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Original article

Material and moisture balance in a full-scale bio-drying MBT system for solid recovered fuel production

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ABSTRACT

Bio-drying MBT is a type of mechanical biological treatment (MBT) system and reduces moisture content of the MSW to improve the separation of combustible fractions. In this study, a full-scale biocell type bio-drying MBT system was investigated. The mass balance of waste component was estimated by composition and characterization of waste and tonnage data. During separation of biodried outputs, 62 % of plastics and 54 % of paper were recovered as RPF material. Wood was decreased by reduction in particle size and 90 % of biodried wood is returned to next reactor. Changes of mixed fine caused by fine wood particle and the loss of organic matters and 60 % of it were returned. Daily water removal during 17-days of bio-drying was simulated through the model by using the operation data. Among the four operation phases, the longest stabilization phase was expected to main water removal period, but half of water removal was occurred at initial two stages and phase IV for only 6

days in total due to the high waste temperature for sanitization (phase I and II) and high airflow rate for cooling. Decreasing waste temperature at phase III resulted in low water evaporation.

Keywords: Bio-drying MBT, Material balance, Water removal, Material recirculation, Solid recovered fuel recovery

1 **Introduction**

2 Formerly, many European Union (EU) countries relied heavily on landfills for waste
3 treatment and disposal. However, as a substantial amount of municipal solid waste (MSW)
4 containing a high portion of biodegradable waste was disposed of at landfills, management of
5 the resulting leachate and gas emissions caused from the biodegradation of organic matters
6 required long-term care [1]. To cope with this problem, the European Commission
7 established the Landfill Directive (99/31/EC), which required the reduction of biodegradable
8 waste in landfills. In response, mechanical-biological treatment (MBT) systems have been
9 developed to reduce the amount of biodegradable waste sent to landfills [2, 3].

10 In a typical MBT system, mixed MSW is mechanically separated and then undergoes
11 sequential biological and thermal treatments to stabilize organics and the remaining
12 combustible fraction. The resulting stabilized residue is then sent to a landfill for final
13 disposal. MBT systems also present an opportunity for energy recovery in the form of biogas
14 from anaerobic digestion (AD) or a solid recovered fuel (SRF). SRF production has been
15 preferred as it enables thermal recovery in a variety of end-uses at cement kilns or other co-
16 combustion power plants as an alternative to fossil fuels [4–7]. In order to recover more SRF,
17 bio-drying MBT systems can be applied to improve the separation efficiency of the
18 combustible fraction by reducing the moisture content of MSW at first process.

19 The reactors used in bio-drying MBT systems can be divided into three types: dynamic
20 reactors, static reactors, and windrows. Dynamic reactors are used in continuous systems with
21 counter flow aeration against the waste movement and consist of an inclined rotary drum that
22 is 2 to 4 m in diameter and a maximum of 45 m long. Typical static reactors are enclosed
23 biocell or biocontainer batch-type reactors. A biocell is an enclosed rectangular reactor that
24 has a volumetric capacity in the range of 100 to 1000 m³ and is, at maximum, 50 m long.
25 Waste is charged to the reactor by a wheel loader or conveyor belt. A biocontainer is smaller

26 box reactor with a volume of 20 to 40 m³ in which waste is loaded from the top of the reactor.
27 A windrow pile system has a pile in a triangular or trapezoidal shape where waste is stacked
28 up to 1.5 or 2.0 m high. In static-type and windrow reactors, forced aeration is accomplished
29 from the bottom. Most full-scale Bio-drying MBT systems in operation are enclosed static-
30 type reactors [2, 3, 8].

31 Tambone et al. investigated the fuel quality and biogas generation potential of biodried
32 outputs from a windrow pile system by analyzing the heating value, respiration index, and
33 biochemical methane potential (BMP) test [9]. Dębicka et al. analyzed the heating value and
34 respiration index of a biocell that has a 150 m³ to determine the moisture reduction and fuel
35 qualities [10]. Evangelou et al. monitored the composting process of a 360 m³ biocell-type
36 bio-drying MBT system for 1.5 years to evaluate organic stabilization and the qualities of the
37 resulting fuel by measuring the dynamic respiration index and heating values [11]. Dziedzic
38 et al. studied a 36 m³ biocontainer to investigate the fuel qualities and biogas generation
39 potential of biodried outputs [12].

40 Unlike the above MSW-treatment facilities, Winkler et al. studied a 1900 m³ biocell-type
41 reactor used to treat sewage sludge [13]. In this system, air was recirculated and biodried
42 sludge was returned as a bulking agent and an inoculum. They evaluated the water removal of
43 sewage sludge and estimated the evaporation during the bio-drying process by using the
44 operation data.

45 In this study, an operating 900 m³ biocell-type bio-drying reactor employing material and
46 air recirculation was investigated. The produced fuel is recovered after 17 days of the bio-
47 drying process, at which point wood and fine residue are returned to the next reactor. Air in
48 the reactor is recirculated to the reactor by changing the fresh air intake ratio.

49 Unlike prior studies, this work focuses on the material balance and model estimation of
50 water removal. The material balance of changes during the bio-drying process and during the

51 separation of biodried materials is estimated by analyzing samples in terms of proximate
52 analysis, total organic carbon (TOC) content, and BMP. Water removal is estimated by
53 analyzing the monitored operation data.

54

55 **Materials and Methods**

56 **Investigation of bio-drying MBT system**

57

58 **Fig. 1** Description of the investigated bio-drying MBT system: (a) process flow and annual
59 mass in the facility and sampling location and (b) schematic diagram of airflow in the reactor

60

61 A process flow diagram of the investigated bio-drying MBT system is shown in Fig. 1a.

62 Six biocell reactors (BR) are installed and operated in turn. Each BR has a capacity of 900 m³,

63 6 m wide, 5 m high, and 30 m long. After 17 days of bio-drying operation, biodried output is

64 separated into three fractions: raw material for refuse-derived paper and plastics densified

65 fuel (RPF), wood residue and fine residue. Residues are transferred to next BR as bulking

66 agent and inoculation. They are mixed with shredded MSW by a shear shredder (Ø 460 mm)

67 and new wood occasionally, then transferred by a wheel loader to the BR. Water (11.5 to 20.7

68 m³) is added to the waste during the first stage of the bio-drying process to promote the

69 biodegradation with mixing generated leachate. Approximately 260 tons of input mixture is

70 processed in one BR. The biodried waste is sorted into heavy, light, and fine matters by a

71 ballistic separator, which is tilted 20 degrees and has screened moving paddles. The heavy

72 and light fractions are separated by gravity with upward airflow. Any inert materials or PVC

73 are removed using an NIR separator.

74 In October 2018, a total of 30 kg of waste samples were collected from five different

75 locations in the facility, indicated with an S in Fig. 1a. All samples except for the BR input

76 was taken from one of the BRs after 17 days of operation. BR output, wood residue, and fine
77 residue samples were taken from each pile, and RPF material samples were taken from bailed
78 products. The BR input sample was collected from next BR that was in the filling stage
79 before the process initiation. Residues contained in BR input sample were the outputs of
80 previous BR. MSW and wood were not sampled because MSW was mixed with return
81 residues just after shredding, and wood was hard to collect representative sample due to large
82 size.

83 The tonnage shown in Fig. 1a indicates the total amount from April 2017 to March 2018.
84 Waste mass was measured either by truck scale (TS) or weighing scale on the wheel loader
85 (LD). Water added to the BR was measured through a flowmeter (FM). The input and output
86 mass of each BR were estimated from the summation of each waste stream.

87 The overall airflow through the reactor is shown in Fig. 1b, in which the mixture of fresh
88 air and recirculating air from the BR is used for aeration of the waste. During the process, the
89 airflow rate and mixing ratio of fresh air is controlled according to the waste temperature.
90 The details of the control mode and variables shown in the figure are explained in final
91 section.

92

93 **Laboratory analyses and procedures**

94 To determine moisture content, the collected samples were dried in a drying oven (hot-air
95 circular drying machine, Toyo) at 50 °C until the changes of sample weight showed below 1%
96 variance. Samples were manually turned two times a day. Moisture content was determined
97 by the difference of weight for each sample. Dried samples were hand-sorted into six
98 components: plastics, paper, textiles, wood, incombustibles, and mixed material. Mixed
99 material was then sieved with a 4.0 mm screen, at which point the oversized fraction was
100 sorted into its components and undersized fraction was categorized as mixed fine.

101 For further analysis, samples were ground by cutting mill (MRK-Retsch Ultra cutting mill,
102 Country). As this mill could not grind textiles, they were cut into small pieces using scissors
103 and then ground with a freeze grinder (AS One Freeze grinder, TPH-02). All samples were
104 analyzed for combustible and TOC content. A BMP test was conducted to determine biogas
105 generation as an index of biodegradable organics.

106 To measure the combustible contents, 5 g of each dried sample were placed in a crucible
107 and ignited using a muffle furnace (Box furnace KBF-894N1) at 800 °C for 2 hours. The
108 TOC content was determined by the difference between the total carbon (TC) and inorganic
109 carbon (IC) of 30 mg dried samples using a TOC-TN analyzer (TOC-V CPH/CPN, connected
110 with TOC-V SSM-5000A Shimadzu Corporation). Pure glucose (C₆H₁₂O₆) and sodium
111 carbonate (Na₂CO₃) were used as a standard carbon source for the TC and IC analysis,
112 respectively.

113 The BMP was experimentally analyzed as done by Hansen et al. and Pantini et al. [14, 15].
114 Here, one gram of the sample was mixed with 40 mL of distilled water and 20 mL of
115 inoculum in a 135 mL vial. The samples were then purged with nitrogen to remove traces of
116 oxygen and sealed with a butyl rubber stopper and aluminum crimps. The test materials were
117 incubated at 55 °C for 28 days; paper and textiles were incubated for 35 days due to
118 prolonged biodegradation. The inoculum used was taken from the liquid sludge of an AD
119 process at Fuji Clean Center in Kagawa, Japan.

120 Generated biogas was extracted by inserting a 50 or 100 mL syringe until the piston
121 movement stopped, at which point the gas volume generated was measured. Then, 0.1 mL of
122 gas was withdrawn using a 0.5 mL gas syringe; the gas composition was determined by gas
123 chromatography (GC-TCD, HITACHI, type 164). Gas was extracted every day during the
124 initial stage and every 2–3 days afterwards. Gas generation was adjusted by subtracting the

125 amount found in a blank containing only water and sludge. After 28 or 35 days (depending on
126 the material) of testing, the gas composition of the air phase of the vial was also measured.

127

128 **Results and Discussion**

129 **Physical composition and waste characterization**

130

131 **Fig. 2** Physical composition of waste on a wet basis

132

133 The physical composition of the five samples taken from the facility is shown on a wet
134 basis in Fig. 2. From the input to the output, the moisture content decreased from 40% to 28 %
135 from BR and rest were shown around 20 % or less. In the wood and fine residue, a
136 considerable amount of wood and mixed fine material was contained. Because MSW and
137 wood were not available for sampling as mentioned in above, the composition of MSW and
138 new wood was assumed based on the literature values. The data of MSW was referred to the
139 report of incineration facility in Mitoyo city [16]. High moisture content of MSW is caused
140 by 61 % of food wastes. Large sized hard wood is introduced to keep porosity [17].

141

142 **Fig. 3** Waste characteristics of (a) combustible content and (b) gasified carbon in biogas

143

144 The determined TOC content, combustible content, and gasified carbon in biogas through
145 BMP test are compared in Fig. 3, where each component is plotted using same symbols, but
146 BR input samples are differentiated by no colored mark, as they contained fresh MSW. The

147 striped mark in each component indicates the characteristics of MSW and new wood as
148 reference [17, 18]. Food wastes that only exist in MSW were marked by star mark.

149 In Fig. 3a, the TOC/combustibles indicate the characteristics of the combustible fraction,
150 where a low combustibility was caused by contaminated inorganic materials. The broken
151 lines in Fig. 3a represent the characteristics of polyethylene and cellulose [17]. The
152 TOC/combustibles of plastics is similar to polyethylene. The TOC/combustibles of textiles
153 that is slightly higher than cellulose suggests that they are composed of synthetic and natural
154 fibers. The TOC/combustibles of the mixed fine component was similar to that the textile
155 component, suggesting that it contained high-carbon-content organics, such as humic
156 substances. Referred characteristics of MSW and new wood are similar to the sample analysis
157 results.

158 The gasified carbon by anaerobic digestion, or gasified carbon in the biogas, determined by
159 the BMP test and TOC content of waste components are compared in Fig. 3b. The ratio of
160 gasified carbon to TOC is 60% for paper, 5%–10% for wood and plastics, and 30% for
161 textiles and mixed fine components. The generated gas from plastics was caused by attached
162 organic material. The determined biogas generation of textiles is reasonable, considering they
163 likely consisted of synthetic and natural fibers. The low ratio of gasified carbon to TOC in the
164 mixed fine components is reasonable, as they consist of stabilized and/or hardly
165 biodegradable organics output from the BR.

166 Comparing each component in Fig. 3a and 3b, no significant differences were seen among
167 the samples, even between the input to and output of the BR. This is likely because the partial
168 reduction of organic matters after the 17-day bio-drying process is only the decrease of the
169 amount and its characteristics including BMP test was not changed.

170

171 **Component mass fraction by waste stream**

172

173 **Fig. 4** Mass fraction of (a) dry solids, (b) gasified carbon in biogas, and (c) combustibles of
174 each waste stream component on a dry basis, normalized by the total dry mass of the INPUT

175

176 Fig. 4 was obtained by multiplying the mass in Fig. 1 with the physical composition in Fig.
177 2 for each waste stream in dry basis excluding moisture. As shown by the arrows on the top
178 of the figure, the sum of RPF materials, wood residue and fine residue were output from the
179 BR, and the input to the BR consisted of wood residue, fine residue, MSW, and new wood.
180 Results of sample analysis for BR input and BR output were not used in the calculation as it
181 was considered less creditable than the summation of other samples in terms of the
182 homogeneity.

183 Fig. 4a was normalized by the total dry mass of the input. In this figure, 76% of wood
184 originates from the return residue, whereas 17% is provided from new wood. For the RPF
185 materials, i.e., plastics and paper, 60% was introduced through the MSW stream, whereas 40%
186 was from returned residue.

187 Fig. 4b–c were calculated similarly for gasified carbon in biogas and combustibles by
188 multiplying their characteristics (Fig. 3) with the dry mass of each component in Fig. 4a. The
189 mass fraction of combustibles in Fig. 4c looks similar to the dry solids in Fig. 4a.

190

191 **Separation efficiency of output from BR**

192

193 **Table 1.** Separation ratio of the output streams on a dry basis

194

195 The separation ratio of the output from the BR into the three streams for total dry mass and
196 each waste component was calculated from the data in Fig. 4a and is shown in Table 1.

197 Of the fuel materials, only 62% of plastics and 54 % of paper was recovered as a fuel and
198 the rest were returned to the next BR. To investigate the cause of unrecovered materials, the
199 size distribution for RPF materials and wood residue was measured and is shown in
200 Supplementary material. The unrecovered ratio was approximately 40% and was considered
201 as the performance of wind separation. The size of recovered plastic particles ranged from 5
202 cm to 65 cm. The recovery rate of plastic particles to RPF materials under 15 cm in size was
203 only 28%. However, the paper components were mostly smaller than 20 cm and had a
204 recovery rate to RPF of 54%. Wood and the mixed fine component were separated into return
205 stream more effectively; 90% of the wood and 60% of the mixed fine components were
206 returned to the mixer before being introduced to the next BR.

207

208 **Mass balances before and after the bio-drying process**

209

210 **Fig. 5** Mass balances of (a) dry solids, (b) gasified carbon in biogas, and (c) combustibles in
211 the bio-drying process on a dry basis

212

213 The difference in dry mass of each component between the input and output of the bio-
214 drying process is shown in Fig. 5a, where the mass fraction of each component was
215 calculated by summing up for INPUT and OUTPUT in Fig. 4a, and they are depicted by bar
216 graph and dot, respectively. The food waste included in the MSW was added to the mixed
217 fine component in the process input stream. Fig. 5b–c, for gasified carbon in biogas and
218 combustibles, were calculated from Fig. 4b–c in the same manner.

219 The small reduction of paper shown in Fig. 5a is reasonable, as paper does not fully
 220 degrade even during the 6 to 8 weeks of the composting process [19]. As wood is a hardly
 221 biodegradable organic, the decrease of wood may have been caused by a reduction in particle
 222 size, allowing the fine particles to be transferred to the mixed fine component. The reduction
 223 in Fig. 5b can be explained by the transfer of wood, and the loss of organic matters can be
 224 another reason. Also, no changes of combustibles of mixed fine in Fig. 5c also proved by
 225 same reason.

226 The gasified carbon in biogas was reduced by 13% between the input and output of the
 227 bio-drying process (Fig. 5b). Among the waste components in the input, food wastes in the
 228 MSW and the mixed fine component could be biodegraded in 17 days of the bio-drying
 229 process, and their fractions were 13% and 9% of the input, respectively. A decrease from 21%
 230 to 13% can be considered reasonable for biodegradation.

231

232 **Water removal during the bio-drying process**

233 To estimate the water removed during bio-drying, the evaporated water mass was then
 234 calculated. As shown in Fig. 1b, the four unknown airflow rates were V_A , V_F , V_{RE} , and V_{EX} ,
 235 where subscripts A, F, RE, and EX refer to aeration, fresh air, recirculating air, and exhaust
 236 air, respectively. A mixture of fresh air and recirculating air from the BR is aerated to the
 237 same BR. The mass and heat balances before and after the mixing of air can be written as Eq.
 238 (1) and (2), respectively.

$$V_A = V_F + V_{RE} \quad (1)$$

$$V_A T_A = V_F T_F + V_{RE} T_{RE} \quad (2)$$

239 The ratio of fresh air to total aeration λ_F can be defined as Eq. (3). The recirculating airflow
 240 rate can be then determined as Eq. (4). Equation (5), derived from Eq. (2) by substituting V_F
 241 and V_{RE} with Eq. (3) and (4), respectively, can then be used to calculate λ_F .

$$\lambda_F = \frac{V_F}{V_A} \quad (3)$$

$$V_{RE} = (1 - \lambda_F) \times V_A \quad (4)$$

$$T_A(^{\circ}\text{C}) = \lambda_F \times T_F + (1 - \lambda_F) \times T_{RE} \quad (5)$$

242 In Eq. (5), T_{RE} was assumed equal with waste temperature (T_S). The exhaust airflow rate,
 243 V_{EX} , was determined to equal V_F by assuming steady state. The value of T_F was assumed as
 244 22.8 °C with 57.9% relative humidity (RH) based on meteorological data [20]. The total
 245 aeration flow rate V_A can then be calculated by Eq. (6) where 16000 is the specified airflow
 246 rate of the fan in m^3/h , and q_A is the fan speed as a percentage.

$$V_A (\text{m}^3/\text{h}) = 16000 \times q_A \quad (6)$$

247 The water removal rate in the BR can then be calculated as the difference between the
 248 water inlet through fresh air from water outlet, shown in Eq. (7), where X represents the water
 249 vapor per unit volume of air in g/m^3 , which is a function of water vapor pressure (pv) and
 250 temperature (T) (Eqs. (8)–(10)).

$$\text{Water removal rate (kg/h)} = (V_{EX} \times X_{EX} - V_F \times X_F) \times 10^{-3} \quad (7)$$

$$X (\text{g}/\text{m}^3) = \frac{217 \times pv}{273.15 + T} \quad (8)$$

$$pv (\text{Pa}) = RH \times pvs \quad (9)$$

$$\text{When saturated: } pvs (\text{Pa}) = 6.1078 \times 10^{\frac{7.5 \times T}{T+237.3}} \quad (10)$$

251 As shown in Eq. (7), water removal rate is mostly governed by air flow rate V_{EX} and X_{EX} ,
 252 which is water vapor per unit volume of air. X_{EX} increases exponentially with temperature by
 253 Eq. (8)-(10). So, airflow rate and temperature are two major factors for moisture removal.

254 Exhaust air was assumed to always be saturated and its temperature (T_{EX}) was equal to the
 255 waste temperature (T_S).

256

257 **Fig. 6** Profiles of (a) operation variables and (b) water removal rate under different
258 operation phase during the bio-drying process

259

260 The profiles of operation variables are shown in Fig. 6a. Points of irregular operation were
261 removed from the figure but considered in the calculation. The process was subjected to four
262 consecutive operation phases for 17 days. During warm-up (I), a low airflow rate was used to
263 warm to and then maintain the waste at 70 °C for sanitization purposes (II). During
264 stabilization (III), moisture was removed by increasing the airflow rate. The ratio of fresh air
265 was decreased with the progress of biodegradation, as indicated by lowering waste
266 temperature. During the cooling phase (IV), a maximum airflow rate of fresh air (i.e., $\lambda_F = 1$)
267 was provided.

268 The daily water removal rate as calculated by Eq. (7) is shown in Fig. 6b, where the
269 numbers on the figure indicate the portion of water removed at each phase over the total
270 water removal. Half of the water removal occurred during phase III, which had the longest
271 elapsed time, and the moisture removal rate declined with the decrease of temperature. Phase
272 I and II showed relatively higher removal rate despite their of short duration and low airflow
273 rate, as the saturated moisture content in air is 198 g/m³ high at 70 °C while 104 g/m³ at
274 55 °C. In phase IV, the water removal rate was lowered due to the decrease in temperature,
275 but the maximum flow rate of fresh air for 3 days enhanced moisture removal.

276 The estimated total water removed, as shown in Fig. 6b, was 86.6 tons. Based on the
277 material flow in Fig. 1 and the moisture content in Fig. 2, the amount of water removed was
278 55.5 tons, from 105.3 ton to 49.8 ton for 260 tons of wet waste per BR. The difference may
279 have been caused by metabolic water generation or sampling error.

280

281 **Conclusions**

282 In this study, a full-scale biocell type bio-drying MBT system was investigated. Mass
283 balance of each waste component was estimated by using composition analysis,
284 characterization of waste samples, and waste tonnage data. During separation of biodried
285 outputs, 62 % of plastics and 54 % of paper were recovered as raw material for RPF. The
286 ratios are not low because unrecovered plastics and paper were returned to next biocell
287 reactor. Due to low biodegradation of plastics and paper for 17-days of bio-drying, almost
288 100 % of the RPF materials could be recovered. Wood was decreased by reduction in particle
289 size during the bio-drying process, 90 % of wood was returned to next BR. Mixed fine seems
290 barely reduced, but reduced wood particle moved to mixed fine and filled up the loss caused
291 by organic degradation and 60 % of mixed fine was returned to next BR.

292 Water removal phenomena was simulated by the model of daily water removal by using
293 the operation data. The process was subjected to four distinct operation phases and was
294 operated according to the waste temperature. The main phase of water removal was expected
295 to be occurred in stabilization phase (III) for 11 days. However, half of water removal was
296 occurred at initial two phases and phase IV for only 6 days in total and this is caused by high
297 waste temperature for sanitization (phases I and II) and high airflow rate for cooling phase.
298 Phase III was long, but decreasing temperature resulted in low water evaporation.

299 The findings of this study which are separation efficiency of biodried outputs and effects of
300 temperature and airflow rate on drying efficiency, can give some contribution on improving
301 the full-scale bio-drying MBT system.

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364 **Figure captions:**

365 **Fig. 1** Description of the investigated bio-drying MBT system: (a) process flow and annual
366 mass in the facility and sampling location and (b) schematic diagram of airflow in the reactor

367 Fig. 1a - S mark, collected waste samples; TS, truck scale; FM, flow meter; LD, weighing
368 scale on the wheel loader; SUM, Mass sum of each waste stream

369 Fig. 1b – Symbols: V, airflow rate; T, temperature; D, degree of openness in damper; q, fan
370 speed in percentage unit

371 Subscript: F, fresh air; A, total aeration; S, solid waste; EX, exhaust air; RE, recirculating
372 air

373 Variables in box are given data by the bio-drying process monitoring

374 (These figures show the details of the investigated bio-drying MBT system.)

375 **Fig. 2** Physical composition of waste on a wet basis

376 Misc. comb.: Miscellaneous combustibles

377 (This figure shows the physical composition of the collected waste sample and referred MSW
378 and new wood)

379 **Fig. 3** Waste characteristics of (a) combustible content and (b) gasified carbon in biogas

380 No colored marks indicate ‘BR input’ samples which includes fresh MSW; Colored marks
381 indicate ‘BR output’ and other separated waste stream; Striped marks indicate referred
382 characteristics of MSW and new wood

383 DS: Dry solids

384 (These figures show the results of sample analysis for its combustible fraction and biological
385 degradable organics indicated by gasified carbon amount in biogas)

386 **Fig. 4** Mass fraction of (a) dry solids, (b) gasified carbon in biogas, and (c) combustibles of
387 each waste stream component on a dry basis, normalized by the total dry mass of the INPUT

388 Misc. comb.: Miscellaneous combustibles

389 (These figures show the weighted mass fraction of dry solids, gasified carbon in biogas and
390 combustibles by each composition in different waste stream on a dry basis. Mass fraction was
391 normalized by the total dry mass of the INPUT.)

392 **Fig. 5** Mass balances of (a) dry solids, (b) gasified carbon in biogas, and (c) combustibles in
393 the bio-drying process on a dry basis

394 (These figures show the mass balance of the bio-drying process in terms of the total dry
395 solids, gasified carbon in biogas and combustibles)

396 **Fig. 6** Profiles of (a) operation variables and (b) water removal rate under different operation
397 phase during the bio-drying process

398 I: Warm-up phase; II: Sanitization phase; III: Stabilization phase; IV: Cooling phase

399 T_s , waste temperature; V_A , flow rate of total aeration; V_F , flow rate of fresh air

400 Numbers in Fig. 6b indicate the water removal portion

401 (These figures show the profiles of operation variables and daily water removal rate under
402 different phase during the bio-drying process.)

403

Table 1. Separation ratio of ‘OUTPUT’ streams in dry basis

	RPF materials	Wood residue	Fine residue	
Total dry mass	0.39	0.42	0.19	
Material	Plastics	0.62	0.31	0.07
	Paper	0.54	0.30	0.16
	Textiles	0.68	0.25	0.06
	Wood	0.06	0.90	0.04
	Mixed fine	0.33	0.08	0.60

Fig. 1 Description of the investigated bio-drying MBT system: (a) process flow and annual mass in the facility and sampling location and (b) schematic diagram of airflow in the reactor

Fig. 1a - S mark, collected waste samples; TS, truck scale; FM, flow meter; LD, weighing scale on the wheel loader; SUM, Mass sum of each waste stream

Fig. 1b – Symbols: V, airflow rate; T, temperature; D, degree of openness in damper; q, fan speed in percentage unit

Subscript: F, fresh air; A, total aeration; S, solid waste; EX, exhaust air; RE, recirculating air

Variables in box are given data by the bio-drying process monitoring

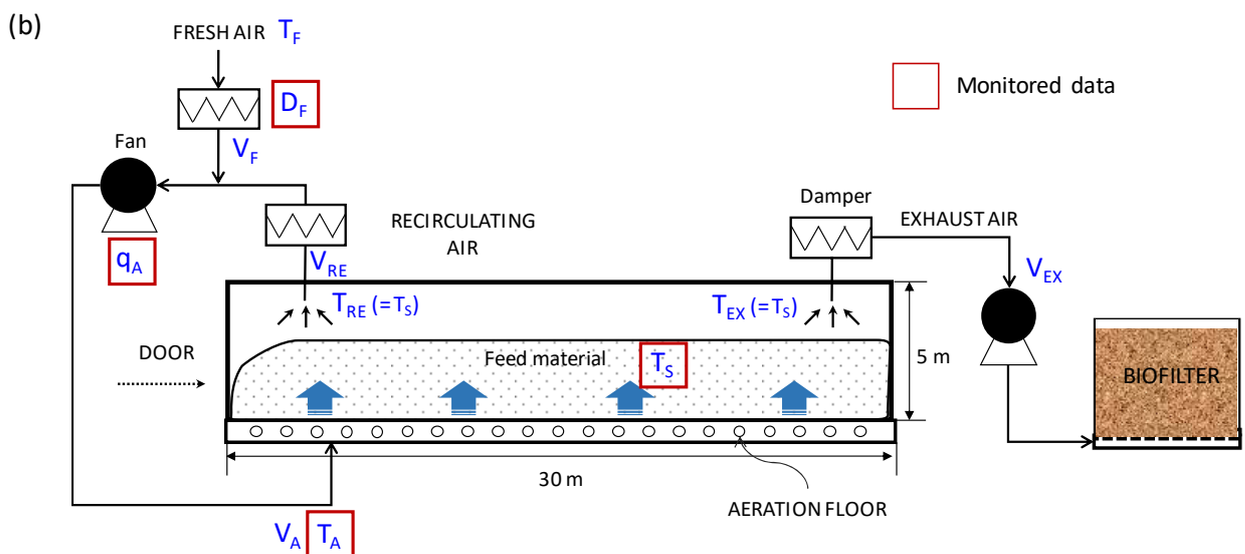
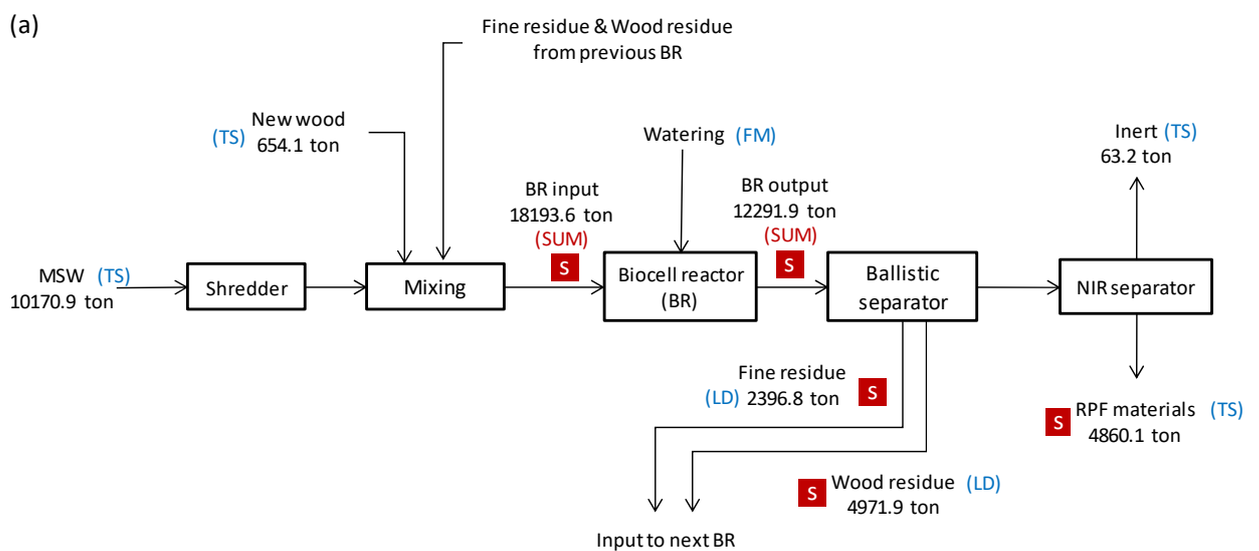


Fig. 2 Physical composition of waste on a wet basis

Misc. comb.: Miscellaneous combustibles

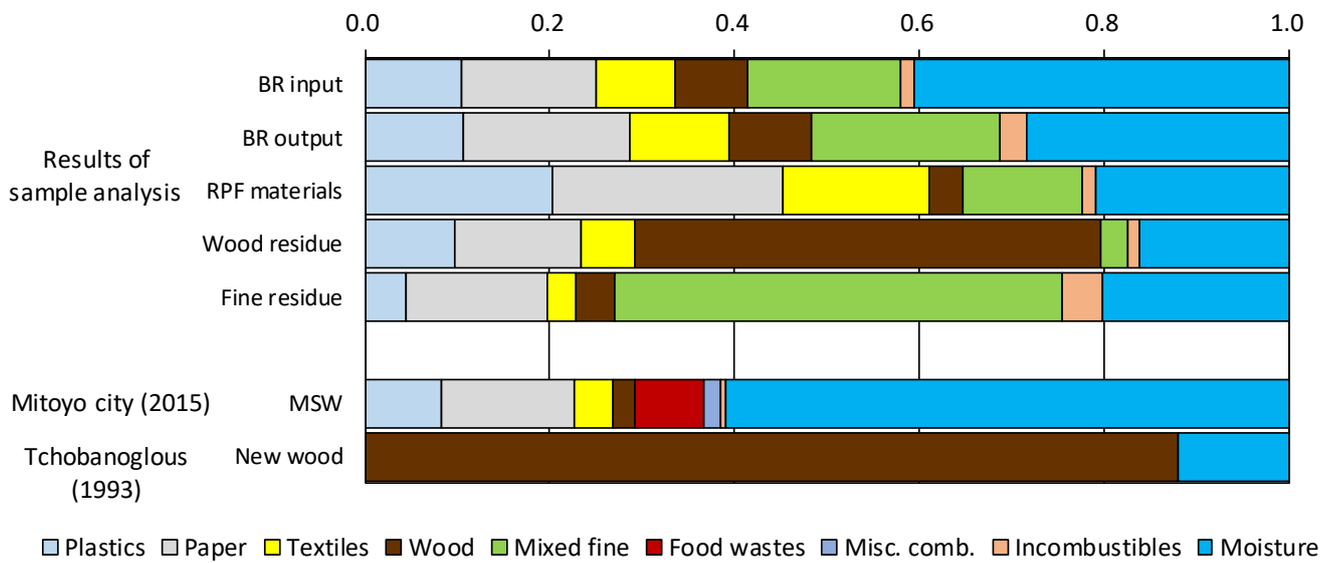


Fig. 3 Waste characteristics of (a) combustible content and (b) gasified carbon in biogas

No colored marks indicate 'BR input' samples which includes fresh MSW; Colored marks indicate 'BR output' and other separated waste stream; Striped marks indicate referred characteristics of MSW and new wood and new wood

DS: Dry solids

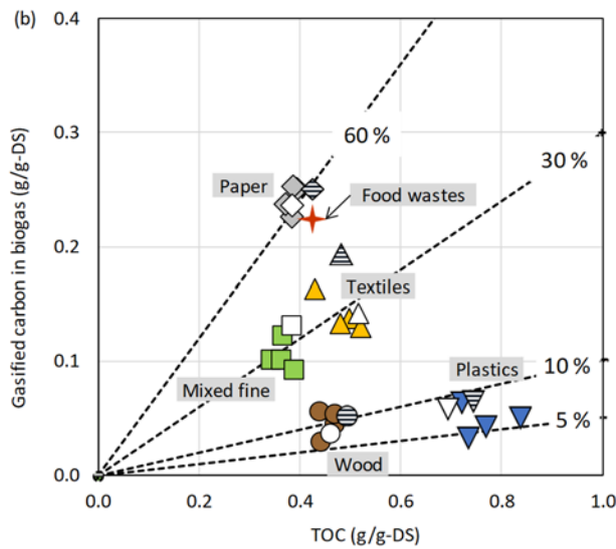
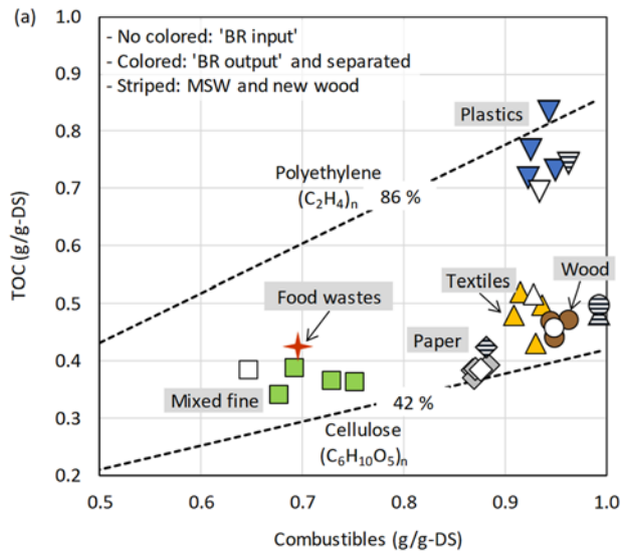


Fig. 4 Mass fraction of (a) dry solids, (b) gasified carbon in biogas, and (c) combustibles of each waste stream component on a dry basis, normalized by the total dry mass of the INPUT

Misc. comb.: Miscellaneous combustibles

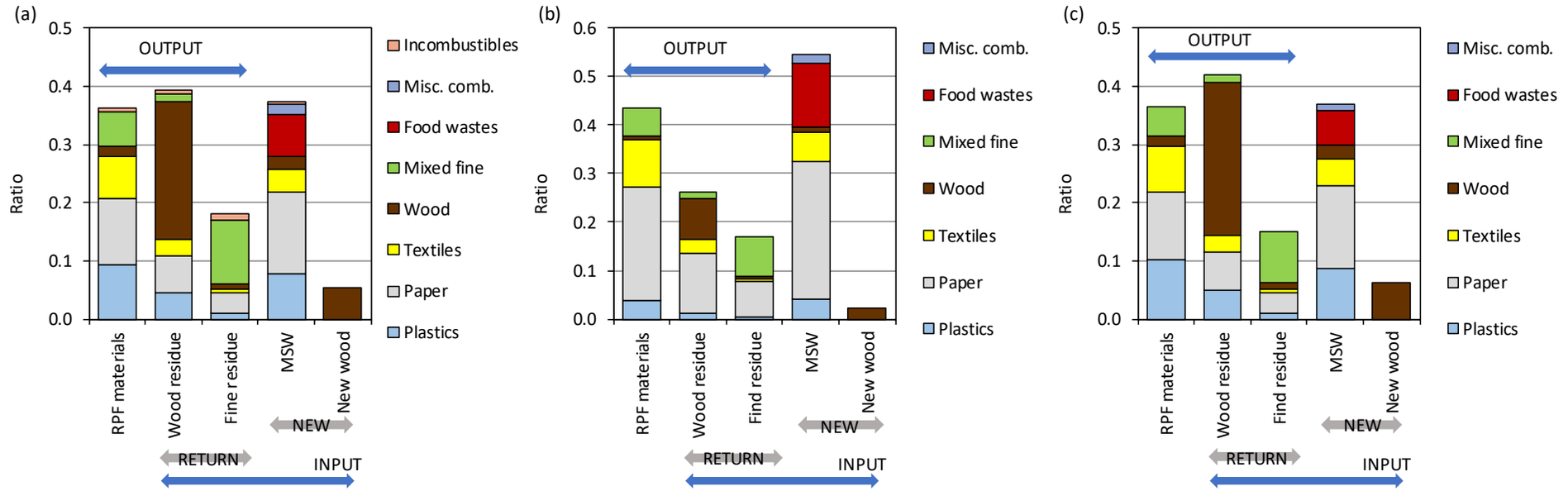


Fig. 5 Mass balances of (a) dry solids, (b) gasified carbon in biogas, and (c) combustibles in the bio-drying process on a dry basis

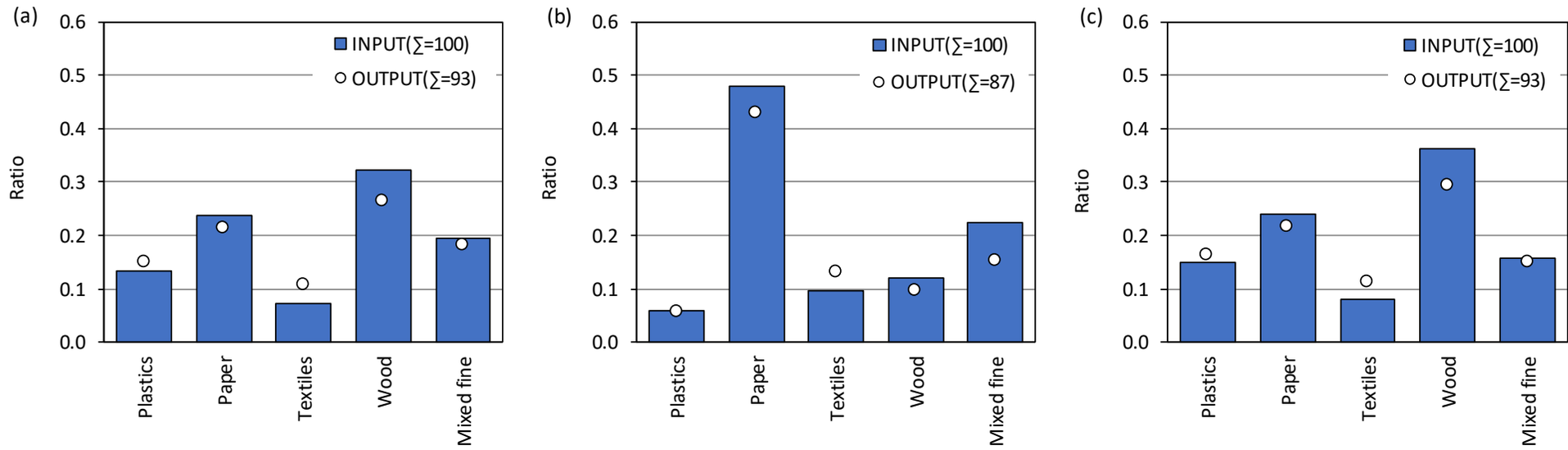


Fig. 6 Profiles of (a) operation variables and (b) daily water removal rate under different operation phase during the bio-drying process

I: Warm-up phase; II: Sanitization phase; III: Stabilization phase; IV: Cooling phase

T_s , waste temperature; V_A , flow rate of total aeration; V_F , flow rate of fresh air

Numbers in Fig. 6b indicate the water removal portion

