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1 **Improving methane production from *Pennisetum* hybrid by**
2 **monitoring plant height and ensiling pretreatment**

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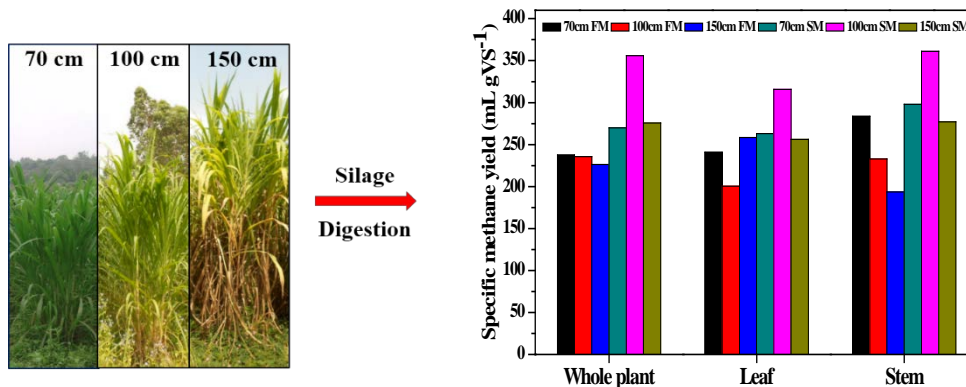
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17 **Abstract:** The biomass of grass-based *Pennisetum* hybrid commonly use for
18 abiogas production via anaerobic digestion. However, it is necessary to determine
19 a method to optimize the plant harvest time for high biogas production. Moreover,
20 ensiling of biomass in the presence of diverse microbes may offer a solution to
21 improve biogas production. In this study, whole plant of *Pennisetum* biomass
22 (including stems and leaves) was collected at different harvesting time (plant
23 heights of 70, 100, 150 cm), and then comparatively assessed for further ensiling
24 and biogas production. Compared to leaves, stems exhibited a significant linear
25 relationship ($R^2 = 0.99$) with whole plants in terms of ensiling quality (i.e. pH and
26 $\text{NH}_3\text{-N}$). Microbial analysis further revealed that *Lactobacillus* was the dominant
27 bacterial genus during ensiling of stems and whole plants, with the highest relative
28 abundance of 50.08% obtained at the height of 100 cm. Ensiling of biomass at a
29 height of 100 cm achieved the best digestion performance, with the methane yields
30 of 316 ± 20 mL/g VS for leaves, 361 ± 43 mL/g VS for stems, and 356 ± 28 mL/g
31 VS for whole plants. A harvesting time at the plant height of 100 cm was the
32 optimal from the silage quality and anaerobic digestion performance.

33 **Keywords:** *Pennisetum* hybrid biomass; plant height; ensiling; *Lactobacillus* bacteria;
34 anaerobic digestion, methane.

35 **Graphical Abstract**



36

37 1. Introduction

38 *Pennisetum* (subfamily: Panicoideae, tribe: Paniceae) is a genus of C4 grasses that
 39 are widely grown in Europe and Asia.¹ *Pennisetum* sp. is economically feasible and
 40 recommended as a promising feedstock for anaerobic digestion, due to its huge biomass
 41 yield and high organics content.²⁻⁴ The annual *Pennisetum* biomass yield was reported
 42 as 88 MT/ha, 210 t/ha of which were produced in China.⁵ The organic components of
 43 *Pennisetum* biomass are mainly composed of cellulose (40–60%) and hemicellulose
 44 (20–40%), which can be easily degraded and used in biological process.¹

45 However, the use of *Pennisetum* biomass may not be straightforward. Plant
 46 harvest time is important for anaerobic digestion performance, because the chemical
 47 composition of grass varies with its growth stage.^{6,7} For example, the specific methane
 48 yields of *Pennisetum* hybrid and switchgrass (*Panicum virgatum*) decreased from 280
 49 to 119 mL/g VS,⁸ and from 266–309 to 191–250 mL/g VS as crops matured.⁹
 50 Lehtomaki et al.¹⁰ observed that harvesting at a younger stage was optimal for Napier
 51 grass (*Pennisetum purpureum*) because it could achieve a higher specific methane yield,
 52 whereas marrow kale (*Brassica oleracea* var. *medullosa*) and Jerusalem artichoke

53 (*helianthus tuberosus*) were optimal at a later harvest, which could obtain higher
54 biomass yields. Dragoni et al. ¹¹ reported that harvesting in September might be the
55 most feasible option for *Phragmites australis*. Similarly, the optimal cutting time for
56 *Miscanthus* was between September and October. ¹² In addition, Surendra and Khanal
57 ¹³ obtained a maximum methane yield of 219 ± 4.9 mL/g VS for *P. purpureum*
58 harvested at 2 months old. Overall, the optimal harvest time varies by species, growth
59 conditions, maturity stage, and planting area. Therefore, establishing a simple method
60 to determine the harvest time is necessary to enhance methane production.

61 Furthermore, the rigid cell wall structures in biomass are strongly recalcitrant to
62 microbial degradation. Therefore, it is critical to pretreat the *Pennisetum* hybrid to
63 improve the specific methane yield. Compared to various pretreatments of biomass,
64 ensiling is a commonly used technology that can destroy the structure of cellulose and
65 hemicellulose, and preserve the nutrient component as effectively as possible. ¹⁴⁻¹⁸ High
66 quality silage can recover 87–98% of methane yield on the basis of methane potential
67 of the biomass. ¹⁹ Vervaeren et al. ²⁰ observed the process of silage could effectively
68 improve anaerobic digestion performance with an increase 10.1–14.7% of biogas
69 production.

70 However, to the best of our knowledge, few researches were reported about
71 combining the aspect of harvest time and ensiling pretreatment to enhance methane
72 production from *Pennisetum* hybrid. Therefore, the present study aimed to (1) improve
73 the silage quality and anaerobic digestion performance by comparing grass at different
74 heights; (2) evaluate the leaf and stem parts in whole plant to study the primary

75 influencing component of the silage process and conversion efficiency; and (3)
76 conclude the feasibility of determining the optimal harvest time by monitoring plant
77 height.

78 **2. Methods**

79 **2.1 Grass materials and inoculum**

80 The biomass, *Pennisetum* hybrid, was sown in Zengcheng district, Guangzhou,
81 China. The *Pennisetum* hybrid planting spacing is 60 cm × 12 cm, and the planting area
82 is 1000 m² (50 m × 20 m). Samples were collected at January 14, 2016, the
83 corresponding grasses at heights of 70 cm, 100 cm, and 150 cm were selected for the
84 study. 5-10 strains were randomly selected from the experimental base for each
85 castration, leaving 10 cm for growth. Before processing the grass, the quality of fresh
86 whole plant was weighted. For the comparison of the main factors for determining the
87 silage quality and anaerobic digestion performance, some of the raw materials were
88 separated and classified into leaves and stems, whereas other materials were classified
89 as whole-plant samples.

90 The inoculum for the anaerobic digestion was obtained from continuously stirred
91 tank reactors operated in the lab. The total solids (TS) and volatile solids (VS) contents
92 of the inoculum were determined as 3.44% and 1.43%, respectively.

93 **2.2 Experimental setup and procedure**

94 The fresh materials were cut into small pieces of 2-3 cm, pulverized, and then
95 stored at -20°C in a refrigerator until further use. The silage materials were prepared in
96 a plastic silo bag. For the ensiling process, about 200 g of fresh sample was placed in a

97 bag, vacuumed-sealed, and then ensiled at ambient temperature for 30 d. After
98 ensiling processing, the silage samples were crushed and then stored at -20°C in a
99 refrigerator for spare. Each treatment was performed in triplicate.

100 The batch anaerobic digestion experiments were carried out using an automatic
101 methane potential test system (AMPTS II, Bioprocess Control Sweden AB) at 35 ± 1
102 °C., the total and working volume of reactor was 500 mL and 400 mL, respectively. In
103 this process, 400 mL of inoculum were used, and the ensiling material was added based
104 on the VS of substrate/inoculum ratio of 1. The experiments were performed in
105 triplicate and were run for 30 d.

106 **2.3 Analytical methods**

107 The TS, VS, pH, total ammonia nitrogen concentration (NH₃-N), carbon (C), and
108 nitrogen (N) analyses were performed according to previously published methods.¹⁶ To
109 determine the microbial community composition in silages of different materials, the
110 collected samples were stored at -20°C until the analysis. Microbial characterizations
111 were based on the method of 16s rRNA high throughput sequencing. The microbial
112 DNA was extracted, amplified, and analyzed according to a previously described
113 method.²¹

114 **2.4 Kinetic analysis**

115 The modified Gompertz equation (Eq. (1)) was used for the kinetic analysis²²:

$$116 \quad M = P \times \exp \left\{ - \exp \left[\frac{R_m \times e}{P} (\lambda - t) + 1 \right] \right\} \quad (1)$$

117 where M , P , R_m , and λ represent the cumulative methane yield (mL/g VS) at a given
118 time, methane production potential (mL/g VS), maximum methane production rate

119 (mL/g VS d), and lag phase (d), respectively.

120 **3. Results and discussion**

121 **3.1 Chemical composition of the materials**

122 *Pennisetum* hybrid as the feedstock for anaerobic digestion mainly includes the
123 parts of stem and leaf, Table 1 presents the TS, VS, C, N, and C/N contents of the stem
124 and leaf in the whole plant obtained at different conditions. Fresh and silage samples
125 typically exhibited significant differences in terms of TS, VS, and N contents. Moreover,
126 samples of different plant parts derived from various plant heights (i.e. 70, 100, and 150
127 cm) also contributed to different chemical compositions. For the fresh materials, the TS
128 contents increased from $13.91 \pm 1.09\%$ to $23.11 \pm 1.65\%$ in the whole plant, from 18.13
129 $\pm 0.10\%$ to $25.73 \pm 1.08\%$ in leaf, and from $11.97 \pm 0.57\%$ to $23.07 \pm 0.03\%$ in stem as
130 the plant height increased. The increase in the TS and VS contents of *Pennisetum* hybrid
131 showed a positive correlation with plant height. These results could be due to the total
132 lignocellulose (including cellulose, hemicellulose, and lignin) content increased with
133 crop maturity.²¹ Moreover, leaf had the highest TS and VS contents, whereas stem had
134 the lowest TS and VS contents in different samples of plant height. No significant
135 difference was observed in the C contents among different biomass parts and heights;
136 however, the highest N content was obtained in leaf and the lowest N content in stem.
137 Correspondingly, the C/N values were higher in stem than those in leaf. Similar trends
138 were previously observed by Erickson et al.²⁴ and Han et al.,²⁵ who reported that the
139 N concentration in sorghum leaf was higher than that in the stem. For the silage
140 materials, the content of TS, VS, and N contents had a decrease compared to the fresh

141 samples, whereas the corresponding C/N values showed an increase. Moreover, the N
142 content in whole plant silage materials decreased from $0.98 \pm 0.02\%$ to $0.64 \pm 0.01\%$
143 with the plant height from 70 to 150 cm. The reason was that the process of ensiling
144 could degrade carbohydrates and proteins into minor molecular such as volatile fatty
145 acids (mainly including lactic acid, acetic acid, and propionic acid) and amino acids.¹⁶
146 In addition, the lowest TS and VS contents were observed in the plant height of 100 cm
147 with different plant parts. Similarly to the fresh materials, higher TS, VS, and N
148 contents were observed in the leaf silage samples, corresponding to lower C/N values.

149

150 **Table 1.**

151

152 Figure 1 presents the pH values and $\text{NH}_3\text{-N}$ concentrations in the silage samples
153 of the stem, leaf, and whole plant. In agreement with the N contents of stem, lower
154 $\text{NH}_3\text{-N}$ concentrations of were obtained for stem silage samples. Meanwhile, lower pH
155 values of 4.15–4.49 were observed in the stem silage samples. By contrast, the leaf
156 silage samples had higher pH values of 4.73–5.54, which increased with plant height
157 from 70 to 150 cm. In addition, the $\text{NH}_3\text{-N}$ concentrations in whole plant silage
158 materials decreased from 44.50 ± 0.64 mg/L to 14.00 ± 0.98 mg/L with the plant height
159 from 70 to 150 cm. Nousiainen et al.²⁶ reported that a negative association was
160 observed between the decreasing crude proteins contend and the certain stage of plant
161 maturity. And the decreasing $\text{NH}_3\text{-N}$ concentrations in the increasing heights of whole
162 plant silage materials were similar to the results of ammonia nitrogen in the dairy cow

163 fed silages harvested at four stages of grass maturity.²⁷ The pH values of the stem and
164 whole-plant silage samples were similar to the so-called critical pH values (range: 4.10–
165 4.45) for silage samples at the dry matter of 15–30%.²⁸ In order to understand the role
166 of plant part in the silage samples, the correlations of pH values and NH₃-N
167 concentration between the silages of stem, leaf, and whole-plant was analyzed in the
168 Figure 2. In a comparison of the pH values among the silage samples, a positive linear
169 relationship between stem and whole plant was observed, following the equation: $y =$
170 $7.8226 - 0.7983x$ ($R^2 = 0.9987$). However, a negative linear relationship between stem
171 and whole plant was obtained by comparing the NH₃-N concentrations of silage
172 samples, and the equation was $y = -3.5975 + 1.3736x$ ($R^2 = 0.9994$). Although the same
173 linear relationship between leaf and whole plant was observed by comparing the pH
174 value and NH₃-N concentration of silage samples, there were not significant linear
175 correlation of pH ($R^2 = 0.0805$) and NH₃-H ($R^2 = 0.3601$) between the leaf and whole
176 plant. In addition, the stem accounted for over 60% of the content of fresh whole plant.
177 Therefore, these results suggested that the part of stem had a greater effect than the leaf
178 on the silage quality of the whole plant.

179

180 **Figure. 1**

181 **Figure. 2**

182

183 **3.2 Bacterial community structure**

184 Figure 3 presents the bacterial communities in the raw material and silage samples.

185 The dominant bacterial compositions at the levels of phyla and genera were similar
186 among the fresh materials. The dominant bacteria were *Cyanobacteria/Chloroplast*,
187 with a relative abundance of 71.03–94.86%, and the major genus was *Streptophyta*,
188 with a relative abundance of 71.03–97.96%.

189 In the silage samples, a dramatic shift in the bacterial compositions at the phylum
190 level was observed in comparison with those in the fresh materials. The relative
191 abundance of *Cyanobacteria/Chloroplast* decreased to 0.72–28.27%, whereas
192 *Firmicutes* (36.26–80.72%) and *Proteobacteria* (6.05–40.79%) became the dominant
193 bacteria at the phylum level after ensiling. Remarkable differences in the relative
194 abundance at the phylum level were observed among the stem, leaf, and whole-plant
195 parts. Most sequences at the phylum level assigned to the genera *Streptophyta*,
196 *Lactobacillus*, *Lactococcus*, *Raoultella*, *Enterobacter*, *Enterococcus*, *Leuconostoc*,
197 *Serratia* and *Weissella*.

198 The most dominant at the phylum level was *Firmicutes*, and a higher relative
199 abundance of *Firmicutes* was obtained in the stem and whole plant. Desirable functional
200 bacteria in silage include *Lactobacillus*, *Enterococcus*, and *Lactococcus*, which are
201 used widely as silage additives.²⁹ These bacteria belong to a major part of the lactic
202 acid bacteria group, which could convert sugars to lactic acid.^{30,31} Since lactic acid was
203 one of the main metabolic intermediates (VFAs) in process of anaerobic digestion, it
204 could easily utilize by the acetogenic bacteria and methanogens.^{32,33} For the stem silage
205 samples, the relative abundance of *Lactobacillus* sp. ranged from 36.41% to 50.08%,
206 reaching a maximum at a height of 100 cm, while the relative abundance of *Lactococcus*

207 sp. decreased from 27.40% to 1.61% as height increased. This was coupled with an
208 increase in the relative abundance of the genus *Enterococcus*. In the leaf silage samples,
209 the relative abundance of *Lactobacillus* sp. ranged from 1.27% to 39.60%, reaching a
210 maximum at a height of 150 cm, while the variations in the relative abundance of the
211 genera of *Lactococcus* and *Enterococcus* were similar to those in the stem. In the whole
212 plant, the dominant genera differed by height. For example, *Lactobacillus* was the
213 primary genus at a height of 150 cm, while relative abundances of 37.62%
214 (*Lactobacillus* and *Lactococcus*) and 46.70% (*Lactobacillus* and *Enterococcus*) were
215 obtained for the silage samples at heights of 70 cm and 100 cm, respectively.

216 The other most abundant at the phylum level was *Proteobacteria* (6.05–40.79%),
217 the genera of *Raoultella* and *Enterobacter* predominated in this phylum. The relative
218 abundance of *Raoultella* in silage samples increased from 1.08% to 9.36% in stem
219 and from 0.71% to 30.73% in leaf, while the relative abundance in the whole plant
220 ranged from 2.42% to 24.52%. *Enterobacter* had a relative abundance of 0.55–
221 30.57%. *Enterobacter* and *Raoultella* have been shown to be deleterious
222 microorganisms during the ensiling process.^{34, 35} Because these bacteria could largely
223 consume sugars and other simple compounds in ensiling process^{34, 35} it is not
224 beneficial to produce more methane for anaerobic digestion. The lowest relative
225 abundance of *Enterobacter* and *Raoultella* in whole plant samples was obtained at a
226 height of 100 cm. Overall, the plant height of *Pennisetum* hybrid harvested at 100 cm
227 for ensiling not only had the highest relative abundance of desirable functional
228 bacteria (*Lactococcus*, *Lactobacillus* and *Enterococcus*), but also had the lowest

229 relative abundance of deleterious bacteria (*Enterobacter* and *Raoultella*) for ensiling.
230 Therefore, these results suggested that grass harvested at a plant height of 100 cm
231 could improve the quality of silage.

232

233 **Figure. 3**

234

235 **3.3 Anaerobic digestion performance**

236 Figure 4 and Table 2 present the cumulative and specific methane yields of fresh
237 and silage materials. For the fresh materials, the specific methane yields decreased from
238 238 ± 12 mL/g VS to 226 ± 8 mL/g VS for the whole plant and from 263 ± 5 mL/g VS
239 to 194 ± 10 mL/g VS for stem as height increased. Meanwhile, the 80% cumulative
240 methane yield was obtained at 9 d for the stem and whole plant at heights of 100 cm
241 and 70 cm, respectively, but required 10–14 d for samples at a height of 150 cm. The
242 specific methane yields of leaf ranged from 206 ± 5 mL/g VS to 258 ± 6 mL/g VS.
243 Ensiling decreased the time required to obtain an 80% cumulative methane yield to 7–
244 12 d for different parts of grass. Moreover, an increased specific methane yield was
245 observed in the silage samples, and their specific methane yields were in the range of
246 263.17-298.04 mL/g VS, 315.75-361.25 mL/g VS, and 256.23-277.11 mL/g VS for the
247 plant height of 70 cm, 100 cm, and 150 cm, respectively. The maximum specific
248 methane yield of 316 ± 20 mL/g VS for leaf, 361 ± 43 mL/g VS for stem, 356 ± 28
249 mL/g VS for whole plant was obtained at a plant height of 100 cm. Since the
250 lignocellulosic structure of *Pennisetum* hybrid was disrupted by the desirable functional

251 bacteria in the process of ensiling, it could be efficiently converted into biogas by the
252 microorganisms of anaerobic digestion.^{32, 33} In addition, the samples harvested at plant
253 height of 100 cm had a better silage quality by the bacterial community analysis. Similar
254 specific methane yield results have been reported elsewhere. For example, the methane
255 yields for tall fescue, cocksfoot, and reed canary grass were between 238 mL/g VS and
256 446 mL/g VS depending on N fertilization and harvest frequency.³⁶ Moreover, specific
257 methane yields of 135 mL/g VS and 185 mL/g VS were reported for switchgrass and
258 giant cane, respectively.^{37, 38} The better performance of biogas production was observed
259 in the silage samples of the plant height 100 cm for preferred bacteria community. These
260 results suggested that harvesting plants at a height of 100 cm might be suitable for
261 biogas production from the perspectives of silage quality and anaerobic digestion
262 performance.

263 The regression analysis showed satisfactory overall agreements between the
264 experimental data and the model, with high regression coefficients ($R^2 > 0.94$) (Table 2
265 and Figure 4). More methane production potential and higher maximum methane
266 production rate were observed in the silage samples. Similar result was observed in
267 anaerobic digestion of the silage *Pennisetum purpureum* with molasses-processed
268 wastewater addition.²¹ The stem, leaf, and whole plant from plants harvested at a height
269 of 100 cm were associated with a higher methane production potential and maximum
270 methane production rate compared with the silage samples harvested at heights of 70
271 cm and 150 cm. It indicated that the silage samples harvested at a height had a better
272 anaerobic digestion performance than the other ensiling samples. These predicated

273 results of the model were consistent with the specific anaerobic digestion performance
274 of the silage samples harvested at the height of 100 cm. A negligible lag time (λ) was
275 obtained for the fresh and silage samples. Allen et al. ³⁹ reported the biochemical
276 methane potential of hay grass varied from 156 mL/g VS to 433 mL/g VS for first cut
277 baled silage, and the kinetics analysis showed the similar results of the methane
278 production potential and lag time. According to the results of the specific methane
279 yields and the bacterial community analysis in the ensiling samples, the optimal
280 harvesting time at the plant height of 100 cm and the pretreatment of silage showed a
281 positive effect on the anaerobic digestion performance of the energy grass.

282

283 **Table 2.**

284

285 **Figure. 4**

286

287 **4. Conclusions**

288 The height of *Pennisetum* hybrid at which it was harvested was demonstrated to
289 have significant effects on silage quality and the subsequent anaerobic digestion. The
290 results from silage quality of different materials concluded a linear relationship between
291 the stem and whole plant. Microbial community analysis revealed that *Lactobacillus*
292 was the dominant genus in stem silage, and reached the maximum at harvesting height
293 of 100 cm. This suggested that the stem had a primary influence on the silage quality.
294 The maximum specific methane yield was 356 ± 28 mL/g VS for the whole plant at a

295 height of 100 cm, indicating that a harvesting height of 100 cm could be the optimal
296 from the perspective of silage quality and biogas production.

297

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429 **Table captions:**

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431 **Table 2.** Anaerobic digestion performance and kinetic parameters of the samples of
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434 **Figure. 1.** The parameters determining the silage quality of different samples: (a) pH
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437 silages of stem, leaf, and whole-plant.

438 **Figure. 3.** Bacterial compositions at the (a) phylum and (b) genus level of the samples
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442 **Figure. 4.** Comparison of the cumulative biogas yields from the samples of
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Table 1. Characteristics of the fresh and silage materials of *Pennisetum* hybrid.

			TS (%)	VS (%)	C (%)	N (%)	C/N
Fresh material	70cm	Whole	13.91±1.07	11.79±0.86	39.14±0.01	1.08±0.01	36.24±0.46
		Leaf	18.13±0.09	15.44±0.30	40.44±0.14	1.43±0.01	28.28±0.18
		Stem	11.97±0.57	10.51±.64	39.72±0.11	0.48±0.11	83.62±1.02
	100cm	Whole	14.58±0.53	12.56±0.28	39.89±0.11	0.88±0.01	45.59±0.50
		Leaf	18.37±0.46	16.11±0.50	40.61±0.08	1.50±0.01	27.16±0.07
		Stem	11.92±0.49	10.56±0.61	39.83±0.10	0.51±0.00	78.10±0.19
	150cm	Whole	23.11±1.65	20.29±2.34	40.98±0.09	1.03±0.01	39.79±0.46
		Leaf	25.73±1.08	22.04±1.33	41.03±0.11	1.35±0.01	30.55±0.08
		Stem	23.07±0.03	21.22±0.03	41.84±0.05	0.43±0.01	98.45±1.75
Silage material	70cm	Whole	15.09±0.52	12.34±0.29	40.72±0.06	0.98±0.02	41.77±0.97
		Leaf	18.82±0.25	15.77±0.48	40.83±0.22	1.34±0.01	30.58±0.00
		Stem	11.79±0.27	10.00±0.32	40.74±0.06	0.54±0.01	75.46±2.09
	100cm	Whole	13.72±0.35	11.55±0.34	39.59±0.04	0.92±0.02	43.28±0.96
		Leaf	16.82±0.78	13.52±0.64	40.48±0.04	1.26±0.01	32.13±0.39
		Stem	10.58±0.71	9.00±0.60	41.12±0.16	0.54±0.00	76.19±0.29
	150cm	Whole	22.05±0.86	18.81±0.86	41.28±0.05	0.64±0.01	64.51±1.50
		Leaf	28.52±0.56	23.83±0.81	41.00±0.16	1.41±0.05	29.20±0.92
		Stem	18.32±3.31	16.04±3.47	41.79±0.05	0.36±0.01	117.73±2.21

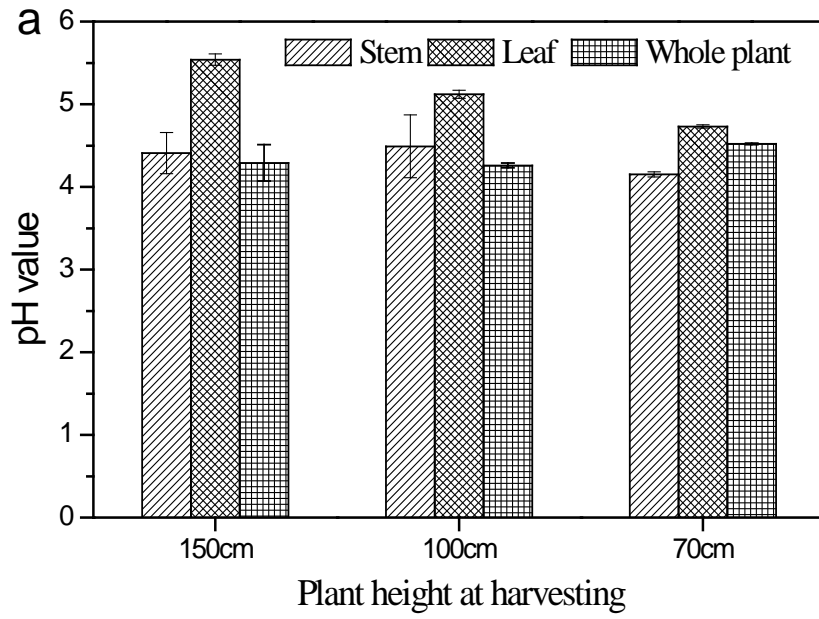
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454 *Pennisetum* hybrid.

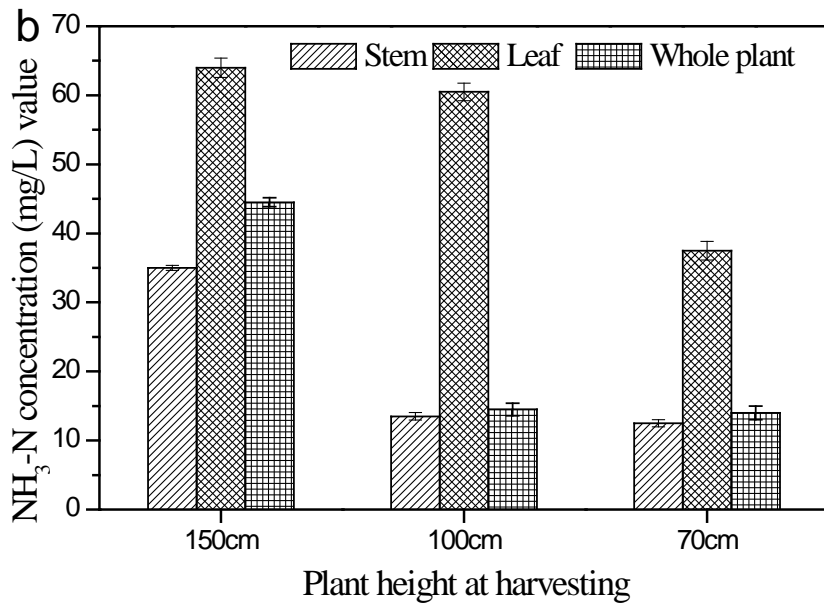
Samples			Anaerobic digestion performance (mL/g VS)	Kinetic parameter			
				P (mL/g VS)	R _m (mL/g VS d)	Λ (d)	R ²
Fresh material	70 cm	Whole	237.62	232.27	29.35	0.33	0.996
		Leaf	240.90	235.87	29.87	0	0.995
		Stem	283.60	273.97	33.15	0	0.988
	100 cm	Whole	235.67	226.60	29.66	0	0.990
		Leaf	200.40	197.12	20.69	0.10	0.998
		Stem	232.85	224.00	30.38	0	0.988
	150 cm	Whole	226.37	219.68	23.15	0	0.984
		Leaf	258.34	249.32	31.28	0	0.987
		Stem	193.70	194.04	13.31	0	0.988
Silage material	70 cm	Whole	270.04	267.75	47.23	0.57	0.999
		Leaf	263.17	259.43	35.26	0.45	0.998
		Stem	298.04	293.52	46.45	0.34	0.997
	100 cm	Whole	355.77	350.56	43.41	0.39	0.997
		Leaf	315.75	312.90	40.74	0.82	0.999
		Stem	361.25	353.73	46.25	0.11	0.993
	150 cm	Whole	275.73	271.60	21.95	0	0.983
		Leaf	256.23	248.56	29.63	0	0.982
		Stem	277.11	271.72	23.13	0	0.981

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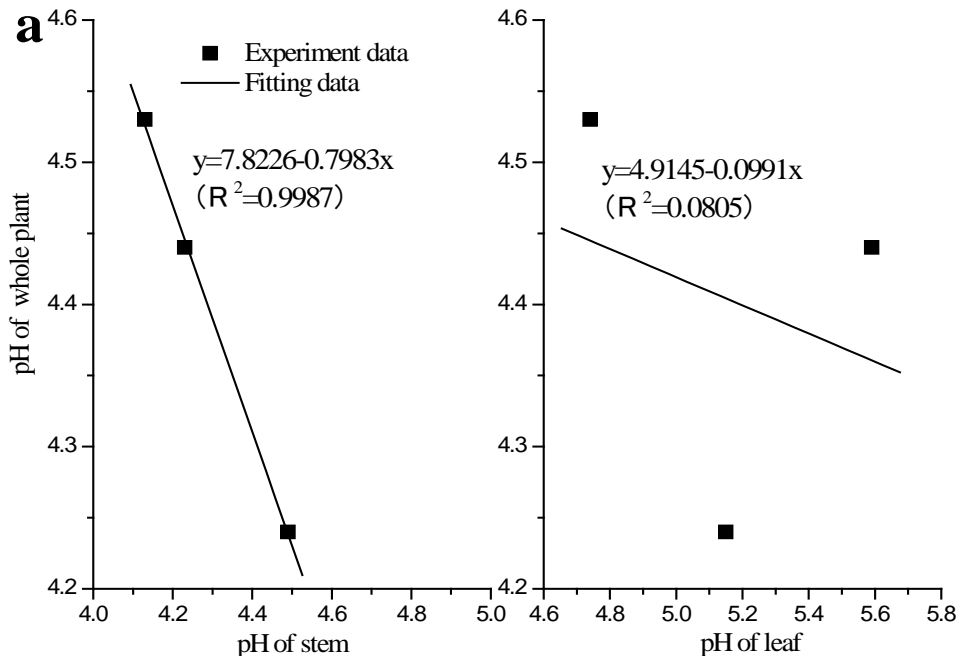
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459 **Figure. 1.** The parameters determining the silage quality of different samples: (a) pH

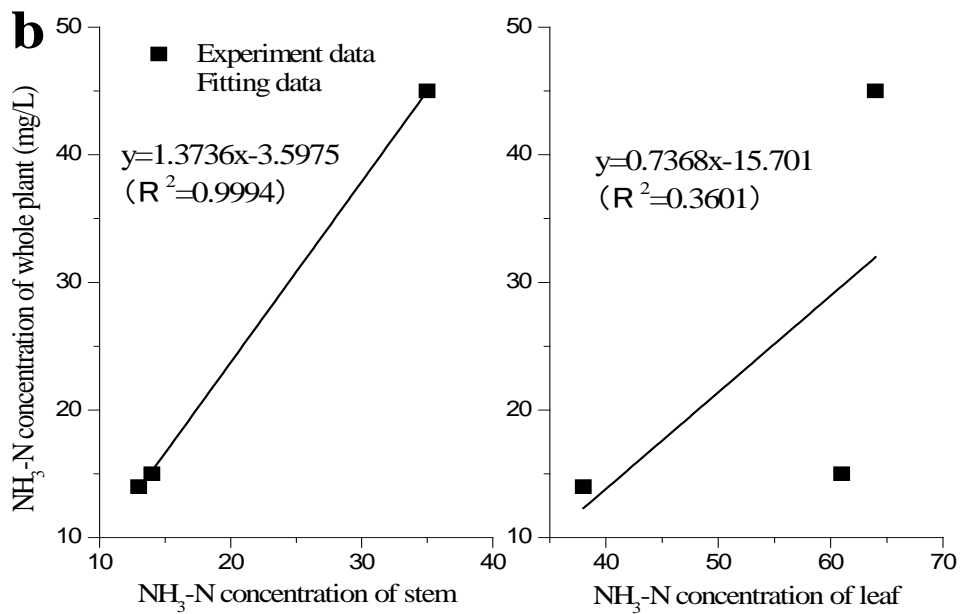
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values, (b) $\text{NH}_3\text{-N}$ concentrations.

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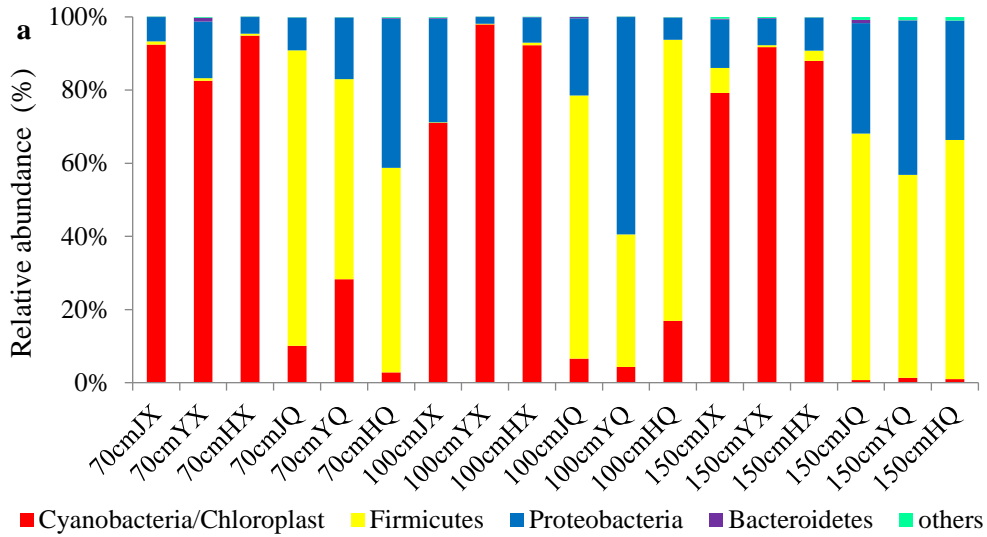
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464 **Figure. 2.** The correlations of (a) pH values and (b) NH₃-N concentration between the

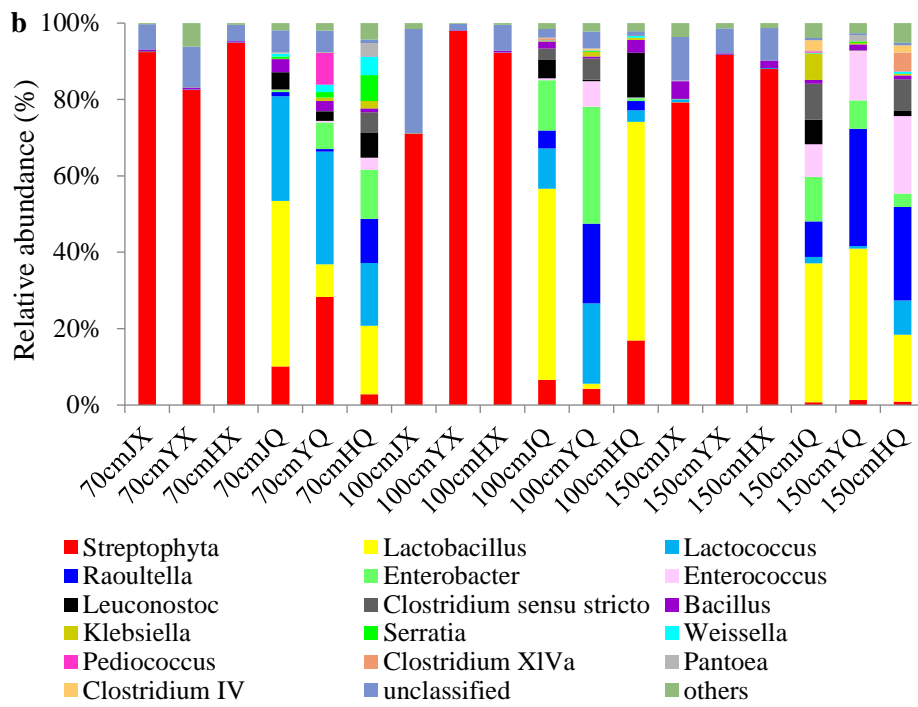
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silages of stem, leaf, and whole-plant.

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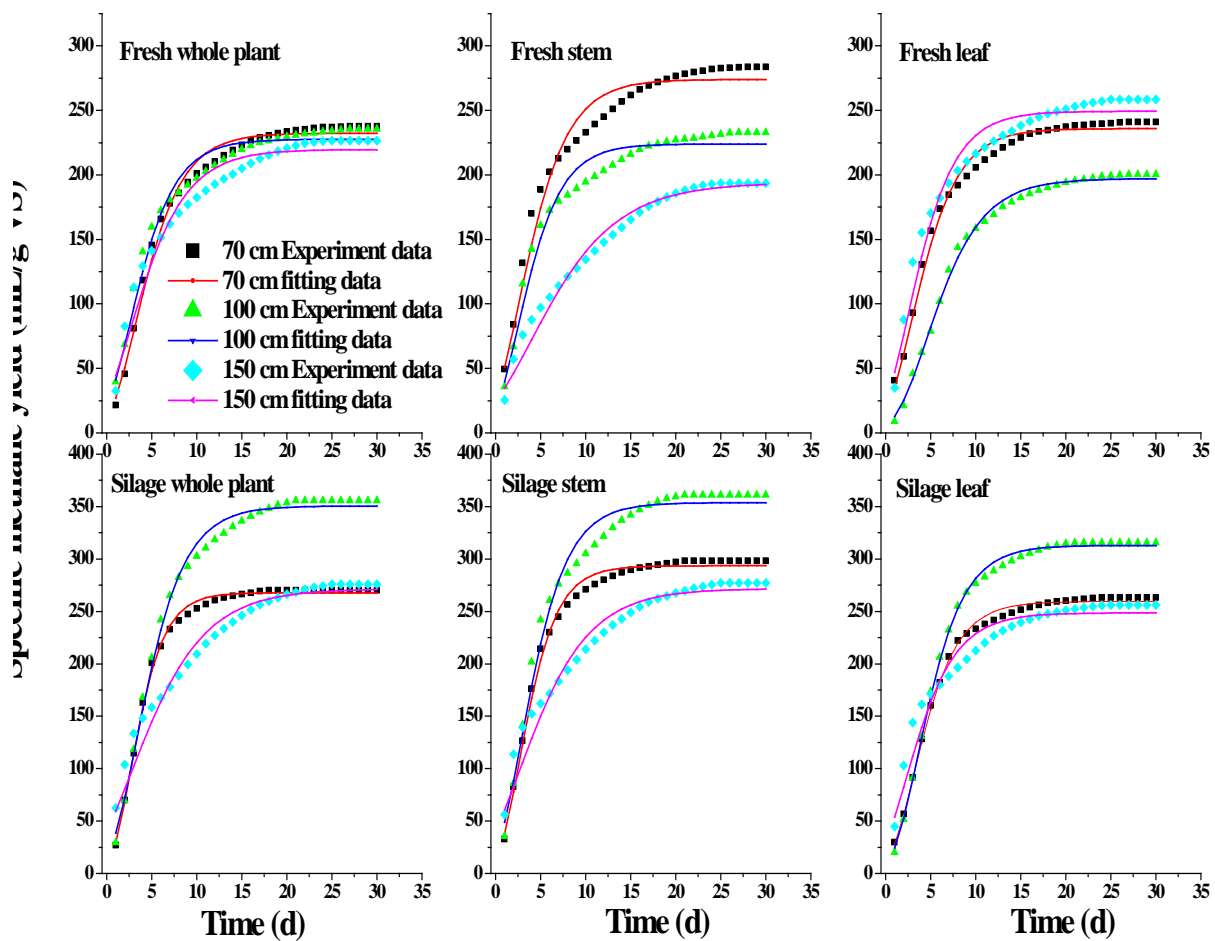


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Figure. 4. Comparison of the cumulative biogas yields from the samples of

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Pennisetum hybrid