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Abstract: Thermal gasification of various biomass residues is a promising technology for combining bioenergy production with soil fertility management through the application of the resulting biochar as soil amendment. In this study, we investigated gasification biochar (GB) materials originating from two major global biomass fuels: straw gasification biochar (SGB) and wood gasification biochar (WGB), produced by a Low Temperature Circulating Fluidized Bed gasifier (LT-CFB) and a TwoStage gasifier, respectively, optimized to energy conversion. Stability of carbon in GB against microbial degradation was assessed in a short-term soil incubation study and compared to the traditional practice of direct incorporation of cereal straw. The GBs were chemically and physically characterized to evaluate their potential to improve soil quality parameters. After 110 days of incubation, about 3 % of the added GB carbon was respired as CO₂, compared to 80 % of the straw carbon added. The stability of GB was also confirmed by low H/C and O/C ratios with lowest values for WGB (H/C 0.01 and O/C 0.14). The soil application of GBs exhibited a liming effect increasing the soil pH from ca 8 to 9. Results from scanning electron microscopy and BET analyses showed high porosity and specific surface area of both GBs, indicating a high potential to increase important soil quality parameters such as soil structure, nutrient and water retention especially for WGB. These results seem promising regarding the possibility to combine an efficient bioenergy production with various soil aspects such as carbon sequestration and soil quality improvements.

Response to Reviewers: Response to Reviewers' comments:

The units and ratios were corrected (L 143, L184).

Treatments description were also improved (L159-64)

The references have been corrected.

Thank you for your comments.

Best regards,

Veronika Hansen

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This manuscript demonstrates the potential for combining of bioenergy production and residual biochar application as soil improving and carbon sequestration agent. This study shows that carbon in gasification biochar is stable against microbial degradation. Furthermore, the liming effect, high porosity and specific surface area of the gasification biochar indicate the ability of the biochar to improve important soil quality parameters such as structure, water and nutrient retention.

Therefore, gasification of crop residues and wood waste is a promising way of producing sustainable bioenergy and reaching the political goals of fossil fuel free society, and at the same time sustaining or even improving of the soil quality, which is crucial for meeting the increasing demand for producing food and energy crops.

We hope that our experimental study might be interesting and relevant for publication in Biomass and Bioenergy. We are looking forward to receive the reviewer's comments and evaluation.

Best regards,

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*Detailed Response to Reviewers

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Highlights

- Biomass gasification has a potential to combine the efficient production of bioenergy with valuable biochar residuals for soil quality improvements.
- The two investigated gasification biochars are recalcitrant indicating soil carbon sequestration potential.
- Gasification biochars have a potential as soil improvers due to high specific surface area, porosity, liming effect and low PAH content.

1 **Gasification biochar as a valuable by-product for carbon sequestration and soil amendment**

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10

11 **Abstract**

12 Thermal gasification of various biomass residues is a promising technology for combining
13 bioenergy production with soil fertility management through the application of the resulting biochar
14 as soil amendment. In this study, we investigated gasification biochar (GB) materials originating
15 from two major global biomass fuels: straw gasification biochar (SGB) and wood gasification
16 biochar (WGB), produced by a Low Temperature Circulating Fluidized Bed gasifier (LT-CFB) and
17 a TwoStage gasifier, respectively, optimized to energy conversion. Stability of carbon in GB against
18 microbial degradation was assessed in a short-term soil incubation study and compared to the
19 traditional practice of direct incorporation of cereal straw. The GBs were chemically and physically
20 characterized to evaluate their potential to improve soil quality parameters. After 110 days of
21 incubation, about 3 % of the added GB carbon was respired as CO₂, compared to 80 % of the straw
22 carbon added. The stability of GB was also confirmed by low H/C and O/C atomic ratios with
23 lowest values for WGB (H/C 0.01 and O/C 0.14). The soil application of GBs exhibited a liming
24 effect increasing the soil pH from ca 8 to 9. Results from scanning electron microscopy and BET
25 analyses showed high porosity and specific surface area of both GBs, indicating a high potential to
26 increase important soil quality parameters such as soil structure, nutrient and water retention
27 especially for WGB. These results seem promising regarding the possibility to combine an efficient
28 bioenergy production with various soil aspects such as carbon sequestration and soil quality
29 improvements.

30

31 **Keywords**

32 Gasification, Bioenergy efficiency, Biochar soil amendment, Carbon sequestration, Soil quality
33 improvement

34 **1. Introduction**

35 Biomass gasification for combined heat and power (CHP) production has the potential to become an
36 efficient and flexible way to generate bioenergy, as a broad variety of biomass residues and other
37 organic resources can be utilized [1, 2]. In Denmark effective gasification platforms for the two
38 major global biomass fuels, wood chips and cereal straw, are currently scaled up and close to
39 commercial application: (1) Low Temperature Circulating Fluidized Bed gasifier (LT-CFB),
40 specifically designed to produce energy from biomasses with high ash contents (such as straw) and
41 (2) TwoStage gasifier, designed for converting woody biomass. The LT-CFB technology has been
42 demonstrated in continuous operation, as a 6 MW demonstration plant, and the first 2 MW
43 commercial plant for continues CHP production with the TwoStage process is about to produce
44 power and district heating for a local community, Hilleroed Municipality, Denmark. This plant will
45 produce approximately 64 tons of biochar residues annually, while the planned 60 MW full scale
46 commercial LT-CFB plant is going to generate approximately 10 000 tons of carbon-rich residues
47 per year. The potential further upscaling and expanding of those processes requires a strategy for
48 the sustainable utilization of a growing amount of biochar residues produced. Recirculation and
49 utilization of those residues to agricultural land, instead of costly disposing as a waste, would
50 improve the sustainability and economy of the bioenergy production. Gasification biochar generally
51 contains a considerable amount of minerals and recalcitrant carbon and is considered an attractive
52 product for soil amendment due to its fertilizer and carbon sequestration potential [3, 4].

53 Carbon sequestration in soil mitigates the effect of climate change [5], and may furthermore help to
54 maintain or even improve the soil fertility. This is of key importance to be able to fulfil the
55 increasing global demand for producing crops for both food and energy [6]. Soil organic carbon
56 (SOC) influences the physical, chemical and biological properties of the soil, and is essential for
57 good soil quality [7]. Increasing SOC has been shown to improve soil aggregation, water

58 infiltration, and water and nutrient retention [8, 9]. Traditional annual incorporation of crop residues
59 such as cereal straw can increase soil organic matter content [10], therefore there is a concern that
60 the removal of residues from the field for energy production may lead to soil degradation [11].
61 Gasification of biomass and returning the residual biochar-carbon to the field is regarded as a
62 promising strategy combining effective bioenergy generation with the maintenance of soil carbon
63 stocks [2]. Utilizing low quality wood and residues from timber harvesting for bioenergy production
64 and subsequent addition of wood biochar to agricultural soils may be another strategy to increase
65 SOC and improve arable soils' productivity, creating novel synergies between the agricultural and
66 forestry sectors. Nevertheless, since there are qualitative differences in the molecular structure of
67 pyrogenic carbon compared to the stable carbon derived from microbial/enzymatic soil processes
68 [12], the impacts of substituting crop residue incorporation with the addition of gasification biochar
69 (GB) on soil services are largely unknown and should be thoroughly investigated before
70 implementing this into practice [8].

71 Several studies have shown positive impacts of pyrolysis biochar, produced at relatively low
72 temperatures (400 – 600°C), on soil properties [13, 14], which are, however, highly dependent on
73 biochar feedstock and thermal processing conditions [15]. The physical properties of biochars, such
74 as high porosity and specific surface area (BET), may result in an increase of not only soil water
75 retention [16], water infiltration, and cation exchange capacity [5, 13], but also soil microbial
76 activity [14]. Chemical properties, such as low hydrogen-to-carbon (H/C) and oxygen-to-carbon
77 (O/C) ratios, result in high stability of biochar against microbial degradation in soil [17]. Compared
78 to pyrolysis biochar, GB is produced at higher temperatures (around 700 – 1100°C), using low
79 amounts of oxygen. Gasification results in higher energy yields compared to pyrolysis and leaves
80 biochar with less, but more stable carbon, compared to pyrolysis biochar [15, 18]. Chemical
81 characterization of GB, showing its stable structure, is well reported [4, 15, 17, 19], however studies

82 on the effect of GB on soil and microbial processes are scarce. Concerns about the use of GB as a
83 soil amendment include its possible content of Polycyclic Aromatic Hydrocarbons (PAH) [20],
84 which proved to be highly variable, as e.g. in the studies of Wiedner [4] and Kloss [20], who
85 measured values up to 15 and 33 mg kg⁻¹, respectively. Especially the wood gasification biochars
86 showed high PAH contents [4, 17].

87 The aim of this study was to evaluate the potential of the biochar residues from two gasification
88 processes to exert a beneficial effect on soil carbon sequestration and soil quality. Through a short-
89 term soil incubation study and physical and chemical analyses, the objectives were to investigate if
90 the gasification biochars: (1) contain carbon recalcitrant to microbial degradation; (2) have a
91 potential to improve soil physical and chemical properties; (3) have any negative effects on
92 microbial biomass and (4) have a potential for higher carbon sequestration rates than those achieved
93 with traditional direct soil incorporation of the feedstock (i.e. straw).

94 **2. Materials and methods**

95 **2.1. Biochar production**

96 The two gasification biochars (GB) used for this study originated from continuously operated pre-
97 commercial gasification demonstration plants. Straw gasification biochar (SGB) was produced in a
98 Low Temperature Circulating Fluidized Bed gasifier (LT-CFB). The straw originated from winter
99 wheat (*Triticum aestivum* L.) grown in Zealand, Denmark, but is of unknown provenance, date of
100 harvest and chain of custody. Commercially produced wheat straw pellets were crushed prior to LT-
101 CFB gasification for optimal gasifier operation. Wood gasification biochar (WGB) was produced
102 from pine wood (*Pinus spp.*) chips in a TwoStage gasifier. The wood chips were commercially
103 produced with an average chip size of 50 mm, which is the optimal size for the TwoStage process,
104 and originated from Zealand, Denmark.

105 The LT-CFB gasifier (Fig. 1), developed at the Technical University of Denmark in cooperation
106 with Danish Fluid Bed Technology, is designed to gasify biomass resources with high contents of
107 low melting ash compounds (e.g. straw, manure or sewage sludge), that have proven difficult to
108 convert in other processes [1]. The process is based on separate pyrolysis and gasification fluid bed
109 reactors with a suitable circulating heating medium to transfer the heat from the gasification process
110 to the pyrolysis. The temperature is kept below the melting point of the ash components, i.e. max
111 process temperatures around 700 - 750°C. In this way, sintering of the ash and subsequent fouling
112 (from e.g. potassium) or corrosion (from e.g. chlorine) of the plant unit operations are avoided, as
113 these compounds will leave the process in solid form as ash particles.

114 Fig. 1 here.

115

116 The char conversion in the LT-CFB gasifier is a combination of sub stoichiometric oxidation of the
117 char and steam gasification. The retention time (few seconds) in the char reactor is relatively short.
118 The char-ash particles are though circulated in the process until they are too small/light to be
119 separated by the primary cyclone, subsequently most of the ash and unconverted biochar is
120 separated out of the hot gas by the secondary cyclone. The LT-CFB technology is now owned by
121 the company Dong Energy and is being commercialized under the name Pyroneer [21].

122 The TwoStage fixed bed process (Fig. 2) was invented and developed at the Technical University of
123 Denmark and has been designed for gasification of woody biomass with low ash content [1]. The
124 TwoStage process is characterized by having pyrolysis and gasification in separate reactors with an
125 intermediate high temperature tar-cracking zone with temperatures of 1000 - 1200°C. This allows a
126 very fine control of the process temperatures, resulting in extremely low tar concentrations in the
127 produced gas, making it suitable for gas engine operation or synthesis of biofuels. Due to the high
128 temperatures, the process is only applicable for woody biomass. The char conversion is
129 predominantly a gasification reaction between carbon and steam. The char is exposed to steam at
130 high temperature, 800 - 1000°C, for a relatively long period (30+ minutes), resulting in an activated
131 char with a high surface area.

132 Fig. 2 here.

133

134 ***2.2. Biochar characterization***

135 The total content of organic C, H and O in feedstock and gasification biochar was measured on an
136 elemental analyzer (FLASH 2000 Organic Elemental Analyzer, Thermo Scientific, Cambridge
137 UK). The WGB and wood chips were ball milled, while the straw was ground prior to this analysis.
138 The specific surface area was determined by the Brunauer-Emmett-Teller (BET) method by

139 nitrogen gas sorption at 77 K (Quantachrome instruments, Boynton Beach, USA). Pore size
140 distribution was obtained by Barret-Joyner-Halenda (BJH) desorption analysis after degassing the
141 samples for 2 hours at 160°C. The WGB was hand sieved in two fractions (0-0.5 and 0.5-1 mm)
142 prior to this analysis. Carbon-coated biochar samples were examined by scanning electron
143 microscope (SEM) JEOL JSM-5900 (Oxford instruments, Japan). The pH of biochar was measured
144 in a 1:5 (w/v) biochar/Milli-Q water suspension. The ash fraction was determined by heating dried
145 biochar at 550°C for 5 hours in a muffle furnace. Nine Polycyclic Aromatic Hydrocarbons (PAHs)
146 were quantified after a soxhlet extraction of 2 g sample with toluene for 48 hours by Eurofins GfA
147 (Hamburg, Germany). The measured PAHs comprised Acenaphthene, Fluorene, Phenanthrene,
148 Fluoranthene, Pyrene, Benzo(bjk)fluoranthene, Benzo(a)pyrene, Indeno(1,2,3-cd)pyrene and
149 Benzo(ghi)perylene. The particle size distribution of the biochars was determined by a vibrating
150 screen method using sieves (Retsch, Germany).

151

152 **2.3. Incubation study**

153 *2.3.1. Soil*

154 A sandy loam soil from a conventional agricultural field at Bregentved estate, Zealand, Denmark
155 (55° 22' N, 12° 05' E) was collected from the plough layer (0-25 cm), air dried and sieved to
156 obtain a fraction ≤ 6 mm. The soil contained 14 % clay, 14% silt, 47 % fine sand and 24 % coarse
157 sand. The total C content was 1.98 % and total N 0.18 %.

158 *2.3.2. Experimental design*

159 We conducted an incubation experiment including 7 treatments with 4 replicates each. In 280 ml
160 PVC containers, 200 g soil (dry weight) were mixed thoroughly with either 2 g (1 %) or 10 g (5 %)
161 straw or wood GB (dry weight). The treatments were: (1) Control soil without addition of organic

162 material (Control), (2) soil amended with 1 % straw (Straw1), (3) soil amended with 5 % straw
163 (Straw5), (4) soil amended with 1 % straw gasification biochar (SGB1) , (5) soil amended with 5 %
164 straw gasification biochar (SGB5), (6) soil amended with 1 % wood gasification biochar (WGB1),
165 (7) soil amended with 5 % wood gasification biochar (WGB5). The straw used for this experiment
166 was from winter wheat (*Triticum aestivum* L.) produced in Zealand, Denmark. After harvest, it was
167 baled and kept dry. The straw material was ground to a particle size of ≤ 5 mm prior the
168 incubation. The water content of the soil mixtures was adjusted to 50 % of the water holding
169 capacity (determined separately for each respective mixture), and kept constant by regular weighing
170 and watering. The containers were sealed with plastic lids with five holes (5 mm) to allow gas
171 exchange while minimizing moisture loss, and incubated in the dark at 22°C for 110 days. The
172 whole experiment was set up in 5 sets, enabling 5 destructive samplings. Soil respiration was
173 measured on the same set each time, which was then used for the last destructive sampling.

174 2.3.3. Soil analysis

175 Destructive soil samplings were taken at day 1, 8, 16, 32 and 110. All treatments were analyzed for
176 nitrate (NO_3^-), ammonium (NH_4^+) and dissolved organic carbon (DOC) content by extracting 10 g
177 of fresh soil with 50 mL 0.5 mol K_2SO_4 L^{-1} . The suspensions were shaken on a horizontal shaker
178 for 1 h (2.5 Hz), filtrated through pleated filter paper with retention of 5-8 μm (Grade 202F,
179 Frisenette Aps, Denmark) and stored at -20°C until analysis. The extracts were analyzed for
180 concentrations of NO_3^- and NH_4^+ on an AutoAnalyzer 3 (AA3 Bran and Luebbe, Norderstedt,
181 Germany), and for DOC on a TOC-VCPH (Shimadzu Corp., Kyoto, Japan). The soil microbial
182 biomass carbon (SMB-C) content in each treatment was determined by vacuum incubation of 10 g
183 soil mixture with chloroform for 24 hours, followed by K_2SO_4 extraction. The SMB-C was
184 estimated from the relationship $\text{SMB-C} = (\text{DOC}_{\text{fumigated}} - \text{DOC}_{\text{unfumigated}})/0.45$ [22]. The soil pH was
185 determined using soil-water suspension of 5 g soil and 25 ml of Milli-Q water.

186 **2.4. Soil respiration**

187 The CO₂ emission from each sample was measured with an infra-red gas analyzer (LI-COR 8100,
188 Lincoln, Nebraska USA). The measuring frequency ranged from daily in the beginning of the
189 experiment to once a month at the end. The emissions were measured at day 1, 2, 3, 4, 5, 8, 10, 12,
190 15, 18, 22, 30, 36, 46, 52, 67 and 110 of the incubation period.

191 **2.5. Statistical analysis**

192 Statistical analysis of the data was performed in R, version 3.0.2. The significant interaction effect
193 between treatment and time (day) was assessed using a two-way analysis of variance (ANOVA).
194 The differences between treatments within each day of measurement were analyzed using the
195 Student-Newman-Keuls (SNK) test from the R-package “agricolae” at $P \leq 0.05$.

196 **3. Results**

197 **3.1. Biochar characterization**

198 Table 1 illustrates that 4 and 10 % of the carbon in wood and straw feedstock, respectively, were
199 retained in the biochar fraction. The chemical characterization of soil, feedstock and biochars is
200 given in Table 2. Gasification of straw and wood chips led to mass loss of H and O, decrease of H/C
201 and O/C atomic ratios and increase of ash percentage. The carbon content was higher, while H/C
202 and O/C ratios were lower for WGB compared to SGB. The total content of 9 PAHs was 5 mg kg⁻¹
203 in SGB and 0.69 mg kg⁻¹ in WGB.

204 The particle size distribution of biochars is shown in Table 3. Generally, the SGB was a fine
205 powder consisting of small particles, whereas WGB was a mixture of both very small and large
206 particles (up to 1 cm).The majority of WGB-particles were larger than 0.045 mm, while the
207 opposite was true for SGB. Table 4 presents results from BET analysis. Specific surface area (SSA)
208 and pore volume were higher for WGB compared to SGB. The particle size of WGB was crucial, as
209 SSA and pore diameter were more than twice as high in particles larger than 0.5 mm compared to
210 particles smaller than 0.5 mm. SEM images illustrated in Fig. 3 show the porous structure of both
211 biochars and the higher proportion of internal pores in WGB compared to SGB.

212 Table 1 here.

213 Table 2 here.

214 Table 3 here.

215 Table 4 here.

216 Fig. 3 here.

217 **3.2. Incubation study**

218 *3.2.1. Soil sampling*

219 The addition of straw resulted in a decrease of soil mineral nitrogen (N_{\min}) content ($\text{NO}_3^- + \text{NH}_4^+$) to
220 almost zero already at the second sampling day and stayed at that level during the rest of the
221 incubation period (Fig. 4). In contrast, the N_{\min} level increased over time in the control treatment
222 and after the addition of GB. The application of the high dosage of GB resulted in about the same
223 N_{\min} content as in the control treatment, while the low dosage of GB decreased N_{\min} significantly.

224 Both straw and SGB amendments caused a significantly increased content of dissolved organic
225 carbon (DOC) in soil compared to the control treatment throughout the incubation period, except
226 the Straw1 treatment at the last sampling day (Fig. 5A). At day 1, an especially high DOC level
227 could be observed in the treatment with 5 % straw. On the contrary, the soil amendment with WGB
228 led to a significantly lower DOC content compared to all other treatments throughout the incubation
229 period.

230 The content of soil microbial biomass carbon (SMB-C) was - in accordance with DOC -
231 significantly increased after addition of straw compared to the rest of the treatments, especially in
232 the beginning of the incubation (Fig. 5B). Subsequently, the SMB-C decreased until day 16 and
233 increased again towards the end of the incubation. After 8 days of incubation, the content of SMB-C
234 in WGB-treated soil was significantly lower than in the control treatment, and this difference
235 became larger with time. On the contrary, there was no consistent effect of adding SGB on SMB-C:
236 only at day 8 and 110 in the high-dosage treatment the SMB-C was lower compared to the control.

237 Addition of both gasification biochars increased the pH of the soil significantly, and the difference
238 remained throughout the incubation period (Fig. 6). After 110 days, the pH increased by 1.13 and
239 1.36 units for SGB5 and WGB5, respectively. By contrast, soil amendment with straw significantly

240 decreased the pH in the beginning of the incubation, whereas there was no difference anymore after
241 110 days.

242 Fig. 4 here

243

244 Fig. 5 here.

245 Fig. 6 here.

246

247 3.2.2. Soil respiration

248 The addition of straw to soil, at both 1 and 5%, resulted in significantly higher CO₂ emissions
249 compared to control and GB treatments throughout the experimental period (Fig. 7A). The peak
250 CO₂ emissions in the straw and control treatments were observed during the first week of
251 measurement. Soil amendment with GB did not result in any initial emissions, and the treatment
252 WGB5 even resulted in negative fluxes during the first week (Fig. 7B). After 110 days of
253 incubation, the cumulative total emissions were highest for straw treatments, reaching 3.51 and 9.17
254 mg C g⁻¹ soil emitted as CO₂ for Straw1 and Straw5, respectively. GB treatments resulted in
255 cumulative total emissions of 1.7 – 2 mg C g⁻¹ soil emitted as CO₂, slightly higher than the control
256 (1.65 mg g⁻¹ soil) (data not shown). Fig. 7C illustrates the cumulative fraction of added carbon
257 respired within 110 days. At the end of the incubation, 78 and 41 % of straw carbon added was
258 respired in treatments Straw1 and Straw5, respectively, while only 1-3 % of added biochar carbon
259 was respired.

260 Fig. 7 here.

261 4. Discussion

262 4.1. Soil carbon sequestration potential

263 A markedly smaller proportion of added carbon was respired in the GB treatments compared to the
264 straw treatments, which reflects the aromatic and recalcitrant structure of the residual carbon in
265 these biochar materials [4] after energy production during the process of gasification (Fig. 7C). The
266 addition of the high dosage of WGB resulted even in an initially negative CO₂ flux, probably
267 caused by binding CO₂ through carbonation of soluble Ca and Mg contained in the biochar, forming
268 CaCO₃ and MgCO₃ [23, 24]. The CO₂ peak after straw soil incorporation was reflected in the high
269 initial contents of DOC and SMB-C in these treatments, confirming that the easily degradable
270 carbon pool in the straw was rapidly decomposed by the soil microbial biomass, followed by a
271 decrease in CO₂ emissions (Fig. 7A). The very high content of SMB-C at day 1 in the high dosage
272 straw treatment was, however, surprising (Fig. 5B), and could be attributed to chloroform-labile
273 substances in the straw itself, as also suggested by Duong [25] observing similar effects.

274 The DOC level in both biochar treatments was – in accordance with their low CO₂ emissions -
275 significantly lower than that in straw treatments (Fig. 5A). WGB-treated soils were even lower in
276 DOC than SGB-treated soils, which could be due to a higher content of stable carbon, probably
277 caused by higher process temperatures during the wood gasification compared to the straw
278 gasification [26]. The DOC content of SGB was higher than that of the control treatment, but did
279 not result in any corresponding CO₂ emissions. This might be due to CO₂-binding by carbonation
280 occurring simultaneously with CO₂ emissions and therefore concealing soil respiration. However,
281 the DOC value in SGB treatments might also have been overestimated due to very small particles of
282 the biochar which were not retained by the filter during the extraction process. The DOC content in
283 WGB treatments was even significantly lower than in the control treatment, which might be

284 explained by a sorption of organic substances to WGB, as the SSA of wood biochar is very high
285 [14, 27]. This was also confirmed by the clear color of WGB extracts in contrast to the brownish
286 color of the other treatments. The DOC sorption by WGB could explain low CO₂ emissions and the
287 low content of SMB-C, as DOC is a carbon source for the microorganisms [27, 28]. However, the
288 adsorption of both DOC and microorganisms to biochar may potentially also result in higher
289 substrate consumption and therefore increase microbial activity [14]. Generally, our results confirm
290 that DOC-related parameters based on soil extraction procedures should be interpreted with caution,
291 as e.g. also Liang et al. [29] showed that the fumigation-extraction method leads to an
292 underestimation of SMB-C in biochar-amended soil due to sorption processes. The high N
293 mineralization observed in the WGB treatments is another indicator that soil microbial activity was
294 not inhibited by WGB (Fig. 4). Further studies are required to assess the effect of GB on soil
295 microbial biomass.

296 The GB carbon stability was also confirmed by their H/C and O/C atomic ratios, that had been
297 decreased compared to the original feedstock to values below 0.6 and 0.4, respectively (Table 2),
298 which is in agreement with the recommended thresholds indicating carbon recalcitrance [17, 26].
299 The H/C and O/C atomic ratios of WGB were even lower in comparison with SGB.

300 **4.2. Improvement of soil quality**

301 Results from BET and SEM analyses illustrated a higher SSA and porosity in WGB compared to
302 SGB (Table 4, Fig. 3). Besides the feedstock itself, the higher process temperature [19, 20, 27] in
303 the wood gasification process could contribute to those characteristics, as WGB and SGB were
304 produced at about 1000° and 700° C, respectively. However, both GBs in this study showed a
305 relatively high SSA in comparison with other studies, where the SSA of GBs ranged from 5 to 62
306 m² g⁻¹ [15, 19] and that of pyrolysis biochars from 1 to 320 m² g⁻¹ [20, 27, 30]. According to

307 Schimmelpfennig and Glaser [17], biochar with a SSA higher than $100 \text{ m}^2 \text{ g}^{-1}$ has the potential for
308 improvement of soil water and nutrient retention and porosity of the soil, which could benefit
309 microbes and plants. This requirement is definitely fulfilled by the WGB with an SSA of the same
310 magnitude as activated charcoal, which is probably due to the steam activation in the wood
311 gasification process [31]. The lower porosity of SGB is probably also caused by the processing, as
312 the straw fuel was pelletized and crushed, and gasified in a circulating fluidized bed (see section
313 2.1.). Cereal straw has about 6 times the amount of minerals (ash) compared to the wood chips used
314 to produce WGB, which might result in mineral matter occupying the pores of biochars or being
315 exposed at the surface of the biochar particles and blocking the pores, thereby causing the lower
316 SSA [32].

317 Addition of both biochars resulted in an increase of soil pH due to their alkalinity (Fig. 6). The
318 frequently described liming effect of biochar can improve plant nutrient availability, especially in
319 case of phosphorus in low-pH soils [3, 9, 27], and may have a beneficial effect on soil fertility and
320 plant growth on acidic soils [33].

321 Soil incorporation of straw with a wide C/N ratio often results in initial N immobilization [34, 35]
322 and subsequent slow N release [11]. The N immobilization was also observed in this study in the
323 straw treatments (Fig. 4). Contrarily, the soil application of GBs led to N levels similar to the
324 control soil, which means that no initial adverse effects on plant growth - as they can occur after the
325 application of pyrolysis biochar [36] - are to be expected after GB soil application. However, there
326 is no obvious explanation for the decreased N_{\min} levels compared to the control soils in the low
327 dosage of both GBs.

328 The total PAH content of both biochars was well below the threshold limit of 12 mg kg^{-1} for bioash
329 soil application according to the Danish Ministry of the Environment (Table 2). Eventual PAH

330 content in GB originates from PAHs in the produced gas, where they are formed as a decomposition
331 product of gaseous pyrolysis tars. If the GB stays in contact with the produced gas at low
332 temperatures, PAHs may subsequently condense on the GB. Although high PAH contents are often
333 reported for wood gasification biochars [4, 17], the WGB in this study showed a value of 0.69 mg
334 kg⁻¹, which is far below the limit, despite the high process temperatures. This is due to the
335 successful decomposition of PAHs during the TwoStage process, as the separation of the pyrolysis
336 and gasification reactors allows for a controlled gas phase partial oxidation of the pyrolysis tars
337 (Fig. 2). Consequently, the PAHs formed during the partial oxidation subsequently react with the
338 activated char in the char bed and are decomposed [37]. As a consequence of the in-process
339 decomposition, the concentration of PAHs in the produced gas is very low and hence no significant
340 PAH condensation on the WGB is possible [38]. Additionally, in the process, the WGB is separated
341 from the produced gas at high temperature (750 °C), which is significantly higher than the dew
342 point of the low PAH concentration in the gas and thus minimizes the possible condensation of
343 PAHs on the WGB.

344 ***4.3. Biomass for both energy and soil amendment***

345 Biomass, such as crop residues and wood waste, is a renewable global energy source, and efficient
346 energy conversion is required to reach the ambitious political goal in many countries to obtain a
347 fossil fuel free society. According to an LCA analysis by Nguyen et al. [2], gasification is - in
348 comparison with the dominating direct combustion - more environmentally friendly due to
349 primarily three main factors: (1) a higher energy efficiency, (2) reduced emission of major air
350 pollutants and (3) a higher carbon content in the residual fraction [2]. The LT-CFB process has
351 some unique features compared to direct combustion, as it can operate on crop residues and biomass
352 related waste, which are normally problematic for direct combustion. The produced gas has a low
353 content of ash alkali and can thus be combusted at high temperatures resulting in very efficient gas

354 utilization and energy conversion. The TwoStage gasification process allows for efficient utilization
355 of wood at small to medium scale. By producing clean and tar free gas, which can be used in a gas
356 engine for combined heat and power production, it is possible, even for a small scale plant, to
357 achieve efficiencies comparable with those of large scale power plants [1].

358 Crop residue removal for energy production can potentially reduce the soil carbon and nutrient
359 content and thereby the soil quality. Powlson et al. [11] concluded that removal or incorporation of
360 straw had a small effect on soil organic carbon content; however, even a small change in SOC could
361 have large negative impacts on soil physical properties. To date, the biochar fraction extracted from
362 the gasification process is not considered a valuable product, though, if it can be developed into a
363 soil amendment of high fertilizer and soil improver value, this will significantly improve the
364 economic feasibility and sustainability of the gasification technology [39]. On future markets, such
365 parameters have increasing importance, and the sustainability of a particular bioenergy chain will to
366 a large extent depend on the possibilities for its by-products recycling potential [40]. Nevertheless,
367 considering the complexity of effects of SOC on soil quality, the question, whether field application
368 of gasification biochar may replace SOC originating from crop residues, requires further research.

369 In contrast to pyrolysis, which is usually engineered to produce biochar with gas and heat as co-
370 products, the main product of gasification is energy in form of syngas, while biochar is considered a
371 co-product. Thus, gasification produces more energy and less biochar compared to pyrolysis [18]. It
372 is, however, important to find a balance in the amount of carbon utilized for energy generation and
373 carbon left in the biochar for soil application. In the present study, we had a focus on both energy
374 and biochar production. In the LT-CFB process, 90 % of the feedstock-carbon was used for energy
375 production, while 10 % remained in the biochar (Table 1). In the TwoStage process, 96 % carbon
376 was utilized for energy and 4 % remained in the biochar. Therefore, LT-CFB gasification of straw
377 and biochar soil amendment could on the longer term have a comparable soil carbon sequestration

378 potential to the TwoStage gasification of wood, despite the fact that WGB carbon showed a higher
379 stability compared to SGB. Currently, the LT-CFB gasification processes are flexible technologies,
380 allowing an energy output of up to 97 % of the carbon input, which would reduce the SGB's carbon
381 content from the present ca. 50 % to 20 – 30 %.

382

383 **5. Conclusion**

384 In this study, we suggest that thermal gasification of biomass residues is able to combine the
385 production of bioenergy and a biochar fraction that can exert a positive impact on soil quality. Our
386 results showed that gasification biochar (GB) carbon is more resistant to microbial degradation
387 compared to straw carbon and has a potential for soil carbon sequestration. Furthermore, the GBs in
388 our study exhibited a potential as soil improving agents due to their high specific surface area,
389 porosity and liming effect, with PAH contents below the threshold limit. However, the differences
390 found between the two biochar materials will probably qualify them to benefit different soil
391 parameters. WGB with higher SSA, lower PAH content and higher carbon stability, caused both by
392 feedstock source but also by process conditions, could increase water holding capacity and nutrient
393 retention on sandy soils, while SGB could be preferably used as a fertilizer or liming agent.
394 Gasification of straw and wood chips and field application of the biochar is therefore an integrative
395 approach combining both agriculture and forestry with the energy sector, which seems to be an
396 attractive option to maximize both energy output and soil carbon sequestration. The results of the
397 present study reveal that it is worthwhile to further test the potential of GB soil amendment, as it
398 has been done for more traditional pyrolysis biochar materials [26, 27, 34]. In this regard, it will be
399 crucial to investigate the soil application of GBs also in longer-term studies, pot and field
400 experiments, to be able to determine the effect on plant yields, soil biota and soil quality.

401

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408

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514

515 Figure Captions:

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517 Fig. 2 – Schematic of the TwoStage gasifier [1].

518 Fig. 3 - Scanning electron microscope (SEM) images; left: straw gasification biochar (SGF) and
519 right: wood gasification biochar (WGB).

520 Fig. 4 – Content of soil mineral nitrogen (N_{\min}) during the incubation period of 110 days. Straw1=
521 soil amended with 1% straw, Straw5= soil amended with 5 % straw, SGB1= soil amended with 1 %
522 straw gasification biochar, SGB5= soil amended with 5 % straw gasification biochar, WGB1= soil
523 amended with 1 % wood gasification biochar, WGB5= soil amended with 5 % wood gasification
524 biochar, Control= untreated soil. Values presented are means with standard error bars ($n =4$).
525 Treatments with different letters are significantly different at the last day of the incubation ($P <$
526 0.05).

527 Fig. 5 – A) Content of dissolved organic carbon (DOC) in soil during the incubation period of 110
528 days. B) Content of soil microbial biomass-carbon (SMB-C) in soil during the incubation period of
529 110 days. For treatment abbreviations, see Fig. 4. Values presented are means with standard error
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532 Fig. 6 – Soil pH at day 1, 8, and 110 of the incubation period. For treatment abbreviations, see Fig.
533 4. Values presented are means with standard error bars ($n =3$). Treatments with different letters are
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535 Fig. 7 – A) CO_2 fluxes from soil during the incubation period of 110 days. B) CO_2 fluxes during the
536 first 8 days of incubation. C) Cumulative fraction of added carbon respired from soil during the

537 incubation period of 110 days. For treatment abbreviations, see Fig. 4. Values presented are means
538 with standard error bars ($n = 4$). Treatments with different letters are significantly different at the
539 last day of the incubation ($P < 0.05$).

540

Table 1 – Carbon and energy balance for TwoStage gasifier and Low-temperature circulating fluidized bed gasifier (LT-FCB) reflecting the carbon loss in the GB used in this study.

	Percentage TwoStage input		Percentage LT-FCB input	
	Fractional distribution			
	Carbon (%)	Energy (%)	Carbon (%)	Energy (%)
Biomass feedstock	100	100	100	100
Product gas output	96	92	90	85-87
Biochar output	4	4	10	10
Loss	-	4	-	3-5

Table 2 - Chemical characterization of soil, feedstock and biochars (SGB = straw gasification biochar, WGB = wood gasification biochar).

	Soil	Straw	Wood chips	SGB	WGB
C (%)	1.98	45.50	52.04	46.80	65.29
H (%)	-	5.52	7	0.97	0.63
O (%)	-	36.85	41.16	13.11	8.99
H/C atomic ratio	-	1.46	1.61	0.25	0.12
O/C atomic ratio	-	0.61	0.59	0.21	0.10
pH (water)	7.9	-	-	11.6	11.1
Ash (%)	-	4.85	0.75	52	33
Σ PAH ^a (mg kg ⁻¹)	-	-	-	5	0.69

^a Sum of Acenaphthene, Fluorene, Phenanthrene, Fluoranthene, Pyrene, Benzo(bjk)fluoranthene, Benzo(a)pyrene, Indeno(1,2,3-cd)pyrene and Benzo(ghi)perylene.

Table 3 - Particle size distribution of straw gasification biochar (SGB) and wood gasification biochar (WGB).

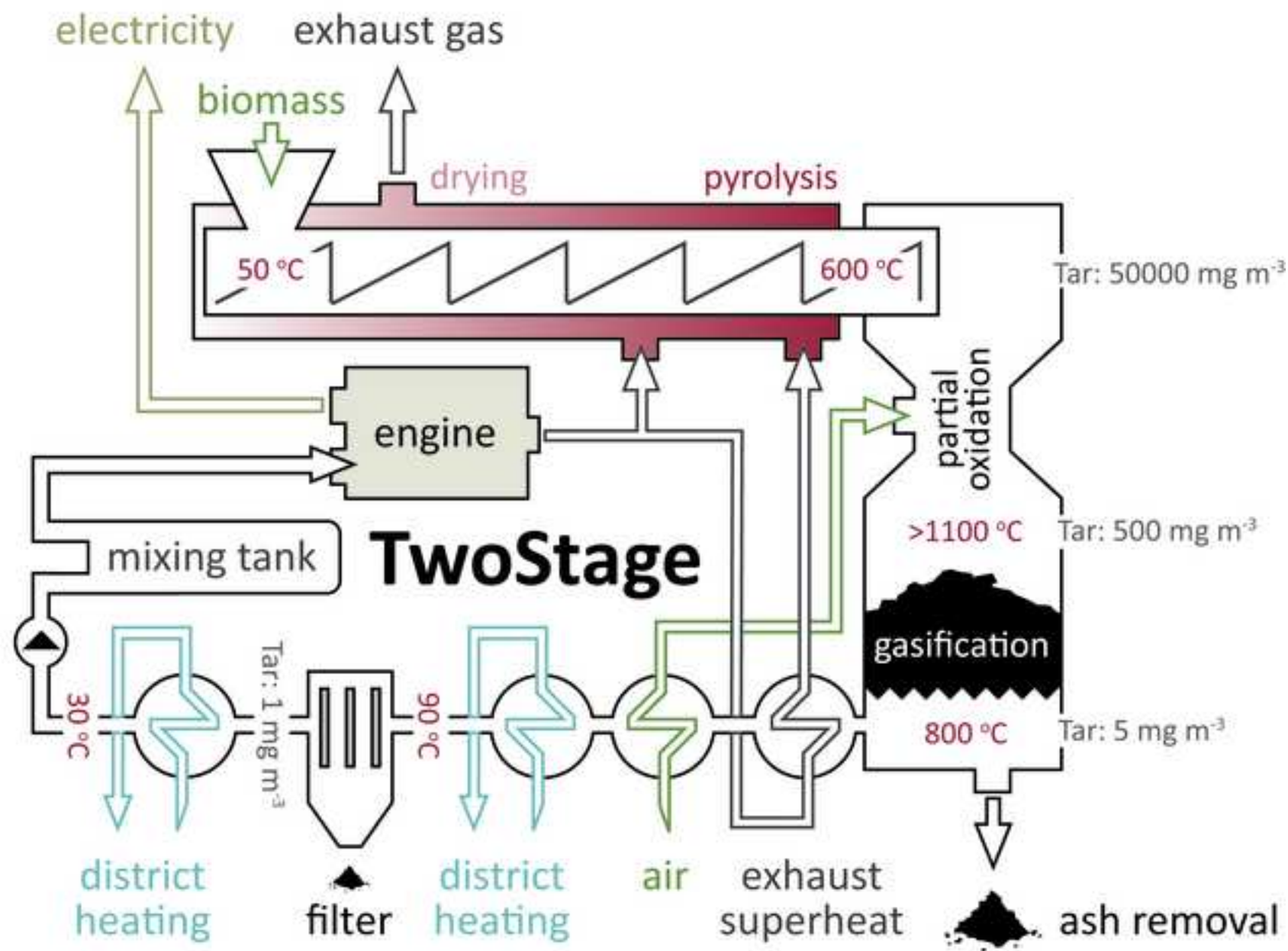
Biochar	Particle size distribution in % of dry mass		
	< 0.045 mm	0.045-0.125 mm	>0.125 mm
SGB	89.4	10.3	0.3
WGB	33.0	13.7	53.3

Table 4: BET specific surface area (SSA), pore volume and diameter of straw gasification biochar (SGB) and wood gasification biochar (WGB).

Biochar	Particle size (mm)	SSA ($\text{m}^2 \text{g}^{-1}$)	Pore volume ($\text{cm}^3 \text{g}^{-1}$)	Pore diameter (nm)
SGB	0-1	75	0.04	3.71
WGB	0-0.5	426	0.52	1.43
WGB	0.5-1	1027	0.58	3.73

Figure2

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All tar concentrations measured per volume gas produced.

All gas volumes measured at Normal Temperature and Pressure i.e. $20\text{ }^{\circ}\text{C}$ and $101325\text{ kg m}^{-1}\text{ s}^{-2}$

Figure4

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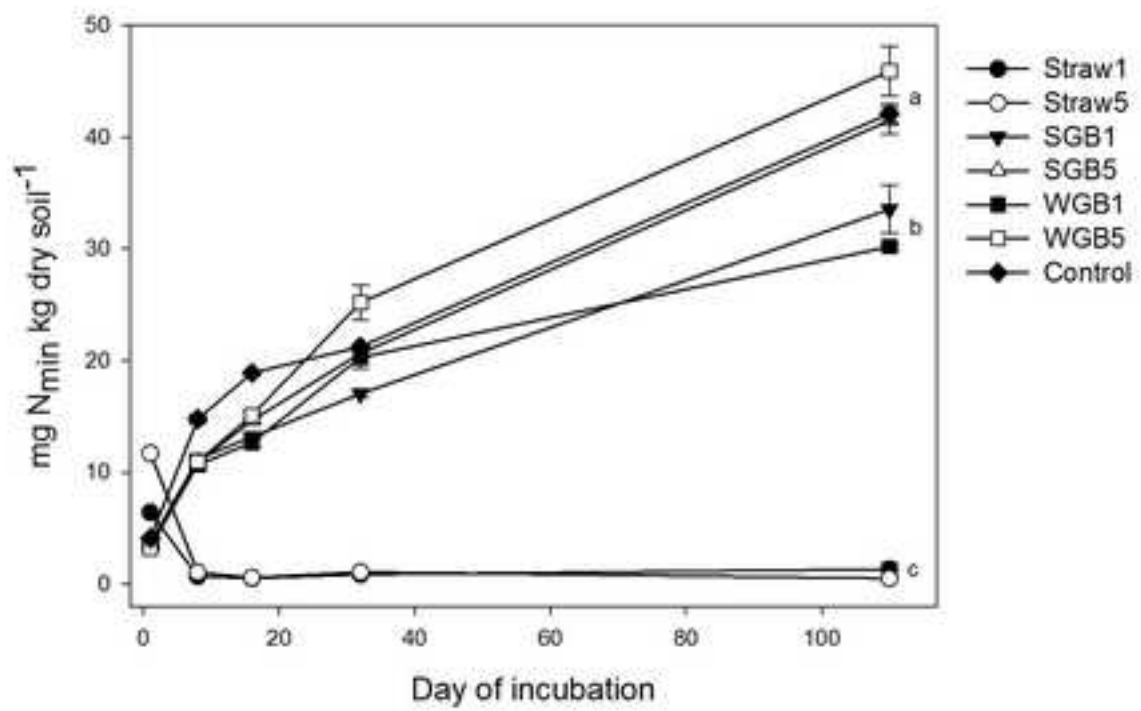


Figure5

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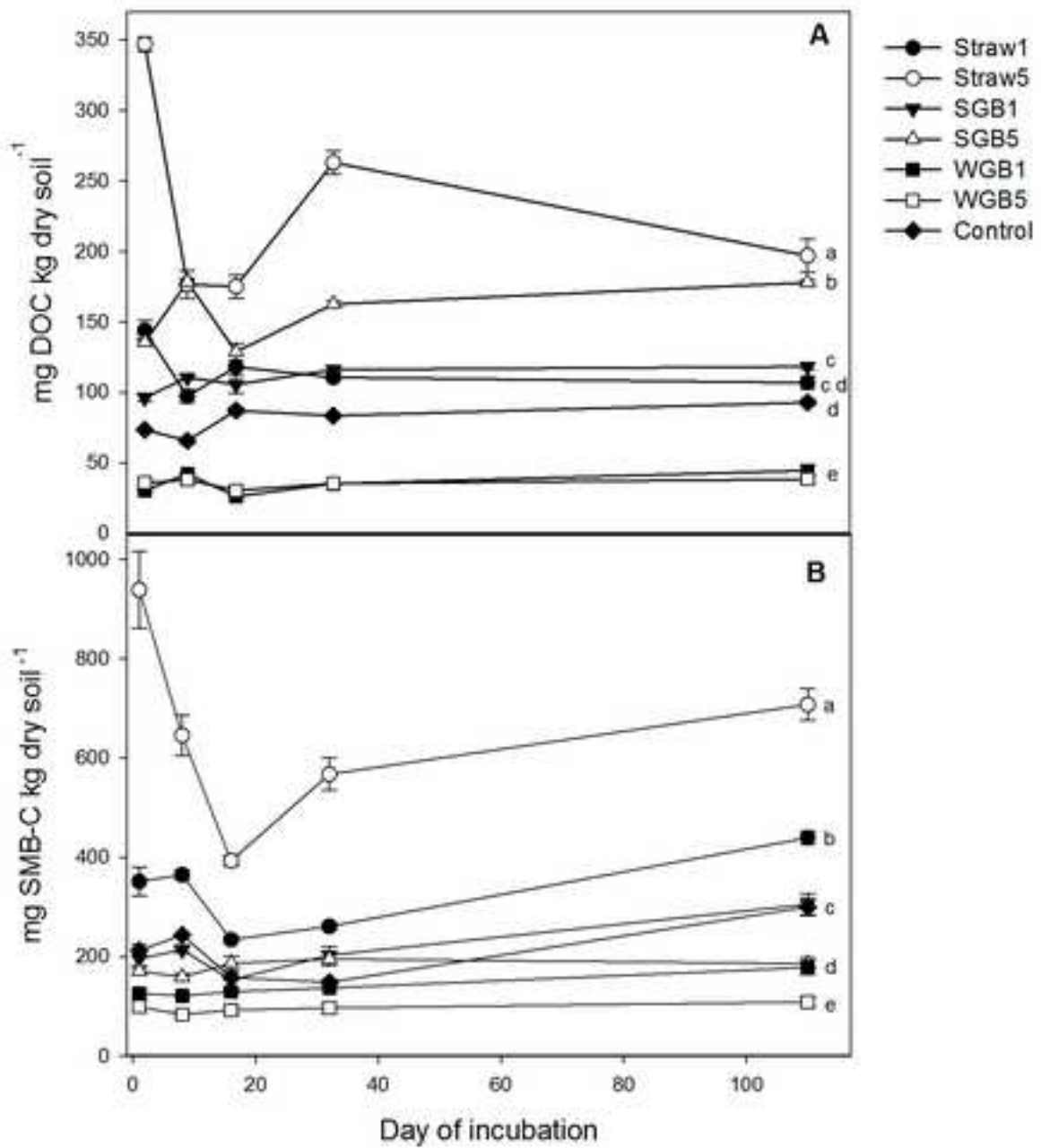


Figure7

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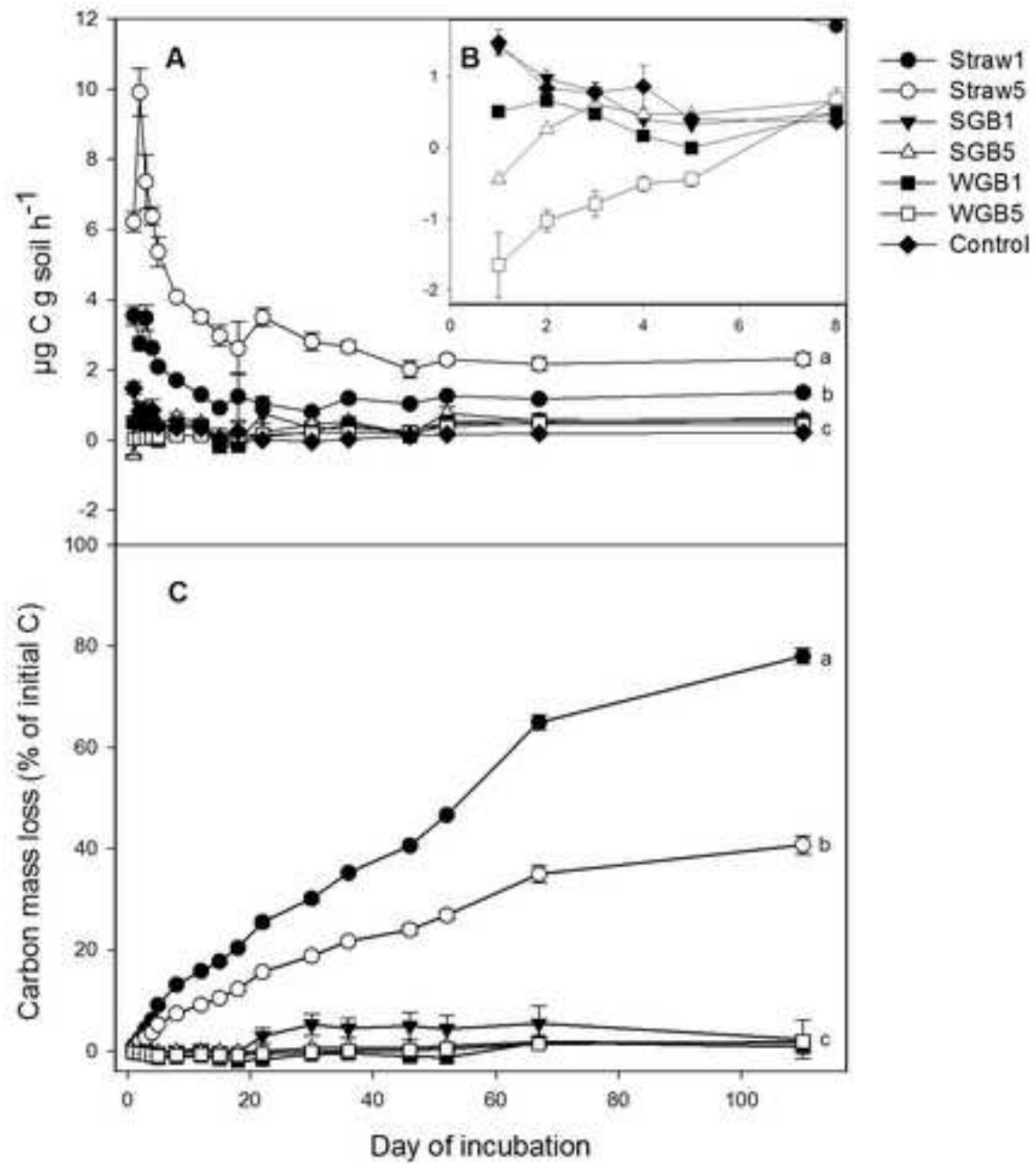


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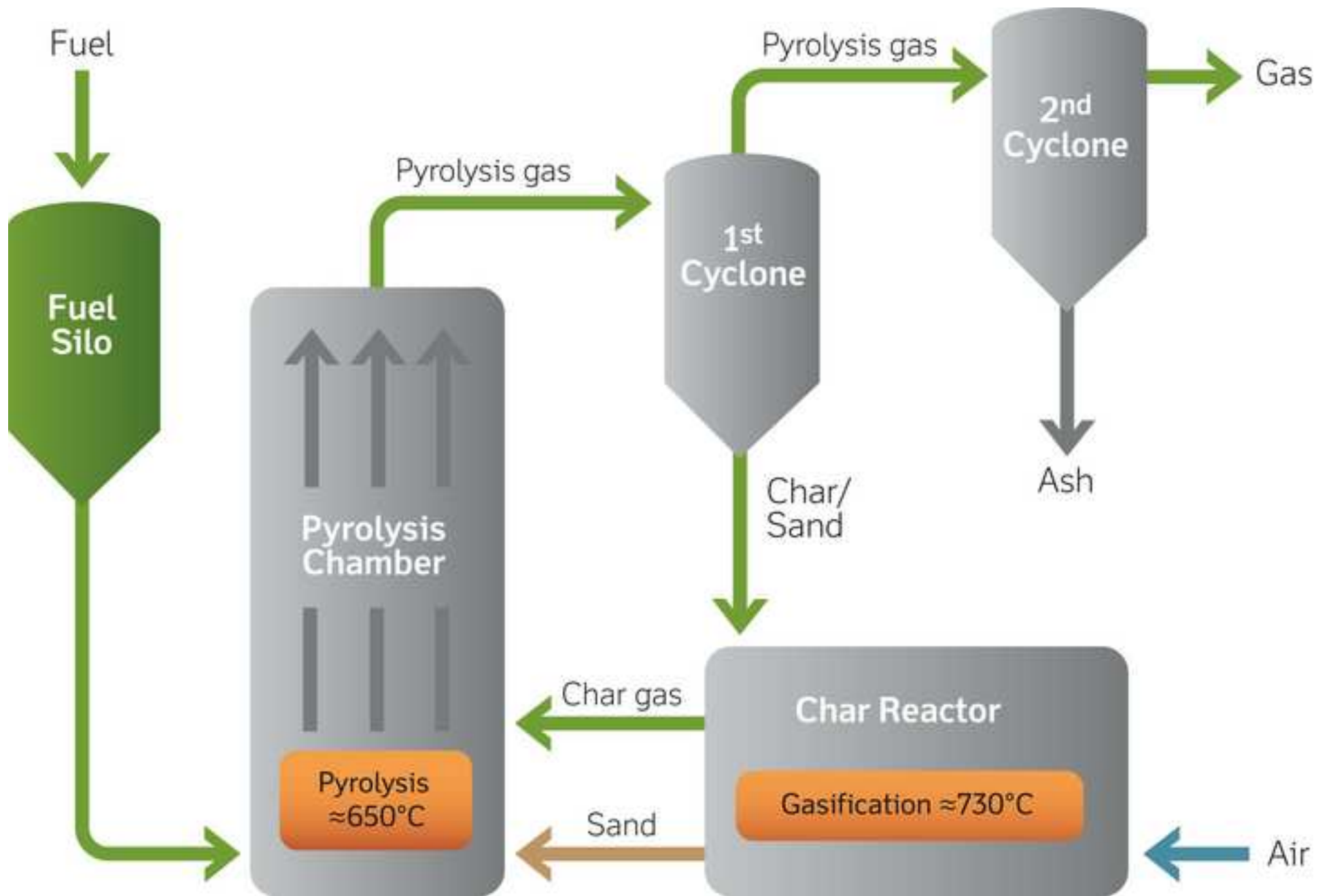
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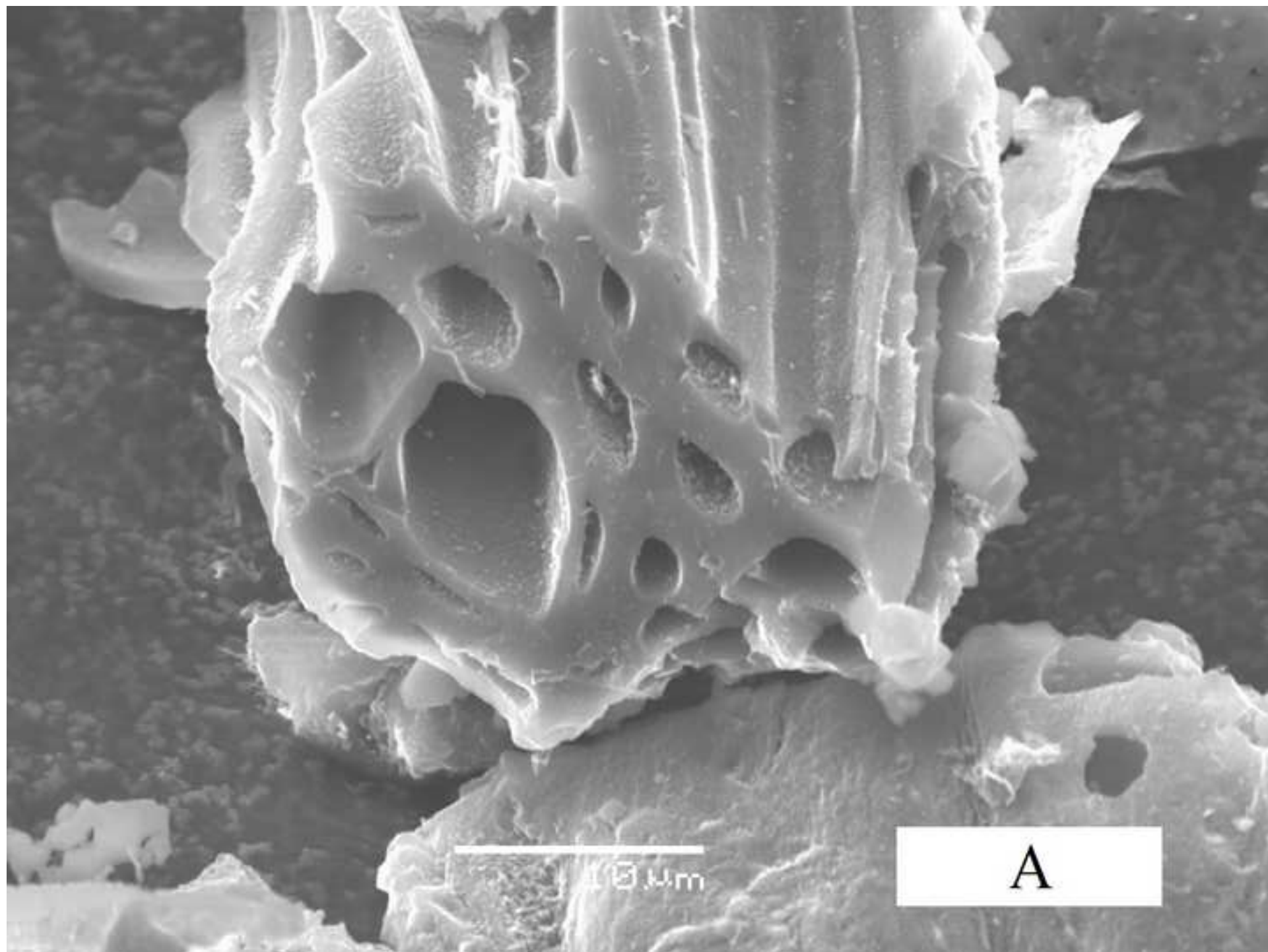
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Figure
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Figure

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