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1 **Tailor-made microbial consortium for Kombucha fermentation:**
2 **microbiota-induced biochemical changes and biofilm formation**

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11
12 Running title: Dynamic follow-up of Kombucha fermentation

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15 **ABSTRACT**

1
2 16 Kombucha is a very distinct naturally fermented sweetened tea that has been produced for
3
4 17 thousands of years. Fermentation relies on metabolic activities of the complex autochthonous
5
6 18 symbiotic microbiota embedded in a floating biofilm and used as a backstop for successive
7
8 19 fermentations. Here, we designed a tailor-made microbial consortium representative of the
9
10 20 core Kombucha microbiota to drive this fermentation. Microbial (counts, metagenetics),
11
12 21 physico-chemical (pH, density) and biochemical (organic acids, volatile compounds)
13
14 22 parameters were monitored as well as biofilm formation by confocal laser scanning microscopy
15
16 23 and scanning electron microscopy. While 9 species were co-inoculated, 4 (*Dekkera*
17
18 24 *bruxellensis*, *Hanseniaspora uvarum*, *Acetobacter okinawensis* and *Lacitilactobacillus nagelii*)
19
20 25 largely dominated. Microbial activities led to acetic, lactic, succinic and oxalic acids being
21
22 26 produced right from the start of fermentation while gluconic and glucuronic acids progressively
23
24 27 increased. A distinct shift in volatile profile was also observed with mainly aldehydes identified
25
26 28 early on, then high abundances of fatty acids, ketones and esters at the end. Correlation
27
28 29 analyses, combining metabolomic and microbial data also showed a shift in species
29
30 30 abundances during fermentation. We also determined distinct bacteria-yeast co-occurrence
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32 31 patterns in biofilms by microscopy. Our study provides clear evidence that a tailor-made
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34 32 consortium can be successfully used to drive Kombucha fermentations.
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41 33

43 34 Keywords: Kombucha, fermentation dynamics, metagenetics, biochemical changes, biofilm,
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45 35 confocal microscopy, SEM
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36 1. INTRODUCTION

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2 37 For centuries, humans have relied on fermentations to preserve foods and beverages but also
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4 38 to diversify products with specific organoleptic, nutritional and health properties. The microbial
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6 39 ecology of many well-known fermented products (i.e. wine, beer, cheese, dry cured meats,
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8 40 olives, sourdough...) has been widely studied by the scientific community to unravel the
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10 41 complex microbial communities involved in fermentation, determine co-occurrence patterns
11
12 42 and, in some cases, the roles of the key fermentation drivers (Bourdichon et al., 2012; Coton
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14 43 et al., 2017; Fremaux et al., submitted; Montel et al., 2014; Mounier et al., 2005; Penland et
15
16 44 al., 2020). For naturally fermented products, solely relying on autochthonous microorganisms
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18 45 for fermentation, these studies are of particular interest as it may lead to novel starter selection
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20 46 and fermentation drivers with specific and desirable traits for a given product.
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25 47 Kombucha is a naturally fermented effervescent beverage with a slightly acidic, refreshing and
26
27 48 distinct taste originating from Northeastern China, about 220 B.C (Jayabalan et al., 2014). It is
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29 49 globally distributed and has become increasingly popular in North America and Europe (Coton
30
31 50 et al., 2017) due to strong consumer demands for more natural and healthier products that
32
33 51 keep their signature characteristics. Kombucha is prepared using sweetened black or green
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35 52 teas and fermented by symbiotic cultures of autochthonous yeasts, acetic acid bacteria (AAB)
36
37 53 and lactic acid bacteria (LAB) embedded in a floating biofilm for, on average, 8 to 15 days
38
39 54 under aerobic and static conditions (Coton et al., 2017) although longer fermentations (~1
40
41 55 month) may occur. The microbially rich Kombucha biofilm is successively used as a starter, or
42
43 56 backslop, for future fermentations and often shared between producers worldwide. Microbial
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45 57 compositions of many Kombuchas has been recently studied using both metagenetics and
46
47 58 metagenomics sequencing approaches (Arikan et al., 2020; Chakravorty et al., 2016; Coton
48
49 59 et al., 2017; De Filippis et al., 2018; Gaggia et al., 2019; Marsh et al., 2014; Reva et al., 2015;
50
51 60 Villarreal-Soto et al., 2020) and, in some cases, linked to culture-dependent techniques (Coton
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53 61 et al., 2017). This is particularly of interest to determine species co-occurrence patterns but
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55 62 also to access, preserve and exploit the identified microbial diversity via culturing techniques.
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63 These studies, regardless of geographical origin, have all highlighted the widespread presence
64 of certain fungal and bacterial species interacting through highly symbiotic relationships and
65 cooperative metabolism. Among yeasts, *Dekkera* spp. (e.g. *D. anomala*, *D. bruxellensis*),
66 *Pichia* spp. (e.g. *P. occidentalis*), *Yarrowia* spp. (e.g. *Y. lipolytica*), *Candida* spp. (e.g. *C.*
67 *zemplinina*), *Saccharomycetales* spp., *Hanseniaspora* spp. (e.g. *H. valbyensis*) and
68 *Zygosaccharomyces* spp. (e.g. *Z. bailii*) were among the most dominant and prevalent species
69 (Arikan et al., 2020; Chakravorty et al., 2016; Chen & Liu, 2000; Coton et al., 2017; De Filippis
70 et al., 2018; Gaggia et al., 2019; Hesseltine, 1965; Jankovic & Stojanovic, 1994; Jayabalan et
71 al., 2014; Liu et al., 1996; S. L. Markov et al., 2001; Marsh et al., 2014; Mayser et al., 1995;
72 Reva et al., 2015; Teoh et al., 2004; Villarreal-Soto et al., 2020), although not all species are
73 systematically identified thus not necessarily part of the Kombucha core microbiota. Among
74 bacteria, *Proteobacteria* phylum is systematically observed with a strong prevalence of AAB.
75 The most frequent genera and species include *Komagataeibacter* (formerly
76 *Gluconacetobacter*) (e.g. *K. xylinus*, *K. europaeus*, *K. rhaeticus*), *Gluconobacter* (e.g. *G.*
77 *oxydans*) and sometimes *Acetobacter* (e.g. *A. tropicalis*, *A. okinawensis*) (Arikan et al., 2020;
78 Chakravorty et al., 2016; Coton et al., 2017; De Filippis et al., 2018; Gaggia et al., 2019;
79 Greenwalt et al., 2000; Jankovic & Stojanovic, 1994; Jayabalan et al., 2010; Kurtzman et al.,
80 2001; Liu et al., 1996; Marsh et al., 2014; Reva et al., 2015; Villarreal-Soto et al., 2020). Some
81 studies have also highlighted LAB, although less abundant than AAB with *Lacitilactobacillus*
82 spp. (e.g. *L. nagelii*), *Oenococcus oeni* and/or *Bifidobacterium* being identified (Coton et al.,
83 2017; Teoh et al., 2004). These microorganisms co-occur in the tea fraction and/or are
84 embedded in the floating cellulosic biofilm regenerated with each new fermentation, also
85 named “mother”, “tea fungus” or “SCOBY”, for “Symbiotic Community of Bacteria and Yeast”
86 (Jayabalan et al., 2014; May et al., 2019; Villarreal- Soto et al., 2018). The stability and
87 maintenance of the biofilm microbiome over time has yet to be characterized despite its intense
88 use as a backstop for fermentations. During fermentation, this microbially rich biofilm floats on
89 the surface of open fermentation tanks thus creating favorable aerobic conditions at the liquid-
90 air interface for microbial interactions and cooperative metabolism. Indeed, yeasts break down

91 the main carbon source, sucrose, initially added to the tea into glucose and fructose, and
92 ferment them into two major end products, ethanol and CO₂ (Coton et al., 2017; Jayabalan et
93 al., 2014; Villarreal- Soto et al., 2018). Then, bacteria, mainly AAB embedded in the biofilm,
94 successively oxidize ethanol into acetic acid but also use glucose to produce other organic
95 acids as acetic, succinic, gluconic or glucuronic acids (Arikan et al., 2020; Coton et al., 2017;
96 Gomes et al., 2018; Siniša L. Markov et al., 2003; Ramachandran et al., 2006). These
97 metabolites all contribute to the slightly acidic and sour taste of the final product but may also
98 contribute to the potential health benefits associated with this product although further studies
99 are needed. Some AAB species also play another key role during Kombucha fermentation as
100 they can produce the cellulosic biofilm from simple sugars (Coton et al., 2017; Villarreal- Soto
101 et al., 2018). LAB, although not systematically found, also produce organic acids and impact
102 product acidity and the overall sensorial attributes. Overall, bacterial metabolic activities not
103 only rapidly reduce pH to values close to 3, but also decrease the ethanol produced by
104 fermentative yeasts to values below 1.2%, which is in accordance with the designation “without
105 alcohol” from the European regulation (UE N° 1169/2011).

106 Changes in biochemical profile during fermentation are related to the Kombucha microbiota
107 and/or biofilm origins but also fermentation conditions, in particular vessel size, oxygen
108 availability, temperature and tea type (Cardoso et al., 2020; Coton et al., 2017; Gaggia et al.,
109 2019; Marsh et al., 2014). A full understanding of the role of this complex microbiota, whether
110 embedded or not in a biofilm, during fermentation is thus needed to better maintain and control
111 production conditions. In this sense, strain selection and associations for directed Kombucha
112 fermentations is of clear interest as recently documented (S. Wang et al., 2020).

113 The goal of our study was to i) design a tailor-made microbial consortium by selecting the main
114 microbial drivers from a previously described Kombucha core microbiota (Coton et al., 2017)
115 to drive this fermentation and ii) determine its impact on fermentation by dynamically
116 monitoring microbial populations (by numerations and metagenetics), physico-chemical
117 properties (pH and density), biochemical parameters (organic acids and volatile compounds),

118 and biofilm formation using fluorescent *in situ* hybridization with microbial group specific probes
119 coupled to confocal laser scanning microscopy and scanning electron microscopy analyses.

120

121 2. MATERIALS AND METHODS

122 2.1. Defined complex consortium for Kombucha fermentations

123 2.1.1. Consortium and culture conditions

124 The defined consortium included five ABB strains belonging to four species: *Acetobacter*
125 *tropicalis* J2-MRN2-BA1.1, *A. okinawensis* J2-BSN3-BA4.1 and J0-MRN1-BA3.1,
126 *Komagataeibacter hansenii* (formerly *Gluconacetobacter hansenii*) J0-MRC3-BA5.2 and
127 *Gluconobacter oxydans* J0-BSD1-BA3.1, two LAB species : *Lacitilactobacillus nagelii* J0-
128 BSV3-BL5 and *Oenococcus oeni* UBOCC-A-315005 and four yeast strains belonging to three
129 species: *Dekkera bruxellensis* J9-MRB2-Lev3.1 and J4-MRN1-Lev2.1, *Hanseniaspora uvarum*
130 J9-MRD1-Lev2.2 and *Zygosaccharomyces bailii* J9-BSB2-Lev5.3. All strains were previously
131 isolated from black or green tea Kombucha fermentations (Coton et al., 2017). Cultures were
132 directly prepared from cryo-conserved glycerol stocks at -80°C by plating on appropriate
133 media. Yeast extract glucose chloramphenicol agar (YGC, Biomérieux, France) was used for
134 yeast cultures (incubation at 25°C for 3-4 days) while AAB were plated on mannitol medium
135 (D-mannitol 25 g/L, yeast extract 5 g/L, universal peptone 3 g/L and agar 15 g/L) supplemented
136 with 0.1 g/L pimaricine and incubated at 30°C for 3-4 days and LAB on De Man Rosaga Sharpe
137 (MRS, Biomerieux, France) acidified with 10% (v/v) citric acid to pH 4.8 and supplemented
138 with 0.1 g/L pimaricine and incubated at 30°C for 3-4 days. Only MRS plates were incubated
139 under anaerobic conditions in closed jars.

140 To calibrate cultures (between 10^7 to 10^8 colony-forming units (CFU)/mL), one colony was
141 resuspended in 10 mL of broth and incubated for 72 h at 25°C (yeast) or 30°C (AAB, LAB).

142 Yeast strains were cultivated in tryptic soy broth (Biomerieux, France) supplemented with 2.5
143 g/L yeast extract while AAB strains were cultivated in mannitol broth (D-mannitol 25 g/L, yeast

144 extract 5 g/L, universal peptone 3 g/L) and LAB in MRS broth (Biomérieux, France). Then, a
145 second culture was performed using a 1% inoculum in 10 mL broth and incubated for 24 h
146 using the same conditions, except for *O. oeni* and *K. hansenii* that were incubated for 48 h.
147 Yeast and AAB cultures were agitated at 120 rpm at 25°C or 30°C, respectively, while LAB
148 were incubated at 30°C under static conditions. All pure cultures were enumerated before
149 inoculation to ensure target values of 10⁵ CFU/mL were obtained for fermentations.

2.1.2. Fermentation conditions

152 For fermentations, the consortium was then prepared by inoculating each strain at a final
153 concentration of 10⁵ CFU/mL in 400 mL sweetened green tea. This was prepared by infusing
154 dried tea leaves (placed in a paper filter) for 30 min in boiled distilled water supplemented with
155 55 g/L organic blond sugar and sterilized at 104°C for 30 min. After homogenization, 800 mL
156 lab scale fermentation vessels were covered with a sterilized linen sheet to create aerobic
157 conditions and incubated at 25°C for up to 27 days. Fermentation time was purposely extended
158 to 27 days for biofilm formation and microscopy observations. For the dynamic follow-up, 16
159 biological replicates were performed and sampling was done at days 7, 11, 14, 20 and 27. An
160 unfermented tea control sample was also included at day 0. For each date, 3 replicates were
161 analyzed and the biofilm was recovered separately. The only exception was at d0, as no biofilm
162 was yet formed.

2.2. Dynamic follow-up

2.2.1. Physico-chemical analyses

166 The indicators used to follow fermentation (pH, density and biofilm wet mass) were measured
167 at each sampling day using 3 biological replicates. pH was determined using an electronic pH
168 meter (Eutech Instruments, The Netherlands) and tea density was measured with a 1.000-
169 1.100 ± 0.001 g/mL range densitometer at 20°C (Fisher Scientific, France). Density

170 measurements were specifically performed as a simple means to monitor sugar consumption
171 during fermentation. Biofilm wet mass was measured using sterile conditions in Petri dishes
172 and samples were kept for further analyses.

173

174 2.2.2. Microbiota monitoring

175 *Microbial enumerations*

176 AAB, LAB and yeast enumerations were carried out on both biofilm and Kombucha tea
177 samples. For tea samples, serial dilutions were performed with tryptone salt (TS, Merck) while
178 for biofilms, 1 g was transferred into 50 mL tubes containing 9 mL TS and homogenized with
179 a sterile Ultra-Turrax^R (IKA, Germany) before being placed in a sterile stomacher bag equipped
180 with a filter membrane and mixed for 180 s using a Stomacher (AES, France). The liquid
181 fraction was collected and 1 mL was used for serial dilutions in TS while the remaining aliquot
182 was conserved for downstream analyses. For both tea and biofilm aliquots, dilutions were
183 plated on YGC for yeast, mannitol for AAB and MRS for LAB counts using an automatic spiral
184 plater method (Interscience, France). Petri dishes were incubated using the same conditions
185 as described above.

186

187 *Metabarcoding*

188 For total DNA extractions from tea samples, 15 mL were collected and centrifuged at 6654 g
189 for 5 min at 4°C. Supernatants were collected, filtered using 0.45 µm PTFE filters (Sartorius)
190 and kept at -20°C until further analyses by LC-MS while cells pellets were stored at -20°C for
191 DNA extractions. For biofilm samples, 4 mL of the homogenate prepared for microbial
192 enumerations were collected and centrifuged using the same conditions and cell pellets were
193 stored at -20°C for DNA extractions.

194 DNA extractions were performed using an optimized version of the protocol described by
1 Coton *et al.* (2017). This included an additional cell lysis step by adding 300 mg glass beads
2 195 (< 212 μm) to cells after incubation with the lysis buffer. Cells were then lysed using a vibro-
3
4 196 crusher (Grosseron, France) twice for 30 s at 30 Hz and kept on ice for 5 min between each
5
6 197 step before continuing the protocol as previously described. DNA extracts were quantified
7
8 198 using a Nanodrop 1000 Spectrophotometer (ThermoFischer Scientific, USA) and adjusted to
9
10 199 20 ng/ μL for downstream MiSeq PE300 sequencing at the Genome Quebec sequencing
11
12 200 platform (McGill University, Canada). PCRs were performed by targeting the V3-V4 region of
13
14 201 the bacterial 16S rRNA gene using the S-D-bact-0341-b-S-17 (5'-
15
16 202 CCTACGGGNGGCWGCAG-3') and S-D-BAct-0785-a-A-21 (5'-
17
18 203 GACTACHVGGGTATCTAATCC-3') primers (Klindworth *et al.*, 2013), and the 26S yeast rDNA
19
20 204 D1/D2 region using the NL1 5'-GCATATCAATAAGCGGAGGAAAAG-3' and NL4 5'-
21
22 205 GGTCCGTGTTTCAAGACGG-3' primers (O'Donnell, 1993). This technology generated 2x300
23
24 206 bp reads and a total of 3.3 Gb of data for both amplicon types derived from the 31 samples.
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31 *Bioinformatics and data analyses*

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33
34 209 The DADA2 library (Callahan *et al.*, 2016) was used in R version 3.5.0 (R Core Team, 2019)
35
36 210 for 16S and 26S rRNA gene reads filtering. For 16S and 26S analyzes, 2 698 218 reads and
37
38 211 2 650 330 reads respectively were conserved with a normalization on the smallest number of
39
40 212 reads found in a sample, 15 301 and 64 750 respectively. For 16S rRNA gene reads, forward
41
42 213 and reverse read pairs were trimmed and filtered, with forward reads truncated at 270 bp and
43
44 214 reverse reads at 210 bp, no ambiguous bases allowed and each read required to have less
45
46 215 than two expected errors based on their quality scores. Amplicon sequence variants (ASVs)
47
48 216 were independently inferred from the forward and reverse reads of each sample using the run-
49
50 217 specific error rates, and then read pairs were merged requiring at least 15 bp overlap. For 26S
51
52 218 rDNA reads, only forward reads were trimmed and filtered, with truncation at 300 bp, no
53
54 219 ambiguous bases allowed and each read required to have less than two expected errors
55
56 220 based on their quality scores. ASVs were directly inferred from forward reads.
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221 16S chimera sequences were removed using UCHIME algorithm (Edgar et al., 2011)
1
2 222 implemented in VSEARCH v1.1.3 (<https://github.com/torognes/vsearch>) against the
3
4 223 ChimeraSlayer reference database (Haas et al., 2011) and the RDP classifier (Q. Wang et al.,
5
6 224 2007) was used for taxonomy assignment, which was made with GreenGenes v13.8 database
7
8 225 (McDonald et al., 2012) available in Qiime (Caporaso et al., 2010). Each 16S ASV was then
9
10
11 226 classified to the species level using the RDP seqmatch tool
12
13 227 (<https://rdp.cme.msu.edu/seqmatch>). 26S rRNA gene ASVs were assigned to the species level
14
15 228 using the BLAST algorithm (<http://www.ncbi.nlm.nih.gov/BLAST/>).
16
17
18 229 Beta diversity analyses based on Bray-Curtis distances were performed using the Calypso
19
20 230 software tool v8.84 (Zakrzewski et al., 2017) after total sum normalization of count data
21
22
23 231 combined with square root transformation (Hellinger transformation).
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26 232

28 233 *2.2.3. Biofilm formation*

31 234 *Fluorescent in situ hybridization and confocal laser scanning microscopy*

34 235 To observe biofilm formation, yeast and bacterial species co-occurrence patterns within the
35
36 236 biofilm, fluorescent *in situ* hybridization (FISH) with different fluorochromes was used to
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39 237 specifically monitor the main microbial groups by confocal laser scanning microscopy (CLSM)
40
41 238 at the microscopy platform "Plateforme d'Imagerie et de Mesures en Microscopie (PIMM)",
42
43 239 University of Bretagne Occidentale. Probes to specifically target yeasts (EUK516) and bacteria
44
45 240 (EUB338) but also AAB (ALF1B) and LAB (LGC and Ooeni) were synthesized with different
46
47
48 241 fluorochromes (Table S1). Preliminary FISH tests were performed on the eleven strains of the
49
50 242 defined consortium using EUB338, EUK516, ALF1B and LGC probes to confirm cell fixation,
51
52 243 probe hybridization and specificity.
53
54

55 244 For biofilm observations by CLSM, cell fixation was performed with protocol described by Roller
56
57 245 *et al.* (1994) with an approximately 15 mm² sample was cut and transferred into a tube with
58
59
60 246 500 µL PBS (PBS; 130 mM NaCl, 10 mM Na₂HPO₄/NaH₂PO₄, pH 7.2-7.4) and 500 µL ice-cold
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247 96% ethanol then conserved at -20°C until hybridization. For cell hybridization, protocol
1 adapted from Manz *et al.*, (1992) was applied. To do so, samples were transferred into 8-wells
2 248
3
4 249 microscopy slides (Ibidi, Germany) and dried and dehydrated by directly adding and removing
5
6 250 300 µL 50%, 80% and 96% ethanol in the well. 100 µL of hybridization buffer (20 % formamide)
7
8 251 was deposited to 100 µL to completely immerse the sample and 12 µL of probe solution (25
9
10 252 ng/µL) were added, then the slides were incubated at 46°C for 4 h (humidity equilibrated).
11
12 253 Afterwards, each slide was washed by immersion in a washing buffer (freshly prepared using
13
14 254 0.225 M NaCl, 20 mM Tris-HCl and 0.01% SDS pre-heated to 48°C) for 10 min at 48°C in a
15
16 255 water bath. Finally, slides were very briefly washed with ice-cold distilled water and dried with
17
18 256 compressed air. One drop of anti-fade solution (ProLong Diamond Antifade Mountant, Thermo
19
20 257 Fisher Scientific, USA) was deposited on the surface of each sample and the slides were kept
21
22 258 at -20°C in darkness until microscopy observations. Labeled biofilm samples were analyzed
23
24 259 by using a confocal microscope ZEISS LSM780 using the different probes at the wavelengths
25
26 260 indicated in Table S1. Three biological replicates were treated per sample and at least two
27
28 261 were analyzed.
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37 263 *Scanning Electron Microscopy*

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40 264 As for CLSM, biofilms were transferred into tubes containing 1 mL of fixing solution (50 mL of
41
42 265 cacodylate buffer pH 5 (cacodylate solution 0.4 M adjusted to pH 5 with 0.2 M hydrochloric
43
44 266 acid), 10 mL of glutaraldehyde 25% and 40 mL of sterile distilled water) for 30 min. Samples
45
46 267 were washed twice with 2-fold diluted cacodylate buffer and conserved overnight. Dehydration
47
48 268 was performed by successive baths in 50%, 70%, 90% and 100% ethanol for 3 min each, with
49
50 269 drying in between. Finally, each sample was placed on a glass slide and incubated at 35°C
51
52 270 until metallization by a gold sputter-coater and Scanning Electron Microscopy (SEM; HITACHI
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54 271 S-3200N).
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273 2.2.4. Metabolome

274 *Organic acids quantification by liquid chromatography-quadrupole-time-of-flight mass*
275 *spectrometry (LC-Q-TOF) and enzymatic kits*

276 Filtered tea samples were transferred into amber vials then 2 μ L were injected into a HPLC
277 1260 coupled to a 6530 Accurate-Mass quadrupole-time-of-flight mass spectrometry Q-TOF
278 (Agilent Technologies, Santa Clara, CA) equipped with a dual electrospray ionization source
279 (Agilent Technologies, Santa Clara, CA) based on the previously described protocol (Ibáñez
280 & Bauer, 2014). Molecules were separated using a Phenomenex (Torrance, CA) Rezex ROA-
281 Organic Acid H+ (8%) (300 x 7.8 mm) column equipped with a Rezex ROA-Organic Acid H+
282 (8%) (50 x 7.8 mm) guard column and analytes were ionized in negative electrospray ionization
283 (ESI-) mode in a scan range of 50 to 1700 m/z and 2 scan/s. The column was maintained at
284 55°C with an isocratic flow rate of 0.3 mL/min of water containing 0.1% formic acid (LC/MS
285 grade Carlo Erba Reagents, France) while samples were maintained at 10°C in a well plate
286 autosampler until injection. Run time was 15 min followed by 5 min post-time to wash and re-
287 equilibrate the column before the next injection. The mass spectrometer conditions were as
288 follows, capillary voltage 4000 V, source temperature 325°C, nebulizer pressure, 50 psig,
289 drying gas, 1 L/min.

290 Stock solutions of five organic acids (Table S2) were prepared in water supplemented with
291 0.1% formic acid and a 10-point linear range was prepared at a concentration between 0.01
292 and 100 μ g/mL. Standard solutions were first injected separately (three injections) to confirm
293 accurate identifications with the theoretical mass-to-charge (m/z) values (Table S2), then in a
294 mixture to validate peak separation and retention times and determine limits of detection and
295 quantification. MassHunter Quantitative Analysis software version B.07.01 (Agilent
296 Technologies) was used for compound identification and quantification using the ions listed in
297 Table S2.

298 For acetic and lactic acids, concentrations were determined using the Acetic acid, UV method
1
2 299 and Enzytec Liquid D/L-lactic acid enzymatic kits (r-biopharm, Germany) according to
3
4 300 manufacturer's instructions.
5

7 301 *Ethanol quantification*

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10 302 Ethanol quantification was performed by gas chromatography using a GC 3900 (Varian
11
12 303 Analytical Instruments, USA) equipped with a CP Sil 8CB LB/MS #CP8752 column (30 m x
13
14 304 0.32 mm, FT 0.25 µm, Chrompack Capillary Column, Varian, USA), an FID detector and an
15
16 305 EFC detector as described by Coton *et al.*, 2017. Results were expressed in g/L.
17
18

20 306 *Volatile compound profiles by gas chromatography-mass spectrometry*

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22 307 Each sampling day, 2.5 ± 0.02 g of Kombucha tea samples were transferred into 22-mL Perkin
23
24 308 Elmer vials, tightly sealed and kept at -20°C until analysis. Volatiles were extracted and
25
26 309 analysed by headspace (HS) trap extraction coupled to gas chromatography-mass
27
28 310 spectrometry (GC-MS) using a Perkin Elmer Turbomatrix HS-40 trap automatic headspace
29
30 311 sampler with trap enrichment and a Clarus 680 gas chromatograph coupled to a Clarus 600 T
31
32 312 quadrupole mass spectrometer (PerkinElmer, France), as previously described (Pogačić *et*
33
34 313 *al.*, 2015), with modifications of the chromatographic conditions according to Penland *et al.*,
35
36 314 (2020). The samples were injected in a random order, with standards and blank samples
37
38 315 (boiled deionized water) to monitor possible carryover and MS drift. Volatile compounds were
39
40 316 identified by comparing their retention index and mass spectra with those from the NIST 2008
41
42 317 Mass Spectral Library (Scientific Instrument Services, Ringoes, NJ, United States) and, when
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44 318 possible, with those of authentic standards (Sigma Aldrich, France) analyzed in the same
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46 319 conditions.
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52 320 *Statistical and correlation analyses*

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55 321 Statistical analyses were performed using the R software (R Core Team, 2019).
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2 322 Analyses of variance (ANOVA) and comparing means were used to determine whether the
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4 323 physico-chemical, microbial and biochemical data significantly changed during the time-course
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7 324 of fermentation.

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9 325 Data from GC-MS were centered and scaled by compound and hierarchically clustered by
10 326 Ward's minimum variance method and Euclidean distance metric with the hclust R function
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12 327 before being plotted by the heatmap.2 function of R gplots package. Principal component
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14 328 analysis (PCA) was first used to identify common biochemical profile changes over
15
16 329 fermentation times using tea and biofilm data. Then, a multiple factor analysis (MFA) was
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18 330 performed using the MFA function of the FactoMineR package (Lê et al., 2008). The data set
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20 331 was constituted of three groups of active variables: pH and density, concentrations of eight
21
22 332 organic acids and log values of abundance of 39 volatile compounds from GC-MS analysis
23
24 333 plus ethanol from HPLC, for the triplicate samples from d7 to d27. The age of samples and the
25
26 334 25 variables from both culture-dependent and -independent microbial analysis of tea and
27
28 335 biofilm were used as supplementary (i.e. illustrative) variables. All analyses were performed
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30 336 with R software using FactoMiner, Factoextra, Hmisc, Psych and gplots packages (Lê et al.,
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32 337 2008; Wickham, 2016).
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40 339 **3. RESULTS**

42 340 **3.1. *Physico-chemical analyses***

43
44 341 pH quickly decreased from an initial value of 5.57 to 3.40 ± 0.04 during the first seven days
45
46 342 then progressively reached 3.02 ± 0.05 by day 27 (Figure 1). Density values also decreased
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48 343 quickly during the first 11 days from 1018 to 1010 ± 0.58 then stabilized between 1010 and
49
50 344 1008 until the end of the fermentation (Figure 1) as sugars were consumed by the consortium.

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54 345 Biofilm formation was also monitored over time and a relatively thin biofilm quickly appeared
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56 346 on the tea surface (first 7 days), with a wet mass of 2.27 ± 0.15 g. It progressively darkened,
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58 347 strengthened and thickened to reach a wet mass of 5.36 ± 1.12 g by d27. The biofilm mass
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348 increased with a rate calculated at 0.32 g/day during the first 7 days, then 0.15 g/day up to d27
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2 349 (Figure 1).

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6 7 8 351 **3.2. Microbial populations monitored by culture-dependent analyses**

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11 352 In tea samples, microbial counts significantly increased by nearly 2 log₁₀ CFU/mL during the
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13 353 first 7 days, from 5.82 to 7.35 ± 0.05 log₁₀ CFU/mL for yeast populations, 6.30 to 8.74 ± 0.11
14
15 354 log₁₀ CFU/mL for AAB and from 6.76 to 7.64 ± 0.08 log₁₀ CFU/mL for LAB (p value ≤ 0.01)
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17 355 (Figure 2). In some cases, slightly higher concentrations than those expected for AAB and LAB
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19 356 can be explained by overestimations using the OD600nm-CFU/mL calibration curves. Yeasts
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21 357 and AAB populations were significantly higher at 7 days compared to the following days, then
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23 358 progressively decreased up to 27 days (6.93 ± 0.06 log₁₀ CFU/mL and 7.39 ± 0.05 log₁₀
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25 359 CFU/mL respectively), with an increasing from 11 to 14 days (p value = 0.06 and 0.01
26
27 360 respectively), while the LAB populations remained stable from 7 to 20 then significantly
28
29 361 decrease up to 27 days (7.40 ± 0.14 log₁₀ CFU/mL, p value = 0.02). Biofilm populations were
30
31 362 significantly higher than those in tea for both bacteria and yeasts. Yeast and LAB populations
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33 363 remained relatively stable during the fermentation at nearly 8.7 log₁₀ CFU/g and 7.5 log₁₀
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35 364 CFU/g respectively. On the other hand, AAB populations were highly active, significantly
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37 365 increased from 7 to 14 days (7.5 ± 0.15 log₁₀ CFU/g to 9.9 ± 0.60 log₁₀ CFU/g, p value =
38
39 366 0.003) then significantly decreased to 8.8 ± 0.07 log₁₀ CFU/g at 27 days (p value = 0.04)
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41 367 (Figure 2). Moreover, no visible contamination (e.g. molds) was observed during all
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43 368 fermentations.
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50 51 52 370 **3.3. Dynamic changes in bacterial and fungal consortium species determined by** 53 54 371 **16S rRNA and 26S metabarcoding analyses**

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57 372 Metabarcoding analyses were performed to monitor the relative abundance of bacterial (Figure
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59 373 3A) and yeast (Figure 3B) species in tea and biofilm samples throughout fermentation.
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374 Concerning bacterial community structure, *A. okinawensis* and *L. nagelii* were dominant in tea
1 samples throughout the fermentation (41 to 56% and 39 to 57 % of the abundance
2 375 respectively) while other AAB and *O. oeni* were at lower relative abundances. In the biofilm, *A.*
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3.4. ***Biofilm formation by the complex consortium monitored during fermentation***

396 Preliminary tests using FISH confirmed that the selected protocol did not require any enzymatic
397 permeabilization (lysozyme) step on the fixed cells, using either pure cultures or biofilm
398 samples, and probe specificities (Figures S2A, B, C). Yeasts and bacteria were differentiated
399 using simultaneous hybridizations with two different fluorochromes. Interestingly, the biofilm

400 negative control (without probes) showed natural fluorescence in CLSM, probably due to its
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2 401 cellulose composition.
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5 402 Dynamic monitoring of biofilm formation by the tailor-made consortium highlighted distinct
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7 403 bacteria and yeast cell clustering patterns as the structure strengthened and thickened
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9 404 (Figures 4A to L). There was a high abundance of bacterial cell clusters, certainly linked to the
10
11 405 highly active AAB populations in biofilm samples, in comparison to yeasts, although no specific
12
13 406 biofilm structure was observed. Microscopy observations at different sampling times showed
14
15 407 yeast cells frequently entrapped and/or completely covered by bacterial cells which can likely
16
17 408 explain the difficulty to observe them in some acquisitions. In Figure 4E, d7 biofilm showed
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19 409 that yeasts were mainly grouped on the outer surface of the biofilm, although impossible to
20
21 410 determine whether this section was in direct contact with the air or tea (due to orientation loss
22
23 411 during treatment). After 11 days fermentation, yeast and bacterial cells were more organized
24
25 412 and a bacterial mat was observed covering yeast cell clusters (Figure 4F), this same
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27 413 observation was made on all 3 biological replicates. This finding reinforced the idea that
28
29 414 symbiotic relationship and cooperative metabolism between these microbial populations may
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31 415 exist in the biofilm during fermentation. Obviously, biofilm density depended on the observed
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33 416 zone and depth as represented in Figure 4G and 4H. At d14, this phenomenon was
34
35 417 accentuated, and bacteria and yeasts were very abundant (Figures 4I and 4J). Then, a major
36
37 418 increase in biofilm, concomitant with a progressive decrease in bacteria and yeast clusters,
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39 419 was observed at days 20 and 27 (Figures 4K and 4L). Specific probes targeting AAB and LAB
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41 420 (with a specific probe for *O. oeni* due to the lack of fluorescence with the LGC probe, which
42
43 421 can be explained by two mismatches on 16S rRNA gene target) were also used on these
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45 422 samples but no further differences were found (Figure S3). Finally, LAB cells were certainly
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47 423 present but at much lower levels or scattered, so less accurately observed (Figure S3).
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54 424 SEM observations were consistent with CLSM observations (Figure 5). Indeed, yeast cells
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56 425 were grouped in clusters (based on cell shapes, consortium species could even be
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58 426 distinguished) (Figures 5A, 5E, 5I and 5M) and progressively covered by increasingly abundant
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427 bacterial cells during the fermentation (Figures 5A, 5B, 5C, 5E, 5F, 5G, 5I, 5J, 5K, 5M and 5N).
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2 428 Biofilm matrix also densified, clearly embedding bacterial and yeast cell clusters within its
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4 429 structure by d11 to the end. Also, between d20 and d27, microorganisms were observed to be
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6 430 in curly shaped fibrils that could be distinctly seen throughout the biofilm samples (Figures 5Q
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8 431 and 5R). Interestingly, several pores were also observed on the surface of the biofilm at d11
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11 432 with fibrous extensions (Figure 5B and 5C) appearing from and around bacterial cells and
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13 433 embedding the yeast cells thus creating a strengthening network from the bacterial cellulose
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15 434 being produced over time (Figures 5D, 5H, 5L and 5P). Finally, on some SEM acquisitions,
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17 435 biofilm production appeared to be linked to yeast cells (Figure 5O), although it could be an
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19 436 artefact of cellulose produced by AAB.
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26 438 **3.5. Changes in organic acids and ethanol content during fermentation**

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28 439 Seven organic acids were quantified in Kombucha tea samples during fermentation with very
29
30 440 significant variations in concentrations for succinic, oxalic, malic, glucuronic and gluconic acids
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32 441 (p value < 0.001). At the start, only three acids were quantified while by d7 all organic acids
33
34 442 were present except glucuronic acid that was only quantified from d11 onwards (Figure 6A).
35
36 443 Concentrations of gluconic, glucuronic, malic, oxalic and succinic acids acids then significantly
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38 444 increased until d20 before stabilizing to concentrations of 0.027 ± 0.006 g/L, 0.021 ± 0.004 g/L,
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40 445 0.003 ± 0.001 g/L, 0.020 ± 0.005 g/L and 0.015 ± 0.004 g/L, respectively (Figure 6A). As
41
42 446 expected, acetic and lactic acids were observed at 10 to 1000-fold higher concentrations than
43
44 447 the other acids (Figures 6B). Acetic acid was quantified at variable concentrations between d7
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46 448 (1.31 ± 0.04 g/L) and d20 (1.45 ± 0.07 g/L) (p value = 0.0906), then increased to 2.41 ± 0.06
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48 449 g/L at d27 (Figure 6B). Lactic acid was quantified at 318 ± 42 μ g/mL at d7 and significantly
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50 450 increased to reach 0.82 ± 0.02 g/L at d20 before decreasing to a final concentration of $0.45 \pm$
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52 451 0.02 g/L (Figure 6B). Finally, ethanol was mainly produced between d0 and d11 and showed
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54 452 a significant increase from 0.237 g/L (± 0.010) (d7) to 0.631 g/L (± 0.126). Concentrations then
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453 progressively decreasing to the end of fermentation to each 0.237 g/L (\pm 0.066). Ethanol
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2 454 content was systematically below 0.789 g/L during fermentation (Figure 6B).
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6 7 8 456 **3.6. Changes in the volatilome during fermentation**

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11 457 A total of 39 volatile compounds were identified and belonged to eight main families: alcohols
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13 458 (n=6), aldehydes (n=6), carboxylic acids (n=4), esters (n=11), fatty acids (n=6), furans (n=1),
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15 459 ketones (n=4) and styrene (Table S3). The heat-map drawn from hierarchical clustering
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17 460 analysis (Figure 7) as well the principal component analysis (Figure S4) show the main
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19 461 changes in volatile compound abundances over time with good repeatability, as the three
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21 462 biological tea replicates were systematically grouped, and samples were distinguished at the
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23 463 six time points with specific compounds being produced or most abundant. Moreover, the heat-
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25 464 map highlighted four clusters (A, B, C, D on Figure 7) also observed on the PCA. Indeed,
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27 465 cluster B (n=4) contained three aldehydes and butan-2-ol that tended to show high
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29 466 abundances at d0 or that did not show significant changes over time, then cluster A (n=8)
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31 467 corresponded to volatiles that were most abundant at d7, d11 and/or d14 before decreasing,
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33 468 such as nonanal and dodecanal. The volatilome complexified after d11 and d14 which
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35 469 corresponds to the standard fermentation time for this product and this is represented as
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37 470 cluster D (n=19) on the heat-map. Different alcohols (e.g. 2-methylpropan-1-ol, 3-methylbutan-
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39 471 1-ol) were produced as well as some fatty acids (e.g. octanoic and hexanoic acids) and ethyl
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41 472 esters (e.g. ethyl heptanoate, ethyl-2-methylbutanoate). In the last fermentation stages, as
42
43 473 highlighted in cluster C (n=8), some volatile compounds became increasingly abundant
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45 474 between d20 and d27, such as different acids (e.g. acetic, 4-hydroxybutanoic acids), ketones
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47 475 (e. g. hexan-2-one and heptan-2-one) and some specific esters (3-methylbutyl acetate, ethyl
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49 476 propanoate).
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57 58 59 478 **3.7. Microbiota and biochemical profile correlation during fermentation**

479 A multiple factor analysis (MFA) was performed to have a global view of the changes occurring
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2 480 Kombucha fermentation and to illustrate the correlations between microbial and biochemical
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4 481 variables correlate changes in the microbial species abundances in the tailor-made consortium
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6 482 or changes in population counts with the determined biochemical profiles of tea samples. A
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8 483 total of 74 physicochemical (pH, density, organic acids, volatiles) and microbial (counts and
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10 484 metabarcoding) parameters were included, which described both the tea and biofilm samples
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12 485 from d7 to d27. Dimensions 1 and 2 respectively explained 49.7% and 18.5% of the total
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14 486 variance and separated samples according to time in three clusters: d7, d11-d14, and d20-d27
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16 487 samples, (Figure 8). The beginning of the fermentation (d7 samples) was characterized by
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18 488 high density and pH values, high concentration of several aldehydes, correlated to the greatest
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20 489 AAB counts in tea and the highest abundances of three of the nine species included in the
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22 490 tailor-made consortium: the yeast *H. uvarum* both in tea and biofilm, the LAB *O. oeni* in tea
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24 491 and the AAB *Z. bailii* in biofilm. Days 11 and 14 were correlated to AAB counts in biofilm and
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26 492 numerous compounds including ethanol, many ethyl esters and diacetyl (2,3-butanedione).
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28 493 The final fermentation stage, between d20 and d27, was associated with higher abundances
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30 494 of glucuronic, gluconic and malic acids and some volatiles, including branched-chain acids and
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32 495 esters (e.g. 2-methylpropanoic acid, 3-methylbutanoic acid, 3-methylbutyl acetate and ethyl
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34 496 propanoate) and methyl ketones and secondary alcohols (e.g. hexan-2-one and hexan-2-ol).
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36 497 These changes were correlated to a high biofilm mass, high abundancies of *D. bruxellensis* in
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38 498 both tea and biofilm and of *A. tropicalis* in biofilm.
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48 500 **DISCUSSION**

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50 501 Kombucha is an ancient naturally fermented beverage that has become increasingly popular
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52 502 in the Western world. Many recent studies have described the microbial communities
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54 503 encountered in the fermented tea and/or biofilm used as backslop using culture-dependent
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56 504 and -independent approaches. The main metabolites produced as well as the potential
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58 505 antimicrobial properties or health benefits of the fermented drink have also been documented.
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506 However, the Kombucha microbiome and related properties depend on several parameters.

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2 507 The biofilm backslop used for successive spontaneous fermentations is portioned and often
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4 508 shared among producers. Its microbial stability and maintenance have yet to be fully
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6 509 investigated and changes in microbial composition over time are likely to occur. Fermentation
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8 510 can also vary due to differences in tea type, production conditions (tank size, oxygen
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10 511 availability, temperature...) and, to a lesser extent, geographical origin. To date, Kombucha
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12 512 fermentations largely rely on the autochthonous microbiota embedded within the biofilm, with
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14 513 new layers generated during each fermentation, or the in-house microbiota coming from the
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16 514 production environment or equipment. During successive fermentations, batch-to-batch
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18 515 variations can be therefore observed and are typically linked to changes in the Kombucha
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20 516 microbiome and abundances of the main fermentation drivers. In this context, selection of the
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22 517 key microbial drivers from the Kombucha core microbiota is of clear interest to drive and better
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24 518 control this fermentation. The main objectives of this study were to design a tailor-made
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26 519 complex consortium including species belonging to this core microbiota, as recently described
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28 520 by Coton et al (2017), and evaluate its impact on fermentation by monitoring multiple physico-
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30 521 chemical, biochemical, microbial and biofilm parameters. The tailor-made consortium included
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32 522 nine yeast, AAB and LAB species considered as the main drivers, including two involved in
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34 523 biofilm formation (*K. hansenii* and *G. oxydans*).

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40 524 Kombucha fermentations typically take up to 15 days and our data were consistent with this
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42 525 timeframe as both density and pH reached target values during the first 14 days (density ~1010
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44 526 and pH ~3.2). However, we extended fermentation time to 27 days to monitor biofilm formation
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46 527 as it was solely produced by the tailor-made consortium. Biofilms can be considered to be
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48 528 complex associations of one or more species interconnected cells embedded within a self-
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50 529 produced matrix and formed at either solid-liquid or liquid-air interfaces (Alexandre, 2013;
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52 530 Costerton et al., 1995; Hall-Stoodley et al., 2004; Kolter & Greenberg, 2006). A thin biofilm was
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54 531 rapidly observed at the liquid-air interface and covered the entire tea surface within 7 days,
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56 532 then progressively thickened and darkened to the end of fermentation.
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533 All microbial groups developed well in tea samples (up to +2 log increase in the first 7 days),
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2 534 as shown by microbial numerations, and these changes were linked to the rapid decrease in
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4 535 pH due to organic acid production, in particular acetic and lactic acids, by the consortium.
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6 536 These findings are in accordance with previous culture-dependent data including the same
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8 537 species described by Coton et al 2017. AAB populations were particularly active in the floating
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10 538 biofilm as the highest population increase was observed for this group. This can be directly
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12 539 linked to their obligately aerobic metabolism and the oxidative transformation of ethanol into
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14 540 acetic acid. Some AAB species as *Acetobacter pasteurianus* can also oxidize lactic acid
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16 541 released by LAB into acetic acid and acetoin (Moens et al., 2014). AAB species involved in
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18 542 biofilm formation are also most likely using this strategy to increase their access to oxygen as
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20 543 recently described by May *et al.* (2019).
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25 544 Similar results were also observed with microscopy observations as the microbiota was
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27 545 progressively embedded in the newly formed biofilm structure, especially AAB and yeasts. To
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29 546 the best of our knowledge, this is actually the first time that Kombucha biofilm formation is
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31 547 dynamically followed and observed using FISH probes coupled to CLSM and SEM. The only
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33 548 studies using CLSM on SCOBY were performed with non-specific probes as calcofluor or
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35 549 thiazine (Podolich et al., 2017; Reva et al., 2015). Here, we used yeast- and bacteria-specific
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37 550 probes with distinct fluorochromes to decipher microbial co-occurrence patterns in the newly
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39 551 created biofilm. One drawback should be noted as there is a loss in biofilm orientation during
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41 552 treatment. Dynamic follow-up by FISH coupled to CLSM showed no particular cell
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43 553 configuration within the biofilm except for AAB, which clearly dominated and progressively
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45 554 surrounded large yeast clusters. These cell interactions most likely enhance the symbiotic
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47 555 relationship and cooperative metabolism that occurs between the main microbial drivers during
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49 556 fermentation. SEM observations confirmed and even completed this observation and the
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51 557 biofilm constituents appeared to be actively secreted from some bacterial cells.
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53 558 *Gluconacetobacter* spp. (Mikkelsen et al., 2009; Zhang et al., 2018) and, in particular, *K.*
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55 559 *xylinus* (formerly *G. xylinus*) (Yamada et al., 2012) or *K. hansenii* (formerly *G. hansenii*) (Hodel
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560 et al., 2020), have all been described to produce biofilms and, as mentioned, our tailor-made
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2 561 consortium included biofilm-producing *K. hansenii* and *G. oxydans* strains (an easily
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4 562 observable trait in liquid cultures). Interestingly, comparable microscopic observations have
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6 563 already been made in other studies. Dima *et al.* (2017) studied a natural biofilm with similar
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8 564 composition (species belonged to *Komagataeibacter*, *Gluconobacter Zygosaccharomyces*,
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11 565 *Brettanomyces*, *Pichia* genera but also included some LAB) and observed a cellulosic cluster
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13 566 although bacteria and yeasts could not be accurately distinguished. In addition, El-Taher,
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15 567 (2011) also performed analyses on a biofilm containing *P. occidentalis* and *K. xylinus* and both
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17 568 were clearly embedded in the biofilm. Interestingly, SEM observations on biofilms produced
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20 569 from pure cultures (*K. xylinus* or *K. hansenii*) clearly highlighted a more organized cellulosic
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22 570 network with several fibrils and no apparent presence of embedded bacteria (Gromovykh *et*
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24 571 *al.*, 2020; Mikkelsen *et al.*, 2009; Zhang *et al.*, 2018). Such observations suggest that other
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26 572 microbial interactions exist between bacteria and yeasts within the Kombucha biofilm and the
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28
29 573 resulting biofilm may be different according to the species present. Yeasts and LAB (Edwards
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31 574 *et al.*, 2000) may also produce biofilms or participate in its production. The biofilm mode of life
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33 575 of the wine spoilage yeast, *D. bruxellensis*, has been recently described and may be linked to
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35 576 a potential resistance strategy to persist in the winemaking environment or in wines (Lebleux
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38 577 *et al.*, 2020). In Kombucha, this species is a key driver of the fermentation and may play a
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40 578 participative role in biofilm formation, together with biofilm-producing AAB. Our microscopic
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42 579 observations suggest biofilm filaments were released from some yeast cells meaning at least
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44 580 one or more yeast species can produce it during fermentation. Yeast biofilm production, in
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46 581 particular for *Dekkera/Brettanomyces* as well as *Saccharomyces* or *Debaryomyces* species,
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49 582 has already been described in co-cultures with LAB species (Furukawa *et al.*, 2010; Kawarai
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51 583 *et al.*, 2007; León-Romero *et al.*, 2016; May *et al.*, 2019). Noteworthy, *L. nagelii* dextran
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53 584 production from sucrose has also been reported (Edwards *et al.*, 2000) and despite the fact
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55 585 we could not observe *L. nagelii* clusters in the biofilm, LAB species, which reached > 8 log
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58 586 CFU/g biofilm, were detected using 16S rRNA gene metabarcoding so its role in biofilm
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60 587 formation cannot be excluded. Furthermore, in a recent study focusing on the interactions of
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588 *L. nagelii* and *Saccharomyces cerevisiae* encountered in water kefir, it was found that *L. nagelii*
1 profits from amino acids (i.e., glutamine, histidine, methionine, and arginine) and riboflavin
2 589
3 released by *S. cerevisiae* (Bechtner et al., 2019). If the Kombucha microbiota is well embedded
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6 591 in this biofilm structure, it can be clearly assumed that cooperative metabolism takes place and
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8 is even facilitated. Further work would therefore be necessary to provide a deeper
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10 understanding of the interactions occurring between yeasts, AAB and LAB.
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13 We also observed many distinct biochemical changes during fermentation. As mentioned,
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15 yeasts hydrolyze the sole carbon source at the start of the fermentation, sucrose, into glucose
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17 and fructose, which the different microbial drivers use and induce many biochemical changes.
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19 Despite alcoholic fermentation by yeasts, ethanol levels remained relatively low during
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21 fermentation, suggesting that almost immediate oxidation of ethanol, by AAB, into acetic acid
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23 occurred. We actually observed a rapid increase in both yeast and AAB population during the
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25 first 11 days which would be concomitant with these findings. In fact, AAB highly abundant in
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27 biofilm samples at the liquid-air interface thus ideal conditions would be encountered for
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29 efficient acetic acid production. Other organic acids also increased during fermentation (lactic,
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31 gluconic, glucuronic acids), remained at relatively stable levels (i.e. oxalic and succinic acids)
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33 or were only detected at low levels (malic acid). Some studies (Jia et al., 2016; Ku et al., 2010)
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35 have already shown that malic acid is present in *Camellia sinensis* (L.) O. Kuntze tea which is
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37 coherent with our results. Interestingly, malic acid could be transformed by the malolactic
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39 transformation by *O. oeni* (Wojdyło et al., 2020), thus leading to an increase of lactic acid as
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41 observed in our study. However, although a significant decrease of malic acid was observed
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43 between d0 and d7, it then increased during fermentation. Lactic acid can also be oxidized by
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45 AAB species into acetic acid and acetoin as observed in the present study (Moens et al., 2014).
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47 Among organic acids, highest concentrations were for acetic acid with levels reaching between
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49 1.3 and 2.4 g/L, as frequently observed during Kombucha fermentations (Cardoso et al., 2020;
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51 Chakravorty et al., 2016; Coton et al., 2017). Lactic acid was the second most abundant acid
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53 with concentrations ranging between 0.3 and 0.8 g/L and most likely results from LAB
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615 fermentations. Among the other organic acids, glucuronic acid is considered as one of most
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2 616 important organic acids in Kombucha in regards to its potentially beneficial health properties.
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4 617 Indeed, this acid, by the glucuronidation process, has a detoxifying effect in humans by
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6 618 increasing the elimination of xenobiotics as well as endogenous metabolites as bilirubin,
7
8 619 oxidized fatty acids and excess steroid hormones (Viña et al., 2014). It also increases transport
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11 620 and bioavailability of polyphenols, which are naturally present in the tea used for Kombucha
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13 621 fermentation, and can therefore increase the antioxidant properties of the beverage (Leal et
14
15 622 al., 2018). Interestingly, this organic acid could be involved in the biosynthesis of other
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17 623 bioactive compounds as vitamin C or D-saccharic acid-1,4-lactone (Leal et al., 2018).
18
19 624 Glucuronic acid production was described in AAB and interestingly (Nguyen et al., 2015)
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21 625 highlighted that symbiosis between *D. bruxellensis* and *Gluconacetobacter intermedius* strains
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23 626 leads to a higher production rate. Gluconic acid is also produced by AAB as *G. oxydans* (Sainz
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25 627 et al., 2016) by oxidizing glucose. It was shown to be produced by some *Acetobacter* species
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27 628 when ethanol is depleted which is well correlated with our results as gluconic acid increased
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29 629 from d11 onwards. This organic acid is also of interest in Kombucha, but for its organoleptic
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31 630 properties. Indeed, this acid is naturally found in food products including fruits, plants, wine
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33 631 and honey and provides a refreshing sour taste (Sainz et al., 2016). Moreover, it is also used
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35 632 as an acidity regulator (E574) (Règlement (UE) N°1129/2011, n.d.) and has a pKa of 3.86, so
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37 633 in Kombucha, it probably leads to an acidity balance with acetic acid that has a pKa of 4.8.
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43 634 Correlation analyses highlighted how the observed microbial shifts could be linked to certain
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45 635 changes in the volatilome profile. All species included in the tailor-made consortium were
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47 636 identified by metabarcoding. Two bacterial species, *A. okinawensis* and *L. nagelii*, dominated
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49 637 during fermentation and were well correlated to the mid or final fermentation stages,
50
51 638 respectively, while among yeasts, *H. uvarum* dominated during the early fermentation stages
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53 639 before a clear shift was observed with *D. bruxellensis* dominating for the remainder of the
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55 640 fermentation. All four species also showed the strongest correlations with the identified volatile
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57 641 compounds. Interestingly, distinct changes in volatilome were directly linked to fermentation
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642 time with aldehydes and branched-chain alcohols most abundant in the first two fermentation
1 stages (i.e. 3-methylbutanal associated with malty odors at day 7 and methylbutan-1-ol
2 643 associated with waxy or soapy odors at days 11-14) followed by a much more complex
3
4 644 volatilome profile between d20 and d27. During this final fermentation phase ethyl esters (i.e.
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6 645 sweet, fruity odors), acids (i.e. acidic, dairy or cheesy odors) and ketones (i.e. fruity or buttery
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8 646 odors) dominated and were well correlated with *D. bruxellensis*, *A. tropicalis* and *L. nagelii*.
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10 647 They may all contribute according to their detection thresholds, together with organic acids, to
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12 648 the overall sensorial properties of the final product. Noteworthy, fermentations using the tailor-
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14 649 made consortium were afterward conducted in a 1.5 L volume containers for sensory
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16 650 evaluation by comparing it to classically back-slop produced kombucha (using a 6-person
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18 651 panel). The performed hedonic tests, using a dedicated sensory evaluation table (overall
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20 652 acidity, acetic taste, sweetness, tannins and overall appreciation), confirmed a satisfactory
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22 653 product with acidic, citrus and slightly fruity odors and taste when compared to the control
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24 654 fermentation (data not shown). Like with other fermented beverages, the perception of some
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26 655 aroma compounds can be masked by the presence of others or enhanced due to synergistic
27
28 656 effects. Among all the volatile compounds detected, acids and esters are most likely to shape
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30 657 the overall organoleptic property of the final product as directly linked to the most common and
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32 658 distinct descriptors of this product (i.e. acidic, refreshing, fruity...).

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34 660 This study provides new insights on a tailor-made complex consortium to drive Kombucha
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36 661 fermentation and has provided a better understanding of the roles of each microbial species.
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38 662 The microbial drivers efficiently recreated a biofilm during fermentation which can be a useful
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40 663 alternative to the classical backslopping procedure used for spontaneous fermentations. In the
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42 664 future, pilot or industrial scale fermentations using this tailor-made complex consortium should
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44 665 be done to confirm our results and further characterize the sensory properties and quality of
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46 666 the final product. It would also confirm that these microbial drivers can avoid sluggish or slow
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48 667 fermentations in large-scale productions thus ensuring consumer satisfaction year-round.
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50 668 Also, different tailor-made consortia (i.e. simple versus complex) could be used to drive this
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669 fermentation and improve not only the sensorial properties but also potentially increase the
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2 670 beneficial health effects by increasing the production of certain key metabolites. Finally, the
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4 671 long-term goal would be to completely replace the classical SCOBY biofilm used as a starter
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6 672 and drive this fermentation with the tailor-made complex consortium.
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11 674 **Conflict of Interest**

13
14 675 The authors declare that the research was conducted in the absence of any commercial or
15
16 676 financial relationships that could be construed as a potential conflict of interest.
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18 677

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25

26 681 **Author contributions**

28
29 682 MC, JM and EC obtained the funding and supervised the study. MC, JM and EC designed the
30
31 683 experiments. OS performed experimental work and analysed the data. EP provided technical
32
33 684 assistance for LC-MS analyses while JJ performed enzymatic organic acid determinations. AT
34
35 685 and MM performed GC-MS analyses on tea samples. Metabarcoding analyses were done by
36
37 686 JM. CD, OS, JJ, JM, EC and MC participated to sensorial analyses. Statistical and correlation
38
39 687 analyses on all data were done by MC, MP and AT. OS and MC drafted the manuscript and
40
41 688 all co-authors edited and proofread the manuscript.
42
43

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55 694 **REFERENCE**

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973 **Figure 1: Results of physico-chemical parameters followed during 27 days**

974 **Kombucha fermentations.** The pH and density were determined in the tea fraction at day
975 0, 7, 11, 14, 20 and 27 as well the biofilm mass. Results correspond to the average of three
976 biological replicates with standard deviations.

978 **Figure 2: Numeration of bacteria and yeasts during Kombucha fermentation in**

979 **tea and biofilm.** Yeasts, acetic acid bacteria (AAB), lactic acid bacteria (LAB) were
980 numerated at days 0, 7, 11, 14, 20 and 27 in tea and biofilm except d0 for this latter. Results
981 correspond to the average of three biological replicates with standard deviations and
982 expressed as log CFU/mL (tea samples) or log CFU/g (biofilm samples).

984 **Figure 3A and 3B: Relative abundances of bacterial (A) and yeast (B) species**

985 **throughout Kombucha fermentation.** Each replicates are represented in both tea (T) and
986 biofilm (B) as r1, r2 and r3 for each sampling dates 0, 7, 11, 14, 20, 27 days (d0, d7, d11, d14,
987 d20 and d27).

989 **Figure 4: Dynamic monitoring of biofilm formation by confocal laser scanning**

990 **microscopy using fluorescent *in situ* hybridization.** The figures A to L correspond to
991 CLSM acquisition changes in biofilm during fermentation at days 7, 11, 14, 20 and 27 (d0, d7,
992 d11, d14, d20 and d27). The natural fluorescence of the biofilm is represented in blue in these
993 acquisitions (e.g. Figure 4D) while specific hybridizations with probes targeting bacteria
994 (EUB338) and yeasts (EUK516) are represented in red (e.g. Figure 4B) and green (e.g. Figure
995 4C), respectively. Figures 4A, 4B, 4C, 4D, 4G, 4H, 4I were obtained on the same focal point
996 (z) while figures 4E, 4F, 4K and 4L were obtained in depth of sample and the figure 4J was
997 obtained with both the methods.

998

999 **Figure 5: Dynamic monitoring of biofilm formation by Scanning Electron**
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21000 **Microscopy.** The figures correspond to SEM acquisitions showing changes in biofilm during
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41001 fermentation at days 11 for figures A to D, 14 for E to H, 20 for I to L and 27 for M to P and
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71002 finally Q and R corresponds to days 20 and 27.
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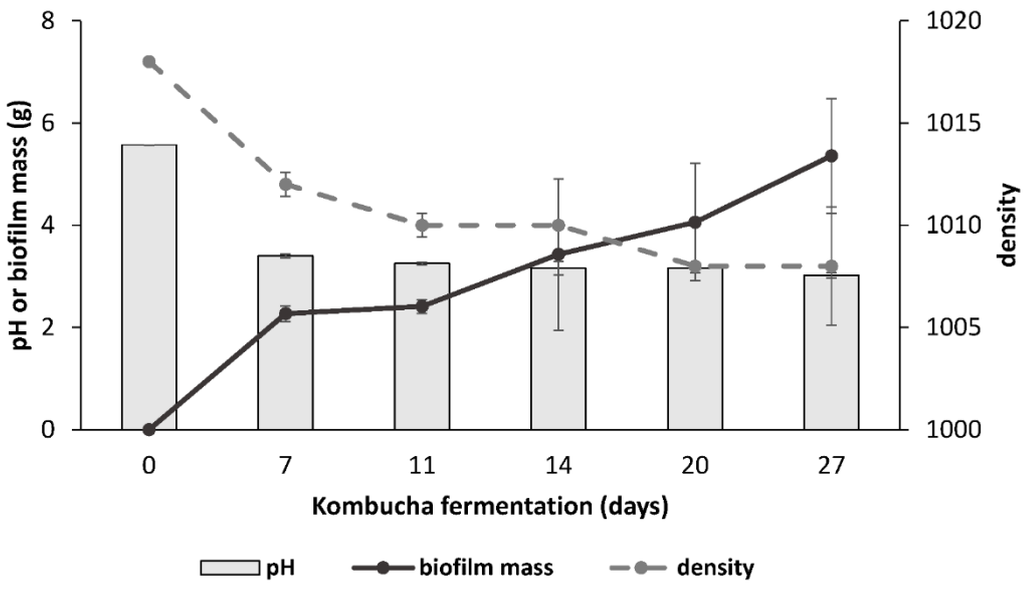
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111004 **Figure 6. Changes in organic acids concentrations during Kombucha**
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141005 **fermentations.** Gluconic, glucuronic, malic, oxalic and succinic acids were quantified by LC-
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161006 MS method (6A) while acetic and lactic acid were quantified by enzymatic kits (6B) and ethanol
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181007 by GC. Organic acids concentrations in g/L of Kombucha (tea) were quantified at 7, 11, 14, 20
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211008 and 27 fermentation days. The letter appearing corresponds to the results of the comparing
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231009 means tests. The results correspond to the averages of three biological replicates.
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251010
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281011 **Figure 7: Normalized heat-map representation of changes in Kombucha**
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301012 **volatilome during fermentation.** Hierarchical clustering was done using Ward's linkage
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331013 and Euclidean distances. Sample names and fermentation times are provided on the bottom
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351014 and volatile compound names on the right side as well as the four clusters. The green to red
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371015 color range indicates low to high compound abundances.
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391016
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421017 **Figure 8. Multiple factor analysis performed on samples collected at d7, d11, d14,**
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441018 **d20 and d27 with three groups of active variables: pH and density, organic acid**
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471019 **concentrations and abundance of 39 volatile compounds.** Individual factor map (A)
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491020 and variable factor map (B). Microbial data from culture-dependent (n= 6) and -independent
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511021 (n=18) analyses of tea and biofilm and biofilm wet mass were used as supplementary
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531022 variables. The ellipses on plot (A) show the three groups from agglomerative hierarchical
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561023 clustering on results from MFA. The variables poorly represented on plot (B) are not shown
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581024 ($\cos^2 < 0.5$).
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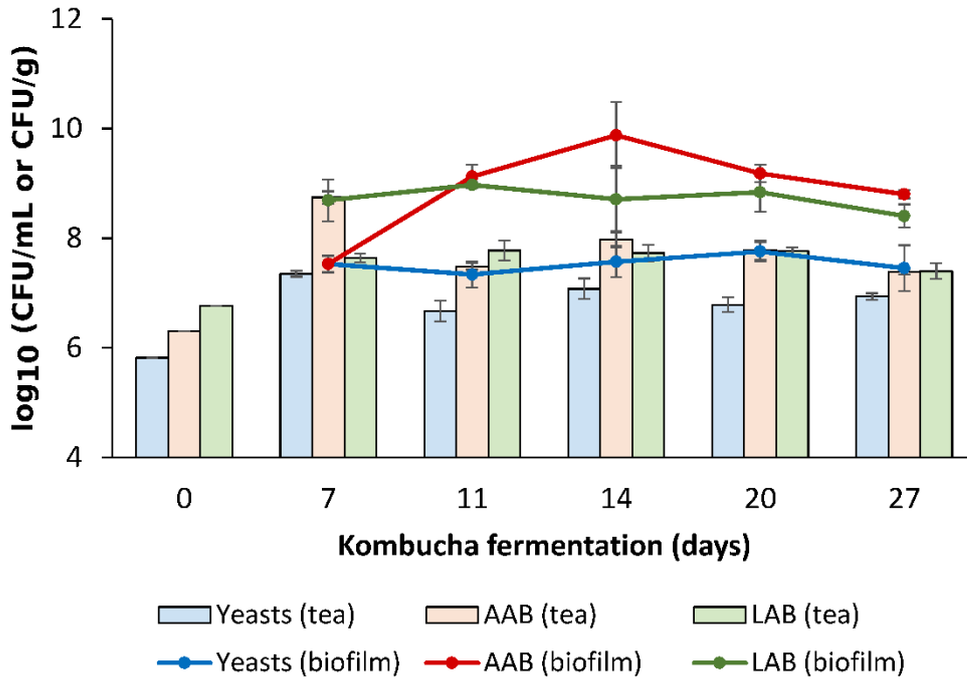
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1026 **Figure 1.**

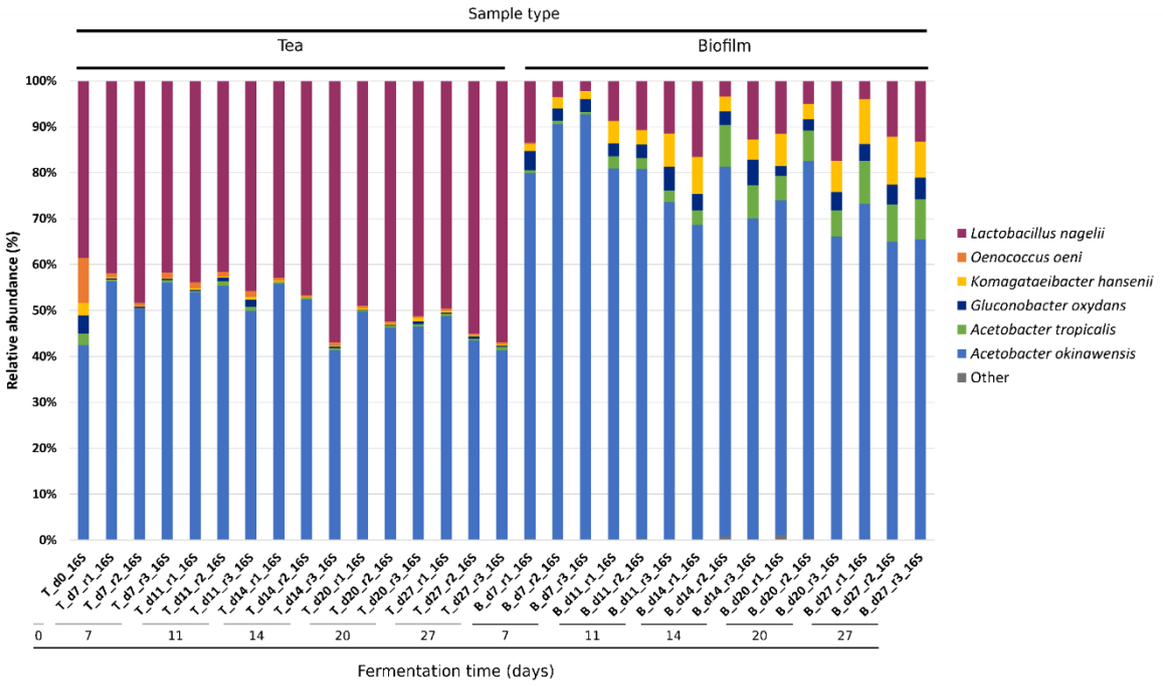


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1028 **Figure 2.**



1030 **Figure 3A.**



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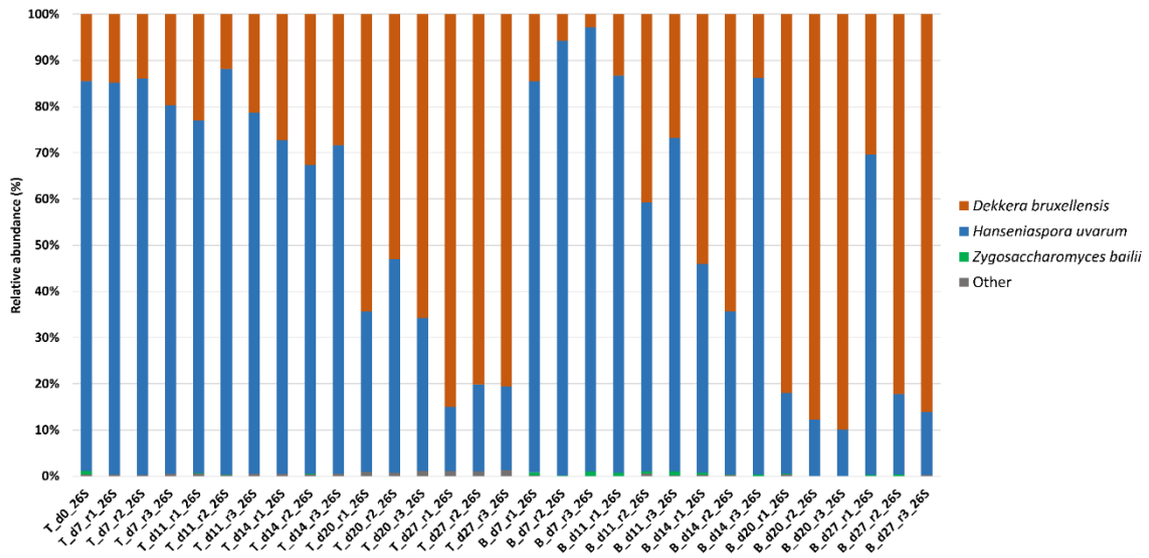
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Figure 3B.



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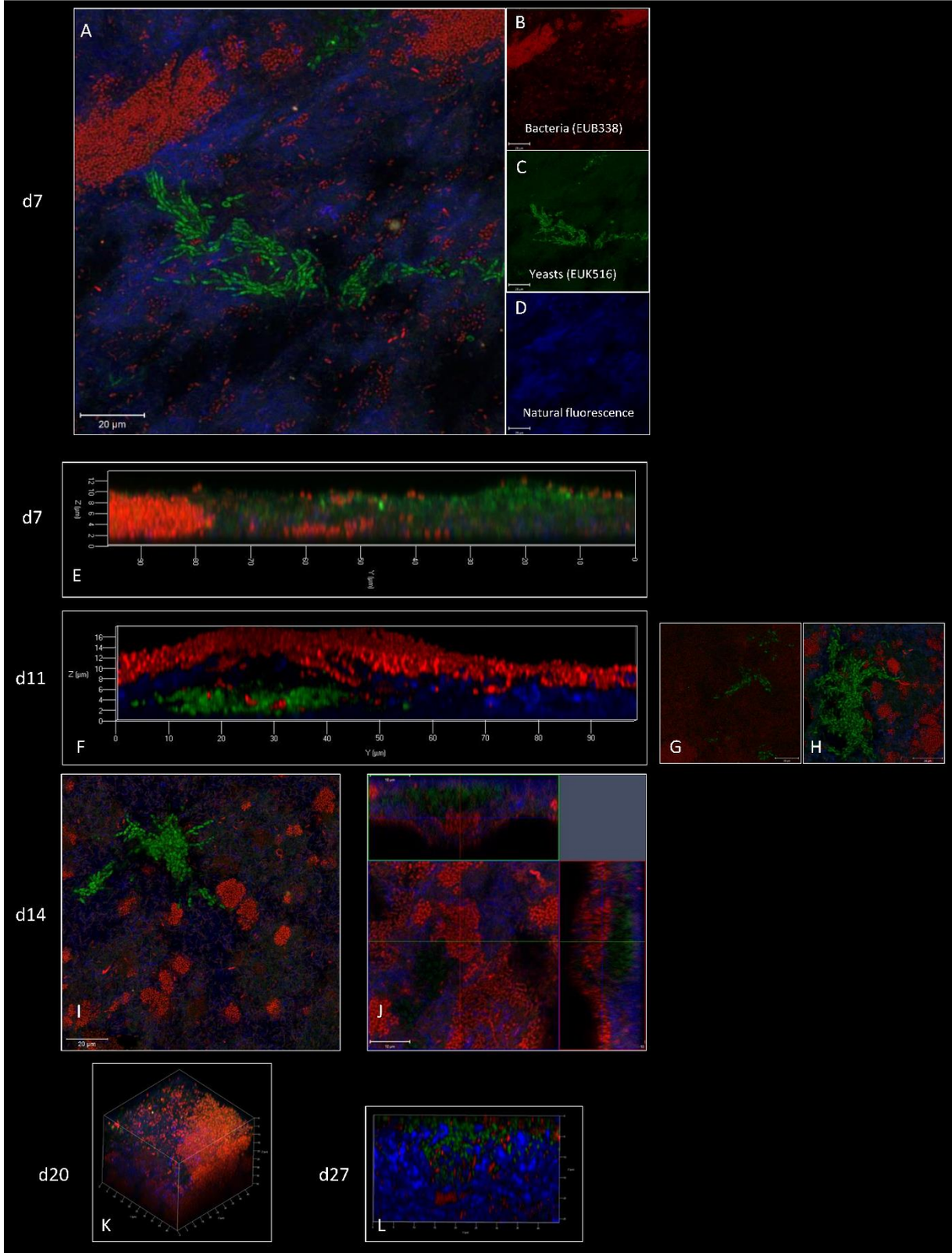
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1034 **Figure 4.**



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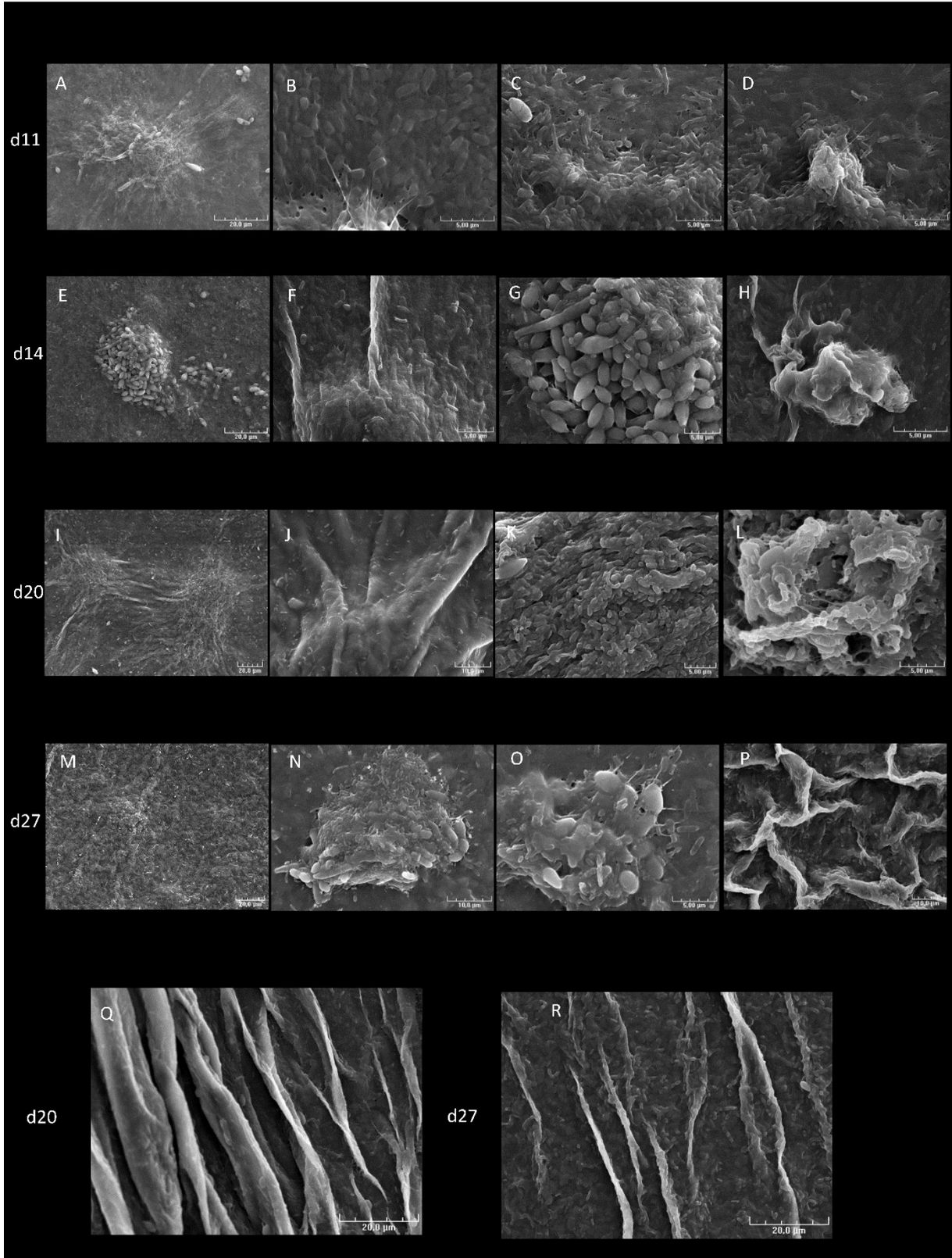
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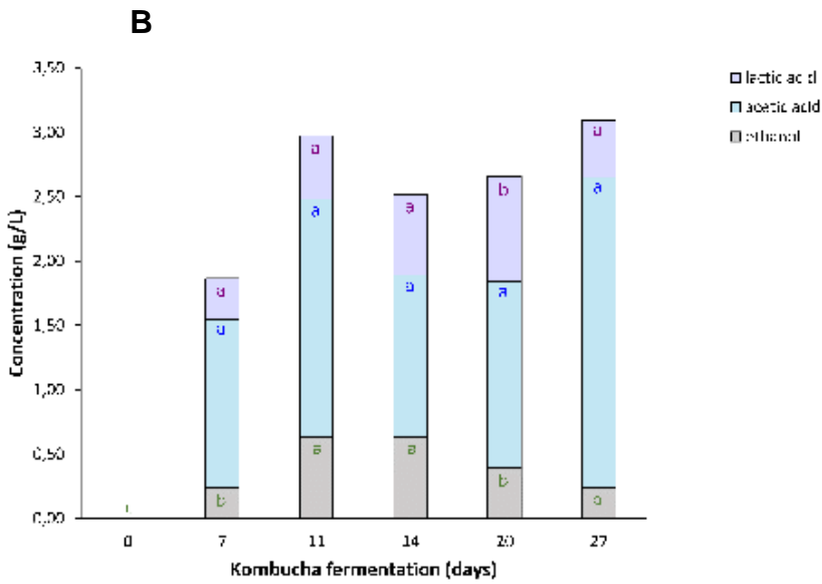
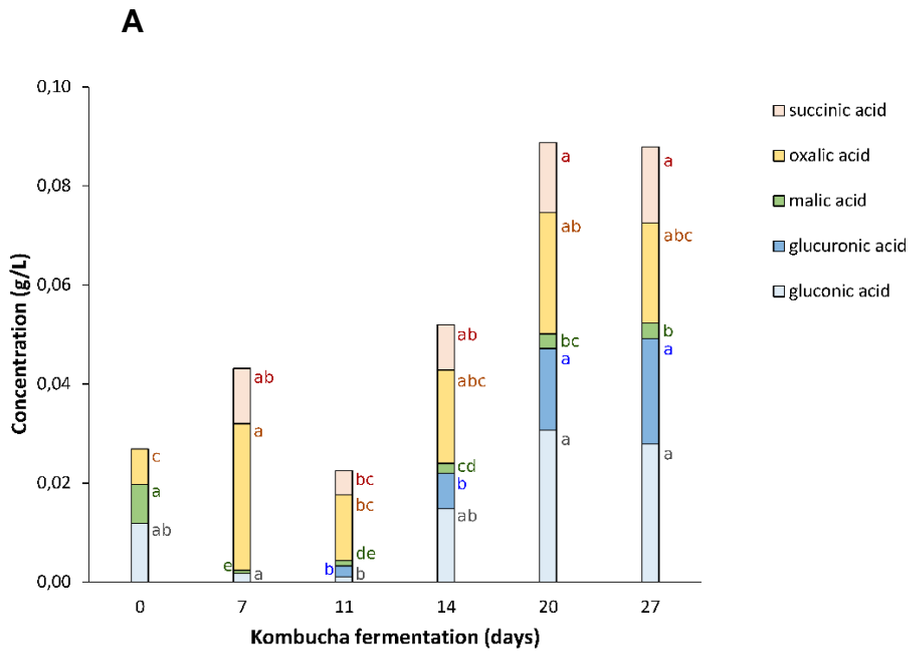
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1038 **Figure 5.**



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1040 **Figure 6.**

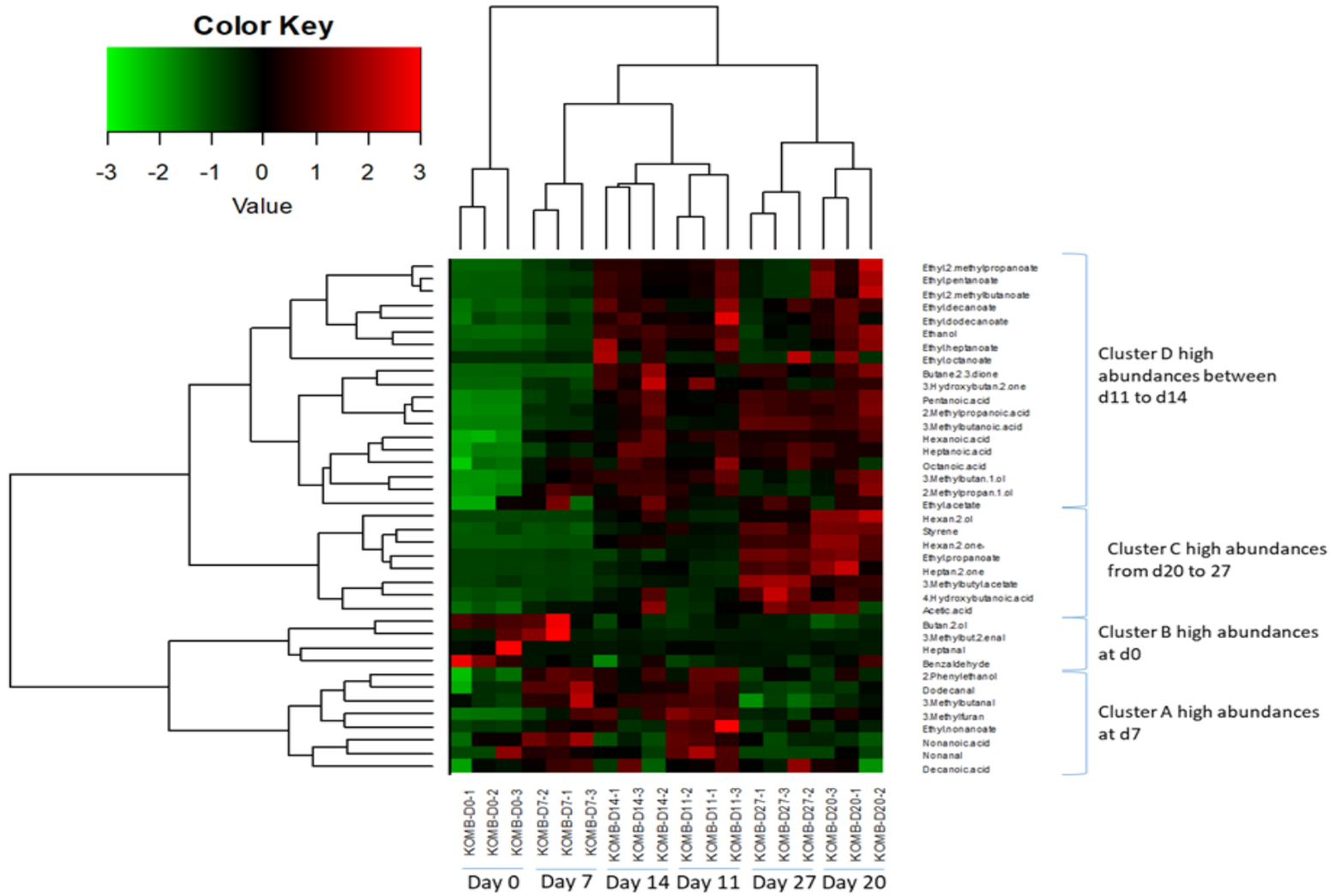


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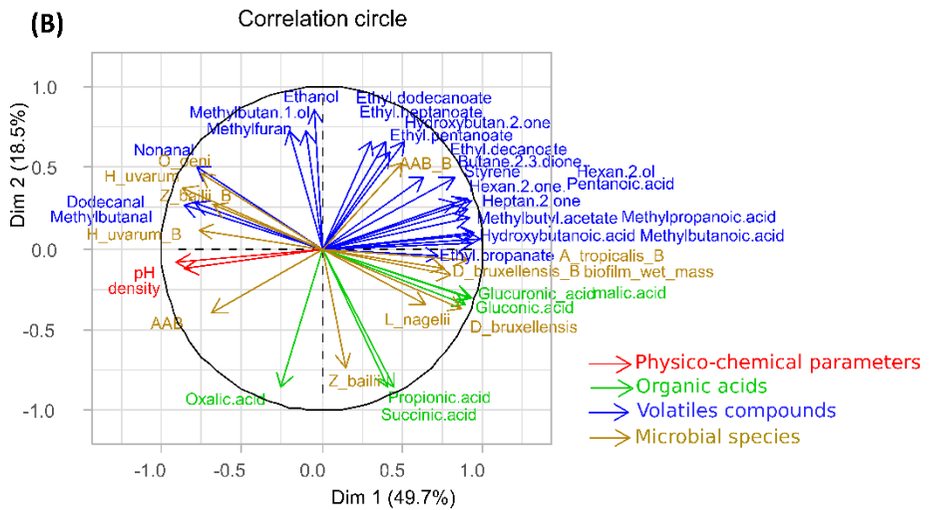
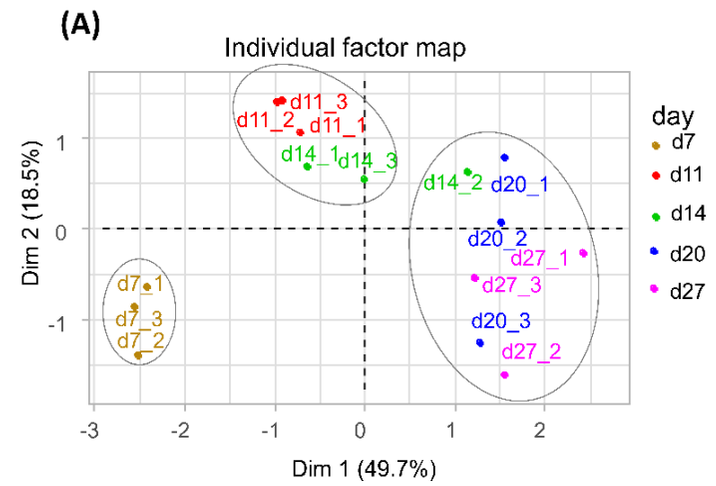
1044 **Figure 7.**



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1046 **Figure 8.**



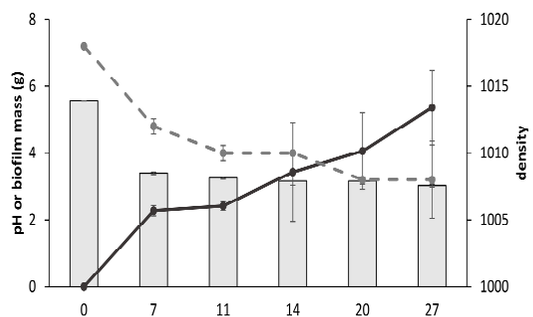


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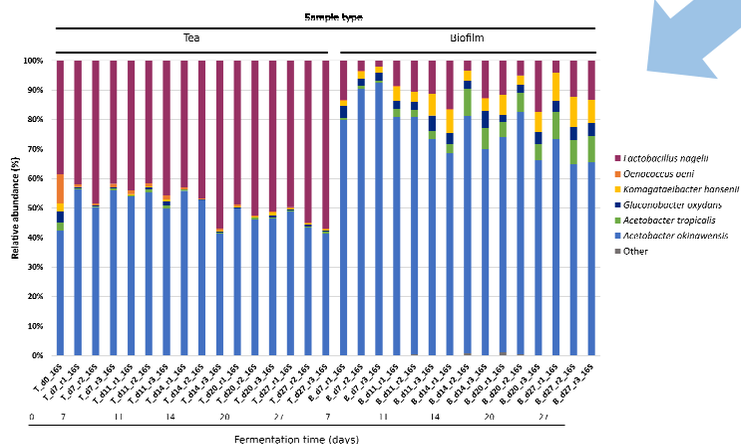
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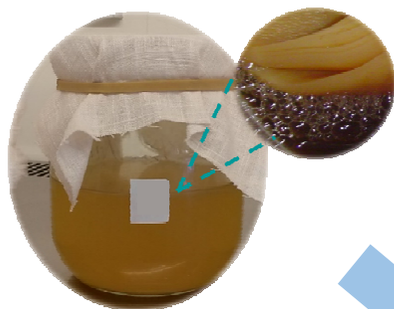
Microbial, biochemical and physico-chemical changes



Species dynamics monitored by metagenetics

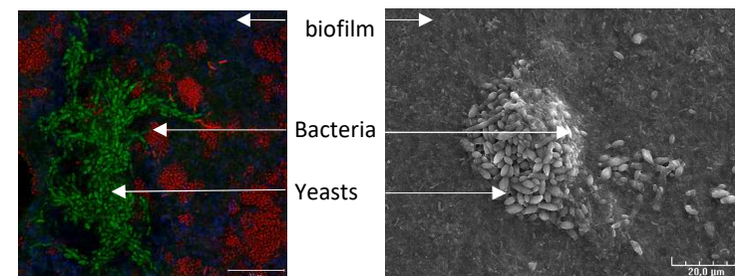


Defined tailor-made complex consortium with key microbial drivers

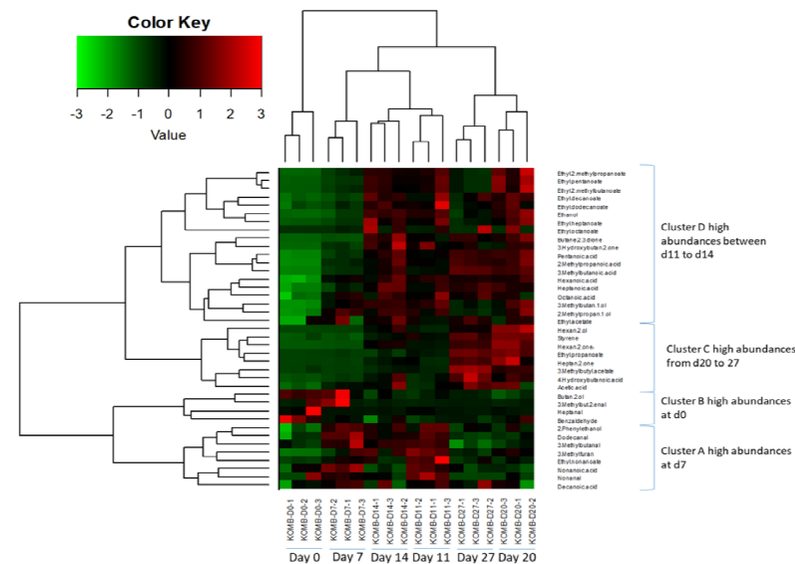


Dynamic follow-up

Biofilm formation and microbial network



Kombucha volatilome elucidated



Author contributions

MC, JM, CD and EC obtained the funding and supervised the study. MC, JM and EC designed the experiments. OS performed experimental work and analysed the data. EP provided technical assistance for LC-MS analyses while JJ performed enzymatic organic acid determinations. AT and MM performed GC-MS analyses on tea samples. Metabarcoding analyses were done by JM. Statistical and correlation analyses on all data were done by MC, MP and AT. OS and MC drafted the manuscript and all co-authors edited and proofread the manuscript.

Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: