples incubated in 245 the dark for 30 min. Then the absorbance was measured using a spectrophotometer at 760 246 nm and p-hydroxybenzoic acid was used to calibrate the absorbance as a standard. A 1 ml 247 portion of the filtered sample was used for soluble organic carbon (sOC) analyses via a 248 Shimadzu TOC-VCPH following the manufacturer's instructions whilst 0.5 ml was used 249 for total and soluble chemical oxygen demand (COD and sCOD) analyses and 1ml was 250 used for determining total/dry solids (TS and DS) for sludge characterisation all according 251 to standard methods (APHA, 2012). 252 Solids: Lignin content and solid organic carbon analyses were carried out on the solid 253 residue collected post-experiment. Lignin content analysis on the solid residue was adapted 254 from Rashid et al., 2017. The samples were dried at 105°C until constant weight, then 0.25 255 g of dry mass was added to 3.75 ml of 95% H₂SO₄ with stirring for 2 h at RTP. Then 140 256 ml of deionised water was added to the resulting solution followed by reflux for 4 h in 257 round bottom flasks. The residual content was collected via Whatman No. 42 ® filter paper 258 $(2.5 \,\mu\text{m})$ and washed with deionised water. Then the residue was dried at 105°C until 259 constant weight. The residue was then volatilised at 550°C to correct for ash. The lignin

260 content was determined by subtracting the mass of ash from that of the dry residue and then

- 261 calculating the ratio of this to the original sample dry mass. The solid total/organic carbon
- 262 (TC/OC) were measured via a Shimadzu TOC-VCPH following the manufacturer's
- 263 instructions. Volatile solids (VS) analysis was carried out according to standard methods.
- 264 <u>Gases:</u> The methane concentration of the biogas was determined via gas chromatography
- 265 (GC) analysis on a Varian 450 GC equipped with a flame ionisation detector and a
- 266 methaniser using a CP SiL5 CB column (50m, 0.33mm diameter, He carrier gas).
- 267 2.4 Statistical Analyses
- 268 Two-way ANOVA and Student's T-test were carried out. Statistical significance was
- attributed at P<0.05 (i.e. ANOVA F-value >F-critical). Correlations between data
- 270 (specifically for the sCOD and sOC values) were identified via Pearson correlation initially
- and plotted against each other to further study the correlations.
- 272 **3 RESULTS & DISCUSSION**

273 **3.1 Waste Characterisation**

274 The composition of the model wastes is shown In Table 2. Volatile solids content of a 275 substrate indicates the fraction that has the potential to be converted to biogas (Barlaz et al., 276 1989; Eleazer et al., 1997; Wang et al., 1994). The wood was almost entirely composed of 277 volatile matter, whereas newspaper had some ash content. This is comparable to recent 278 work studying the chemical composition of different wastes (Chickering et al., 2018; 279 Krause et al., 2017; Krause et al., 2016). As the aforementioned studies also suggested, ash 280 content likely comes from fillers and ink constituents within the newspaper. The wood was 281 very rich in lignin and also has a higher organic carbon content compared to the newspaper. 282 Bearing in mind the variability in waste composition that arises around the world, in

relation to recent work, these values fall within the typical ranges reported (De la Cruz etal., 2014; Wang & Barlaz, 2016).

285 **3.2 Bacterial Enhancement**

286 Small-scale: Total carbon release profiles from the preliminary small-scale experiments 287 conducted on the non-autoclaved and autoclaved ≤ 0.15 mm and ≤ 2 mm wood are shown in 288 Figures 1a and 1b. In the presence of the Agrobacterium sp., a steady and significant 289 increase in the carbon present in the liquid phase occurred for the non-autoclaved ≤ 0.15 290 mm samples (black circle, Fig. 1a)) (~20% relative to the control, P<0.05, ANOVA F-291 values>F-critical) during the 7-day experiment. Since the only source of carbon in these 292 experiments was lignocellulose from wood, and the soluble carbon content was determined 293 post-filtration through 0.2 µm pore-size filters (even bacterial cells should not be passing 294 through into the supernatant), it is highly likely that the release of carbon corresponds to 295 biodegradation due to the bacterium. The profiles for the non-autoclaved < 0.15 mm wood 296 controls (grey circle, Fig. 1a)) remain steady and virtually unchanged, suggestive of no 297 activity in the absence of the bacterium. No significant impact on the sTC release profiles 298 due to the Agrobacterium sp. on 2 mm particles (Fig. 1b, autoclaved and non-autoclaved) 299 was recorded (P>0.05, ANOVA F-values<F-critical). The sTC profile in the presence of 300 the Agrobacterium sp. is consistently lower for 2 mm samples (non-autoclaved and 301 autoclaved, dotted lines denote bacterium augmentation), which could be explained by the 302 bacterium metabolising leached carbon from the wood. 303 It is noted that the <0.15 mm autoclaved wood containing the *Agrobacterium sp.* (black 304 dotted line, Fig. 1a) starts at a lower sTC on day 0 (black dotted line, Fig. 1a), however, the

305 increase in total sTC released in solution is ~30% higher in comparison to the increase

found in the control, hence indicating lignocellulosic breakdown of the solid matrix due tobacterial enhancement.

308	To confirm the liquid phase carbon release results, the organic carbon analysis of the
309	solid residue post-treatment with the Agrobacterium sp. for wood under different
310	conditions is shown in Figure 1c. The organic carbon in ≤ 0.15 mm samples is significantly
311	lower (two-way ANOVA, P<0.05, F-values>F-critical) in the treated samples for both
312	autoclaved and non-autoclaved wood samples (by 8.6 and 7.5% respectively), indicating
313	higher levels of breakdown of the lignocellulosic structure due to Agrobacterium sp.
314	treatment. With 2 mm wood there is no significant difference due to bacterial enhancement.
315	It is expected that the much higher surface area to volume ratio, and accessibility of the
316	<0.15 mm samples is the key cause of the much better solid OC breakdown due to
317	Agrobacterium sp. treatment in this set of samples in comparison to the 2mm size.
318	Overall, the preliminary small-scale data suggest that Agrobacterium sp. can break down
319	woody lignocellulose, and that the particle size is an important factor in the rate of reaction
320	when applying this Agrobacterium sp. strain for biodegradation. Treating wood by
321	autoclaving does not significantly impact the biodelignification ability of this bacterium.



323 Figure 1. Assessing the impact of Agrobacterium sp. treatment on wood lignocellulose

324 under different conditions. (a) Soluble liquid total carbon release results for the small-scale

325 0.15 mm tests. (b) Soluble liquid total carbon release results for the small-scale 2mm tests.

326 (c) Solid phase organic carbon results for the small-scale experiments. All error bars

327 represent +/- 1 standard deviation. Abbreviations: Agrobacterium, A; Autoclaved, AT;

328 Wood, W; M9 solution, M9.

329 <u>Bioreactor-scale (Experiment 1):</u> The cumulative biogas profiles released from

newspaper in Experiment 1 are shown in Figure 2a. After a lag-phase of ~7-8 days,

331 Agrobacterium sp. nearly doubled the biogas production in newspaper reactors compared to

332 controls without Agrobacterium ~92% enhancement, P<0.05) whilst no response was

observed without waste materials. At the same time, the sCOD and sOC profiles (Figure 2c,

2d) exhibit a consistently lower amount of organic carbon in solution with newspaper and

335 Agrobacterium sp. than in newspaper controls. These data suggest that the Agrobacterium

sp. enhance the release of carbon from the solid phase and increase the conversion of

dissolved carbon to biogas. In controls with newspaper, the total amount of OC released

into solution is ~ 0.32 g whilst the OC released in total as biogas is 0.04 g, giving a total of

 ~ 0.36 g. In specimens with *Agrobacterium sp.* and newspaper, ~ 0.31 g of OC was released

into solution by the end of the experiment, but the amount of biogas was 0.08 g, giving a

total of ~ 0.39 g. These data indicate greater release of solid organic carbon into solution

342 and greater conversion of that soluble OC to biogas in the presence of Agrobacterium sp.,

- 343 leaving less in solution. With 1.4 g of solid OC present initially, in the presence of the
- 344 Agrobacterium sp. there is an $\sim 8.2\%$ increase in the total release of OC in relation to the

345 control. The observed effects most likely result from the Agrobacterium sp. breaking down 346 either solid phase or leached organic materials into a form utilisable by the methanogens. 347 For the newspaper reactors, the sOC and sCOD profiles increase gradually with 348 increasing biogas production and result in overall accumulation towards the end of the 349 experiment of organic carbon in the liquid phase. It is hypothesised that the cause was 350 accumulation of recalcitrant organics and other degradation products which are not 351 converted to the gas phase and which hinder the activity of the microorganisms. Therefore, 352 an attempt was made to 'restart' the experiments by replacing the liquid fraction with fresh 353 autoclaved M9 on Day 35. Following this, however, little to no additional activity was 354 observed (Figure 2). As such, the flat-lining of the biogas and sCOD/sOC curves is likely to 355 be due to a different limiting factor. 356 No significant effect was observed on biogas production due to the addition of the Agrobacterium sp. strain to wood (Figure 2b). Similarly, no significant increase in sCOD or 357

358 sOC was observed in this experiment. A possible explanation for this, and an important

difference between the preliminary small-scale and bioreactor experiment 1 is the

360 concentration of wood in the overall volume. For the small-scale wood biodegradation

tests, the S:L ratio was 0.05 g/ml, and a significant impact of bacterial treatment on the

- 362 woody lignocellulose was observed, whereas here it was 0.006 g/ml (~8.3 times more
- 363 dilute). It is possible that the wood was not sufficiently concentrated in the bioreactors to
- 364 either observe a measurable response or to stimulate microbial activity.



367 Figure 2. Assessing the impact of Agrobacterium sp. treatment on newspaper and wood 368 lignocellulose in the presence of sludge. (a) Cumulative biogas volume results for 369 newspaper. (b) Cumulative biogas volume results for wood. (c) Soluble chemical oxygen 370 demand profiles. (d) Soluble organic carbon profiles. Note: system 'restarted' on Day 35, 371 last liquid sampling point was at day 26 before the 'restart'. All error bars represent +/- 1 372 standard deviation. Abbreviations: Agrobacterium, A; Methanogens, M; Newspaper, N; 373 Wood, W; M9 solution, M9. 374 Impact of increasing solid:liquid (S:L) ratio (Experiment 2): The lack of a response from 375 wood exposed to Agrobacterium sp. in the first bioreactor experiment was hypothesised to 376 be due to a combination of a slow biodegradation rate and a low S:L ratio. Increasing the 377 S:L ratio from 0.006 g/ml to 0.017 g/ml had a significant impact on the cumulative biogas 378 volume for both lignocellulosic wastes (Figure 3a -wood, Figure 4a -newspaper). 379 In the case of wood, the biogas for the higher S:L ratio was increased by $\sim 205\%$. 380 Changing the S:L ratio has been shown to have a significant impact on the biogas kinetics 381 and cumulative volume of the organic fraction of MSW (Krause et al., 2017; Krause et al., 382 2016; Raposo et al., 2006). Although, to the best of our knowledge, this has not been 383 investigated in the application of bacterial biodelignification systems on wood 384 lignocellulose. The wood sCOD profile (Figure 3c) increases and reaches a peak around 385 day 14 then decreases gradually. The sOC profile (Figure 3b) does not seem to strictly 386 follow this exact trend of reaching a peak followed by gradual decrease. sCOD is indicative 387 of the amount of oxygen required to oxidise the carbon present, and the more oxygen 388 required, the more carbon is present in solution or roughly the same amount is present, but 389 in a more complex structure which is more difficult to oxidise. The general trend in sCOD

390 is an increase then a gradual decrease. The sOC increases and then stays reasonably steady 391 with a higher value measured in the wood cases. A possible explanation of these 392 observations might have to do with the form of the carbon released. The fact that sCOD 393 lowers but the sOC does not perhaps indicates that the form of the soluble compounds 394 might have gone from more complex structures, to more simple structures that may be 395 relatively easier to convert to biogas, i.e. end-product of anaerobic digestion. 396 It is important to note that at the very high S:L of 0.05 g/ml in the small-scale tests, a 397 very flat line for the total carbon profile for the ≤ 0.15 mm wood controls was recorded. 398 Whereas here, the sCOD profile due to the addition of the bacterium is transient and 399 follows a logical pattern of lignocellulosic breakdown (i.e. ~31% increase in the first two 400 weeks) and methanogenesis to convert that carbon to the gas phase (i.e. evidence from 401 biogas production profile, and decrease of sCOD by ~16% relative to the maximum peak-402 value reached in the first two weeks). As such, it is suggested that the liquid analyses 403 results discussed here are most likely due to the Agrobacterium sp. acting in synergy with 404 the microbes from the sludge for depolymerisation of wood. 405



407	Figure 3. Assessing the impact of increasing S:L ratio on Agrobacterium sp. treatment of
408	wood lignocellulose with sludge. (a) Cumulative biogas volume for wood. (b) Soluble
409	organic carbon profiles for wood. (c) Soluble chemical oxygen demand profiles for wood.
410	All error bars represent +/- 1 standard deviation. Abbreviations: Agrobacterium, A;
411	Methanogens, M; Wood, W; M9 solution, M9; 4 grams mass of waste, 4g; 12 grams mass
412	of waste, 12g; PrecedingRun, data from the preceding run from Fig.2, identical
413	experimental conditions to S:L 0.006 and included to account for inter-experimental

414 variability, since sludge for the S:L experiments was sampled at a later date.



416	Figure 4. Assessing the impact of increasing S:L ratio on Agrobacterium sp. treatment of
417	newspaper lignocellulose with sludge. (a) Cumulative biogas volume for newspaper. (b)
418	Soluble organic carbon profiles for newspaper. (c) Soluble chemical oxygen demand
419	profiles for newspaper. All error bars represent +/- 1 standard deviation. Abbreviations:
420	Agrobacterium, A; Methanogens, M; Newspaper, N; M9 solution, M9; 4 grams mass of
421	waste, 4g; 12 grams mass of waste, 12g; PrecedingRun, data from the preceding run from
422	Fig.2, identical experimental conditions to S:L 0.006 and included to account for inter-
423	experimental variability, since sludge for the S:L experiments was sampled at a later date.
424	In Figure 4b and 4c the newspaper sOC and sCOD profiles are shown respectively. In the
425	case of newspaper, the biogas volume was 280% greater than that at the lower S:L ratio.
426	The cumulative overall volume for newspaper at the lower S:L ratio was significantly
427	different (P < 0.05; approximately double) to that obtained previously with the same
428	conditions (Figure 4a). This may have arisen due to variability in the newspaper
429	composition (a different sub-sample was milled for this test) or variability in the microbial
430	community from the sludge, since this was sampled 4 months after the previous test, albeit
431	from the same site/digester/sampling point. However, it is clear that with all else equal the
432	higher S:L ratio gives a significantly greater degree of biodegradation. For newspaper, the
433	sOC and sCOD profiles follow a similar trend to the previously reported Figure 2b, 2c.
434	To study the impact of S:L ratio on the activity of the Agrobacterium sp. and overall
435	biogas generation as a continuum, this ratio could be further increased or decreased under
436	similar experimental conditions. This would perhaps allow for optimisation of the
437	microbial processes studied. However, since the aim of this paper has been to gain
438	mechanistic insights, optimisation has been beyond the scope of this study.

439 From the above bioreactor experiments, it is obvious that under all conditions, newspaper

440 clearly produces much higher biogas yields than wood and shows a greater degree of

441 enhancement (Fig. 2, 3, 4). This is likely due to a combination of two main factors. Firstly,

442 due to the much higher lignin content of softwood (~1.5 times more than newspaper)

443 (Table 2), which results in more recalcitrant biomass in comparison to wood. Secondly,

444 newspaper, is a mechanical pulp which in relation to softwood is more processed, this

445 likely also results in better accessibility to key polymers for degradation (Baldwin et al.,

446 1998; Barlaz, 2006; Eleazer et al., 1997; Stinson & Ham, 1995; Wang et al., 1994).

447

3.3 Enzymatic Enhancement

In Figures 5 and 6 biogas profiles/values and net biomethane yields per gram of VS due to the action of LiP are shown. A small positive effect on biogas/biomethane yields due to the enzyme for just the bioreactors containing sludge and no waste took place (P<0.05) (Figures 5 and 6). This is likely a result of the enzyme breaking down suspended and/or dissolved recalcitrant organics present in the sludge which would otherwise not be broken down by the microbial communities present, which can then be utilised for metabolism and eventually be converted to biogas.

455 A significant (P<0.05) enhancement in biomethane potential from newspaper of \sim 41%

456 was achieved with LiP present. LiP is thought to form reactive free radicals that attack the

457 non-phenolic parts of the lignocellulosic structure (Bugg et al., 2011a; Bugg et al., 2011b;

458 Cragg et al., 2015). The enhancement is possibly a result of the enzyme attacking the

459 lignocellulosic structure of newspaper, thereby providing additional substrate to the

460 microorganisms from the sludge which can then be converted to biogas.

- 461 In the enzyme-containing bioreactor cases discussed above, in addition to the overall
- 462 biogas volume being enhanced, the proportion of methane also seems to be higher in the
- 463 presence of the enzyme. This observation suggests that enzymatic enhancement in these
- 464 reactors not only increased the overall biogas yield, but also the specific methane
- 465 production, thereby increasing its concentration in the biogas mixture.



- 467 **Figure 5.** Cumulative biogas generation profile results. Assessing the impact of lignin
- 468 peroxidase application on newspaper and wood. (a) Sludge only. (b) Newspaper. (c) wood.
- 469 All error bars represent +/- 1 standard deviation. Abbreviations: Sludge, S; lignin
- 470 peroxidase enzyme, E; Newspaper, N; Wood, W. Inset figures included for closer
- 471 examination of the gas generation profiles towards the end of the test.



473	Figure 6. Biogas and methane yield results. Assessing the impact of lignin peroxidase
474	application on newspaper and wood. (a) Cumulative biogas volume comparison for
475	bioreactors. (b) Cumulative methane generated in bioreactors over the 30-day experimental
476	run. (c) Net methane yield per gram of VS. All error bars represent +/- 1 standard deviation.
477	In bioreactors containing wood, there was a significantly lower amount of biomethane
478	production (P<0.05, Figures 5,6) in the enzyme-augmented reactors compared to the
479	control. This effect was also found by Schroyen et al. (Schroyen et al., 2017; Schroyen et
480	al., 2014; Schroyen et al., 2015), where not all lignocellulosic substances responded
481	positively to pre-treatment through peroxidase enzymes prior to anaerobic digestion, in
482	some instances the treatment negatively impacted methane production (e.g. with corn
483	stover, wheat straw, maize). No significant changes in phenolics (which at high
484	concentrations can inhibit methanogenesis (Hernandez & Edyvean, 2004)), TCOD or any
485	transient changes in pH were observed within the first two weeks in comparison to the
486	controls (data not shown). At low concentrations of LiP and similar fungal peroxidases in
487	relation to the amount of lignin, the peroxidase enzyme has been said to catalyse
488	repolymerisation of lignin alongside depolymerisation (Cragg et al., 2015; Rahmanpour et
489	al., 2017; Rahmanpour et al., 2016; Rashid et al., 2017). Repolymerisation of lignin means
490	that lignin would precipitate back onto the surface and recent work has shown that this has
491	a negative impact on the accessibility of the lignocellulosic structure to biodegradation (Li
492	et al., 2007; MacAskill et al., 2018; Oliva-Taravilla et al., 2016; Wiman et al., 2012). Due
493	to the very high lignin content of softwood (nearly double that of newspaper), it is possible
494	that the concentration of LiP was low for these reactors, and that depolymerisation and
495	simultaneous repolymerisation led to the lower methane yield in the enzyme-amended

496 reactors. The lignin content of the wood residue is slightly higher (38.17 ± 0.42) in the

497 enzyme reactors in relation to the controls (37.22 ± 0.02) , one possible reason for which

498 could be re-polymerisation, although this difference is not statistically significant (two-way

499 ANOVA P>0.05, F-values<F-critical). It might be that lignin droplets (post-

500 depolymerisation) coalesced onto the surface of the wood (re-polymerisation) and resulted

501 in slightly higher lignin contents in relation to the controls and lesser access for the

502 microbes to the cellulose fibres resulting in lower methane yields as shown in Figures 5 and

503 **6**.

504 **3.4** General Discussion of Bacterial and Enzyme Experiments

505 Agrobacterium sp., isolated from MSW soil, was tested(Rashid et al., 2017). for its 506 ability to break down softwood bark chips in compost under aerobic conditions, showing an 507 approximately 2.3-3.5-fold enhancement in biogas production The enhancement of a ~2-508 fold enhancement reported here is comparable to this, despite the methanogenic microbial 509 communities from organic compost used by (Rashid et al., 2017) and the anaerobic sewage 510 sludge (this study) being very different. In addition, compost certainly contains aerobic 511 strains which could result in much better initial hydrolysis of the lignocellulose making it 512 easier for the various strains tested to then attack the structure. Much of the work carried 513 out on bacterial breakdown of lignocellulosic wastes in the literature (Chandra et al., 2007; 514 Mathews et al., 2016; Mathews et al., 2015; Mathews et al., 2014; Mnich et al., 2017; 515 Tsapekos et al., 2017) has not specifically paired this process with simultaneous 516 methanogenesis as has been done in the current present study and that of (Rashid et al., 517 2017) as such this study has helped to shed more light into this gap in understanding.

518 Schroyen et al. applied peroxidase enzymes to lignocellulosic materials (Schroyen et al., 519 2017; Schroyen et al., 2014; Schroyen et al., 2015). Interestingly, the case where they were 520 able to obtain maximum enhancement of biogas from corn stover (4.5% lignin) was with a 521 laccase treatment, not peroxidase or their mixture, and it resulted in a 17% increase in 522 methane yield. In comparison in this study a $\sim 41\%$ enhancement was obtained in 523 biomethane potential (25.7% lignin in this newspaper, Table 2). In Schroyen et al.'s 524 mixture of laccase and peroxidase pre-treatments, they studied 7 substrates and obtained 525 biogas enhancement only in 4 (hemp: 4.8%, miscanthus: 9.5%, flax: 14%, willow: 40%), 526 whilst the pre-treatment had a negative impact on the biogas production of the rest of the 527 substrates (in ensilaged maize, the pre-treatment instead of enhancing biogas production, 528 nearly halved it). Their work suggested that enzymatic biodelignification is a treatment that 529 depends highly on the nature of the substrate, it may actually have a negative impact in 530 some lignocellulosic wastes, the reasons for which are not clear as of yet. Interestingly, the 531 substrates studied in their study and ours suggest that it is not entirely clear as to why the 532 peroxidase enzyme treatment works with some wastes, and not with others. Looking at the 533 lignin contents, there does not seem to be a relationship between high lignin contents and 534 proportionally negative impact on peroxidase enzyme action. However, lignin is 535 predominantly made up of guaiacyl and syringyl monomers, where the former is less cross-536 linked than the latter and in comparison, is an easier lignin-monomer to breakdown. 537 Syringyl-rich lignin has also been shown to negatively impact the growth of some lignin-538 degrading fungi (Hooker et al., 2018). It might be the case that the monomeric composition 539 of lignin plays an important role in the success of peroxidase treatment, whereby wastes

with high syringyl monomeric units (e.g. softwood) may be more difficult to attack thansay wastes made up predominantly of guaiacyl units (e.g. hardwood).

542 Jayasinghe et al. applied LiP and similar fungal peroxidases to partly degraded 30-year 543 old excavated MSW (Hettiaratchi et al., 2014; Hettiaratchi et al., 2015; Jayasinghe et al., 544 2011; Jayasinghe et al., 2014; Jayasinghe et al., 2013). They also observed a positive 545 impact of LiP on biomethane generation. However the increase in biogas yields recorded by 546 them was nearly an order of magnitude larger than the control compared to the 41% 547 increase in biomethane potential in the case of newspaper in this study. This may be due to 548 the age of their waste, since for very old landfilled waste, significant degradation of the 549 biomass likely occurred making the structure of the waste more accessible to enzymatic 550 attack. On the other hand, this study used virgin materials and since newspaper is made 551 from a mechanical pulp, much of the lignocellulosic structure is still intact (Eleazer et al., 552 1997; Wang et al., 2015; Wang et al., 1994).

553 From the experimental work carried out in this paper, it has been obvious that both the 554 lignocellulosic wastes, wood and newspaper, responded to treatment by the Agrobacterium 555 sp., whilst only newspaper responded positively to the peroxidase enzyme treatment. 556 Possible reasons for these results and biogas yields obtained have been discussed. It has 557 been highlighted that one of the major factors resulting in the different response exhibited 558 by wood and newspaper has been due to the lignocellulosic structure and particularly, 559 lignin content of the individual wastes. Finally, in comparison with enzymatic 560 biodelignification, delignifying bacteria are present in landfills (the strain used here was 561 isolated from landfill soil) and so there may be the potential to enhance their activity to 562 encourage enhanced biogas recovery.

563 4 CONCLUSIONS

564 Agrobacterium sp. enhances the biodegradation of lignin-containing wastes, specifically 565 newspaper and softwood, under idealised small-scale conditions, containing numerous 566 other microbial communities. The solid:liquid ratio is a potentially important variable for 567 the application of Agrobacterium sp. and should be considered for pilot-/field-scale trials 568 (e.g. by adjusting leachate table). Lignin peroxidase enhances the biodegradation of 569 newspaper (not wood), in conjunction with methanogenic bacteria. These results suggest 570 that enhanced breakdown of real wastes in MSW landfills and processes such as anaerobic 571 digestion is feasible using either technique but that the waste form is an important factor in 572 the rate and extent of breakdown.

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