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Optimization of orange peel waste ensiling for sustainable anaerobic digestion / Paolo, C., Fazzino, F., Sidari, R., Zema, D.A.. - In: RENEWABLE ENERGY. - ISSN 0960-1481. - 154:(2020), pp. 849-862. [10.1016/j.renene.2020.03.047]

Availability:

This version is available at: <https://hdl.handle.net/20.500.12318/59061> since: 2024-10-04T09:33:29Z

Published

DOI: <http://doi.org/10.1016/j.renene.2020.03.047>

The final published version is available online at: <https://www.sciencedirect>.

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3 ***Calabrò, P. S., Fazzino, F., Sidari, R., & Zema, D. A. (2020). Optimization of orange peel waste***
4 ***ensiling for sustainable anaerobic digestion. Renewable Energy, 154, 849-862.***

5
6 *which has been published in final doi*

7
8 10.1016/j.renene.2020.03.047

9
10 (<https://www.sciencedirect.com/science/article/pii/S0960148120303724>)

11
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14 **Optimization of orange peel waste ensiling for sustainable anaerobic digestion**

15

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17

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22

23 **Abstract**

24

25 Today, orange peel waste (OPW) is mainly used as cattle feed, often after ensiling. This storage
26 phase can increase the efficiency of anaerobic digestion, since it allows both a better management
27 of possible co-digestion and a reduction in the high content of essential oils (mainly composed of d-
28 Limonene a well-known inhibitor of anaerobic digestion). The effects of ensiling on the methane
29 potential of OPW have been little studied, particularly its microbiological profile. This study has
30 simulated, at laboratory scale, OPW ensiling under three different conditions. Ensiled OPW
31 samples were then either directly anaerobically digested or subjected to simple pretreatments aiming
32 at the further removal of d-Limonene. The microbiota evolution during ensiling and the species of
33 microorganisms present during the aforementioned process were also identified. After ensiling, up
34 to over 70% of the initial d-Limonene content of OPW was removed and biomethane yield was
35 preserved up to about 90%.

36

37 **Keywords:** *anaerobic digestion; d-Limonene; ensiling; microbiota; molecular identification;*
38 *orange peel waste.*

39

40 ***Highlights:***

41

- 42 • OPW ensiling under dry or uncontrolled conditions are the most suitable techniques
- 43 • During ensiling up to 63% of volatile solids in OPW are lost
- 44 • Ensiling allows d-Limonene removal up to 75%
- 45 • Up to about 90% of the methane potential of fresh OPW is preserved by ensiling
- 46 • The microbiological population shows a high biodiversity.

47

48 **1. Introduction**

49

50 Orange peel waste (OPW), the residue of orange juice production, is about 50-60% of the processed
51 fruit (in weight). The specific physical and chemical properties of OPW (the high amount and
52 seasonal nature of the production, the low pH, the high water content, the presence of essential oils -
53 mainly d-Limonene, 80-95% of the total) make its management difficult [1,2]. OPW has a high
54 biorefining potential [1] both for the extraction of added-value products (e.g., pectin and bioactive
55 compounds) and biofuels (e.g. biogas from anaerobic digestion - AD) [3–9]. However, the
56 biomethane yield of OPW is curbed due to its high content of d-Limonene [6], which is highly toxic
57 to microorganisms [2,10]. Three alternatives are available to overcome this limitation: (i) the
58 preliminary removal of d-Limonene [5,7,11–13]; (ii) the co-digestion with other substrates [14–18];
59 (iii) the digestion of OPW alone adopting moderate organic loading rates (OLRs) and/or using
60 additives [6,8,9].

61 Since the advanced removal of d-Limonene from OPW is expensive and the digestion of OPW
62 alone due the aforementioned problems reduces the overall economic convenience of the process,
63 co-digestion is a more promising management option. However, its present application for energy

64 conversion of OPW is limited, since AD plants located in citrus production areas are not able to
65 treat the high amounts of residues produced during the limited time of the harvesting season (from
66 November to April in the Mediterranean climate) and long distance transportation is economically
67 unsustainable. Therefore, OPW is traditionally used as animal feed [19–22] and ensiling [17,23] is
68 commonly used, as for forages, for conservation throughout the year.

69 The ensiling process is commonly divided into subsequent four steps [24–26]: (1) an aerobic phase,
70 beginning immediately after process start, when aerobic bacteria and yeasts predominate, thanks to
71 the air entrapped in the biomass; (2) a fermentation phase, when anaerobic and facultative
72 microorganisms use the available substrates for their metabolism, producing mainly organic acids;
73 (3) a steady storage phase in the silage silo, when the reduced pH after the previous phase allows
74 the substrate preservation; and (4) the feed-out phase, when the material is exposed to air for the
75 subsequent use (the latter stage is not considered in this paper).

76 During ensiling, the properties of raw OPW (e.g. pH and volatile solids) are quickly made stable
77 due to a spontaneous lactic fermentation. Stabilisation is normally completed in about two weeks
78 [27,28], when pH becomes slightly higher than three. The changes are also macroscopically
79 evident: in 10-20 days OPW can not be visually recognizable, since the original substrate becomes a
80 dense homogeneous slurry and only the seeds remain intact (Figure 1 - SI).

81 According to the literature, the main product of fermentation is lactic acid and, secondarily, ethanol
82 and acetic acid. .

83 However, until now the effects of ensiling on the methane potential of AD of OPW have been little
84 studied, despite the potential increases in methane yields that can be expected. Previous results of
85 experimental tests of AD of ensiled OPW [7,28,29] have shown, beside the viability of the process,
86 that the methane production per unit of digested biomass weight is similar to the energy yield of the
87 raw substrate. The d-Limonene is partially removed during the process, but a noticeable loss of
88 volatile solids (VS) is observed. Overall, ensiling provides preliminary homogenization, hydrolysis
89 and acidification of OPW.

90 However, until now OPW ensiling has not been optimized in view of using the ensiled material as a
91 substrate for AD, whose objective is the maximization of the methane yield. Therefore, more
92 research is needed in order to identify the most sustainable ensiling technique to be used as OPW
93 pre-treatment in AD plants. Moreover, little has been reported in the literature about microbiota of
94 OPW fermentation during ensilage [27].

95 To fill these gaps, this study explores a set of possible conditions and treatments for OPW ensiling,
96 targeted to maximise d-Limonene removal and, at the same time, limiting the biomass loss. In more
97 detail, the ensiling process is simulated at the laboratory scale under (i) natural, (ii) wet (adding
98 20% water to raw OPW), and (iii) dry (in a drainage system purposely prepared) conditions. In
99 order to remove as much d-Limonene possible, all samples of ensiled OPW are then subjected to (i)
100 simple centrifugation and (ii) ethanol extraction and centrifugation. Moreover, the microbiota
101 evolution of OPW and the species of microorganisms involved in the ensiling process are evaluated.
102 Finally, the overall loss of VS and the bio-methane potential (BMP) of the samples have been
103 evaluated.

104

105 **2. Materials and methods**

106

107 *2.1. OPW sampling*

108

109 OPW was sampled from an orange processing factory in Reggio Calabria (Southern Italy) and
110 immediately frozen (-20°C). According to [30], freezing is not expected to affect the biological
111 activity of the biomass. Before starting ensiling, the samples of OPW were thawed at room
112 temperature

113

114 *2.2. OPW ensiling*

115

116 OPW was ensiled in hermetically sealed batches. Each batch (made of glass, with a volume of 1.1
117 L) is provided with a central neck, closed with a stopper, and two side openings closed with rubber
118 septa that allow the biogas withdrawal. Three ensiling conditions were tested: (i) natural conditions
119 (hereinafter indicated as “ENS”), as usually carried out by agro-farms of the Mediterranean Basin
120 (ii) “wet” conditions, where water (20% w/w) was added to OPW (“WET”) to try to improve d-
121 Limonene leaching; (iii) “dry” conditions (“DRY”), placing OPW over a drainage system (quartz
122 gravel), in order to remove by gravity the liquid released by the biomass in order to reduce the
123 moisture and speed the stabilization process.

124 For each ensiling condition, six batches were prepared: three batches were opened after 7, 14 and 21
125 days respectively, in order to evaluate the changes (weight loss, TS, VS, COD, microbiota) in OPW
126 throughout the process. The remaining three batches were opened after 28 days, when ensiling was
127 stopped. In fact, in previous studies a substantial stability of the ensiled biomass was observed after
128 2-4 weeks [25–27]. In these three batches, the volume of the biogas produced during ensiling was
129 measured three times per week using a graduated 100-mL syringe.

130

131 *2.3. Treatments on ensiled OPW*

132

133 After ensiling, the OPW was extracted from the batches and subjected to two treatments, in order to
134 further remove d-Limonene: (i) a chemical treatment followed by centrifugation and (ii) a simple
135 centrifugation. As regards the first treatment, each sample of ensiled OPW (under ENS, WET and
136 DRY conditions) was chemically treated (hereinafter the treated samples were referred as “CHEM”)
137 using ethanol as solvent for d-Limonene extraction and then centrifuged at 9000 rpm for three
138 minutes. Solvent was dosed at 10% w/w with a contact time of one hour under continuous mixing
139 in a rotary shaker, Stuart Scientific Rotator Drive STR/4).

140 Moreover, OPW after ensiling (also in this case under ENS, WET and DRY conditions) was also
 141 subjected to simple centrifugation (CEN) only (that is, without a previous treatment with ethanol),
 142 in order to evaluate the efficiency of d-Limonene removal by the chemical treatment.

143 In both treatments, the liquid from the centrifuge was disposed of, while the solid biomass was used
 144 as substrate for BMP tests.

145 The OPW samples were weighed before and after ensiling, and before and after each treatment, in
 146 order to estimate the various mass flows. Table 1 reports a scheme of the experimental tests carried
 147 out on the nine samples.

148

149 **Table 1.** Acronyms of the OPW samples subjected to the experimental tests (ENS - naturally
 150 ensiled OPW; WET - OPW ensiled in wet conditions; DRY - OPW ensiled in dry conditions).

151

		Treatments		
		<i>Natural ensiling</i>	<i>Centrifugation</i>	<i>Chemical (ethanol addition)</i>
Condition	<i>Natural</i>	ENS	ENS+CEN	ENS+CHEM
	<i>Wet</i>	WET	WET+CEN	WET+CHEM
	<i>Dry</i>	DRY	DRY+CEN	DRY+CHEM

152

153

154 2.4. Physico-chemical measurements

155

156 Before and after ensiling and treatments, pH, contents of total (TS) and volatile (VS) solids, and
 157 chemical oxygen demand (COD) of OPW were measured following standard methods [31]. As
 158 suggested in [32,33], we cared to prevent the loss of as much of volatile compounds as possible,
 159 such as some components of the essential oils (EO), acetic acid and, if present, ethanol and other

160 alcohols. To this aim, during TS measurement we usually limit oven temperature at 60 °C. Under
161 this temperature, water evaporation can be considered complete when stable weight is reached.

162 For COD measurement first each OPW sample was dried. Subsequently, it was milled and the
163 powder was then mixed to distilled water. Finally, COD was measured by the potassium dichromate
164 method using pre-dosed cell tests (WTW 114555) the method complies with the DIN ISO 15705
165 and is similar to APHA 5220 D method.

166 As regards the determination of the concentration of d-Limonene before and after the experimental
167 tests, the analysis is difficult for OPW, since the concentration is strongly influenced by the
168 extraction conditions and the degradation level of the substrate. Moreover, the complexity increases
169 if the possible inhibition of AD process must also be measured. Since during ensiling the biomass
170 was homogenised, presumably most of the EO was released throughout the process due to the
171 breaking of the small flavedo sacs which contain it. Following previous tests [9], which used a
172 “mild” EO extraction only the d-Limonene that was available immediately after substrate feeding
173 was determined .

174 d-Limonene was extracted from the biomass by mixing 1.5 g of sample with 3 mL of a solution of
175 toluene (Sigma-Aldrich, St. Louis, MO, USA) and cyclohexane (0.1M, Sigma-Aldrich, St. Louis,
176 MO, USA), which was used as internal standard, for two hours. This blend was then injected into a
177 gas chromatograph (Agilent 6890) equipped with a wide-bore capillary column and a flame
178 ionization detector (FID), the latter set at 250 °C. The capillary column (J&W DB-WAXetr 50 m x
179 320 mm x 1 mm) used nitrogen as gas carrier with a flow rate of 10 mL/min. The temperature,
180 initially kept 50 °C for 8 min, was then raised to 230 °C (at 5 °C/min) for 2 min and finally set at
181 240 °C for 4 min during the post run.

182 The liquid recovered after centrifugation of ENS and WET samples and that collected at the bottom
183 of the DRY ensiling reactor were analysed for propionic, butyric and lactic acids. The liquid
184 samples were filtered (1.2 µm) twice and then 2.5 mL of filtrate were mixed with 2.5 mL of ethyl
185 acetate (Sigma-Aldrich) and shaken to allow organic acids extraction.

186 The amount of organic acids extracted was determined with a gas chromatograph (Agilent 6890)
187 equipped with a wide-bore capillary column (CPWAX52CB, 50 m, i.d. ¼ 0.53 mm) and a flame
188 ionization detector (FID). The injector was settled at 250°C. The temperature program started at
189 50°C, held for 5 min, the temperature was raised to 230°C at 5°C/min, held for 8 min, raised to
190 240°C and held for 2 min during the post run.

191

192 *2.5. Biochemical methane potential (BMP) tests*

193

194 Three series of BMP tests were carried out in triplicate for each sample under mesophilic conditions
195 (35 ± 0.5 °C) as follows (Figure 1):

196

- 197 1. BMP1: ENS; WET; DRY.
- 198 2. BMP2: ENS+CEN; WET+CEN; DRY+CEN.
- 199 3. BMP3: ENS+CHEM; WET+CHEM; DRY+CHEM.

200

201 In each series the blank (that is a batch to assess the biogas production of the inoculum) and, as
202 additional internal controls, two other batches were added; the first fed with cellulose and the other
203 with raw OPW. The cellulose-fed reactor is suggested by UNI/TS 11703:2018 (Italian standard
204 procedure for BMP tests) in order to verify inoculum activity. The second was designed as an
205 internal control to verify the response of the different inocula to the substrate.

206 As inoculum of the AD process, a liquid digestate was used collected in three separate sampling
207 operations from a full-scale anaerobic digester fed with cattle manure and agro-industry residues.
208 After collection, the inoculum was sieved and stored for less than a week at 35 °C to reduce non-
209 specific biogas production (i.e. the production of the inoculum itself). The TS of the inoculum of
210 the three BMP tests was $5.5 \pm 0.2\%$, the VS $69.5 \pm 2.4\%$ and the pH was 7.5 ± 0.05 .

211 For each BMP test 1.1-L bottles with a central neck and two other lateral necks equipped with
212 perforable septa (WTW-Germany) were used. Each bottle was placed on a magnetic stirrer, and the
213 digestion blend was continuously mixed in a thermostatic cabinet kept at a preset temperature
214 (35 ± 0.5 °C).

215 In each batch the substrate was mixed with 200 mL of inoculum at a ratio (on a VS basis) equal to
216 0.3, this value being in the range suggested by UNI/TS 11703:2018. According to the same
217 regulation three nutrient solutions were also added, to supply nutrients and micronutrients for the
218 bacterial metabolism. The three solutions (indicated as A, B and C) contained KH_2PO_4 ,
219 $\text{Na}_2\text{HPO}_4 \cdot 12\text{H}_2\text{O}$, NH_4Cl (A, 5% final volume), $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$, $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$, $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ (B,
220 5% of final volume) and $\text{MnCl}_2 \cdot 4\text{H}_2\text{O}$, H_3BO_3 , ZnCl_2 , CuCl_2 , $\text{Na}_2\text{MoO}_4 \cdot 2\text{H}_2\text{O}$, $\text{CoCl}_2 \cdot 6\text{H}_2\text{O}$,
221 $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$, Na_2SeO_3 (C, 1% of final volume). Finally, water was added to the batch, in order to
222 reach final volume (600 mL) and to keep the TS content at about 35 gTS/L, which is consistent with
223 the limits (10-50 gTS/L) required by the aforementioned UNI/TS regulation. In accordance with
224 this regulation, the BMP tests were stopped when the daily methane production of a batch was
225 lower by 1% than the cumulated volume from the process start.

226 About three times per week, the biogas produced in each batch was withdrawn using a 100 mL
227 syringe and transferred with care into an alkaline trap through a tube. After the injection, the carbon
228 dioxide in the biogas was absorbed by an alkali solution (NaOH 3M), while the methane bubbles,
229 increasing the pressure in the trap, displaced the same volume of the alkali solution, measured in a
230 graduated cylinder. The test was stopped when daily production was lower than 1% of the
231 cumulated value since test start. The net specific methane production (that is, the methane volume,
232 normalised to standard conditions, per unit of VS depurated by the blank production) was calculated
233 as follows:

$$234 \quad BMP = \frac{(V_{CH_4,s} - V_{CH_4,blank})}{VS_s \cdot V_s} \quad [1]$$

235

236 where:

237

238 - $V_{CH_4,s}$ = final cumulated methane production (NmL_{CH_4})

239 - $V_{CH_4,blank}$ = final cumulated methane production of the blank (NmL_{CH_4})

240 - V_{S_s} = initial VS concentration of the substrate ($g_{VS} \cdot L^{-1}$)

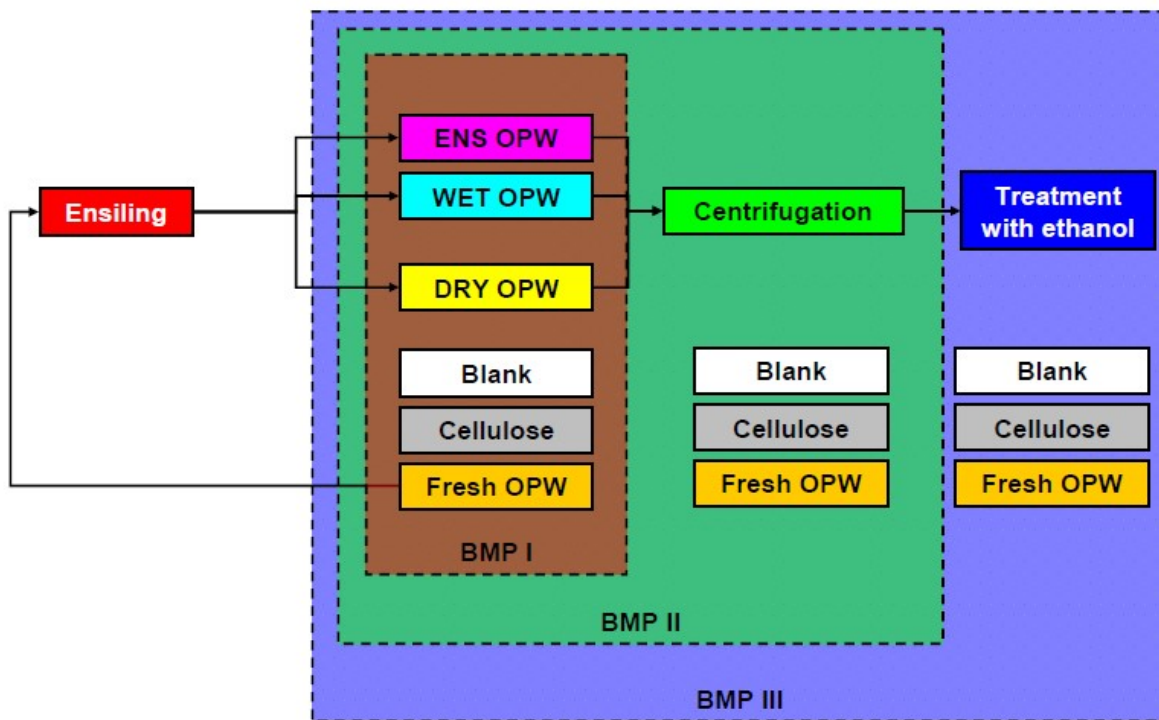
241 - V_s = total volume of the batch (L)

242

243 According to the aforementioned Italian standard procedure, the test was accepted, if the batch fed

244 with cellulose in the same BMP series produced $335 NmL \cdot g_{VS}^{-1} \pm 25\%$.

245



246

247 **Figure 1.** Experimental scheme of the BMP (I, II and III) tests carried out after OPW (orange peel

248 waste) ensiling subjected to different ensiling conditions and treatments (ENS - naturally ensiled

249 OPW; WET - OPW ensiled in wet conditions; DRY - OPW ensiled in dry conditions).

250 2.6. BMP kinetic modelling

251

252 The net specific cumulative methane production of each BMP test was modelled using the modified
253 Gompertz equation [34], in order to verify its prediction capacity under the experimental conditions:

254

$$255 \quad B = P \cdot \exp \left\{ -\exp \left[\frac{R_m \cdot t}{P} \cdot (\lambda - t) + 1 \right] \right\} \quad [2]$$

256

257 where:

258 - B ($\text{NmL} \cdot \text{g}_{\text{VS}}^{-1}$) = specific methane production at time t (d)

259 - P ($\text{NmL} \cdot \text{g}_{\text{VS}}^{-1}$) = specific methane production at $t = \infty$

260 - R_m ($\text{NmL} \cdot (\text{g}_{\text{VS}} \cdot \text{d})^{-1}$) = maximum methane production rate

261 - λ (d) = lag phase duration.

262

263 P , R_m and λ were calculated using iteratively the least square method of the routine “Solver” of
264 Microsoft Excel until to the highest r^2 between the modelled and experimental data.

265

266 2.7. Statistical analyses

267

268 First, the statistical significance of the final values of the weight loss as well as TS and VS contents
269 after the ensiling tests was investigated using the t-test (at $p < 0.05$).

270 Then a two-way Analysis Of Variance (ANOVA) along with Tukey’s test (designed for the
271 pairwise comparisons) was used to evaluate the statistical significance of the net cumulated specific
272 methane yields of the batches, assuming as variability factors: (i) the ensiling conditions (ENS,
273 WET and DRY); (ii) the treatment (raw OPW, natural ensiling, chemical treatment and
274 centrifugation); (iii) reciprocal interaction of ensiling condition and treatment. At $p < 0.05$ level of

275 significance was adopted. It was not necessary to perform data transformations for the analysis.
276 ANOVA assumes normality and this assumption was checked using QQ-plots.
277 All the statistical analyses on the samples were carried out using the XLSTAT (release 2017)
278 software.

279

280 *2.8. Microbiological analyses and strains isolation*

281

282 The microbiota associated with raw and ensiled OPW (ENS, WET and DRY modes) was analysed
283 at day 0 (raw OPW), 7, 14 and 28 of the ensiling period, the leachate collected during DRY ensiling
284 was also considered. To this end, each type of solid sample was firstly homogenized to allow the
285 microorganisms release from the solid matrix; more specifically, 10 g of each solid OPW sample
286 was homogenized in a solution of 0.9% NaCl. Then, the obtained homogenates and the leachate of
287 DRY OPW were diluted ten-fold and inoculated by spread-plate method in triplicate onto Petri
288 plates, containing: (i) Plate Count Agar (PCA) (Sigma-Aldrich), for total microbial count (TMC);
289 (ii) de Man–Rogosa–Sharpe (MRS) agar (VWR International srl, Italy), supplemented with 15
290 mg/L cycloheximide (Oxoid), to count lactic acid bacteria (LAB); and (iii) Yeast Peptone Dextrose
291 (YPD) agar (VWR, International srl, Italy), supplemented with 100 mg/L chloramphenicol
292 (Liofilchem Diagnostici, Italy), to count yeasts. All the plates were incubated at 30 °C for two days
293 under aerobic conditions for yeasts and TMC, and under anaerobic conditions for LAB. At day 0, 7,
294 14, 21, and 28 during ensilage, the colonies grown on YPD and MRS agar were randomly picked
295 from the highest dilution sample [35]; then, the isolates were purified by streaking on the
296 corresponding isolation medium and stored as glycerol stock at – 80 °C until use. The isolated
297 bacteria were tested for catalase and for Gram by KOH method [36].

298

299 *2.9. Restriction analyses and sequencing*

300

301 DNA from overnight grown yeasts (101 isolates) and bacteria (97 isolates), isolated throughout the
302 ensilage and from the different treatments, was extracted by InstaGene Matrix (Bio-Rad
303 Laboratories, USA), according to the manufacturer's instructions. Then, yeasts were analyzed by
304 PCR of the 5.8S-ITS regions using the primers ITS1 (5'-TCCGTAGGTGAACCTGCGG-3') and
305 ITS4 (5'-TCCTCCGCTTATTGATATGC-3') and amplification conditions, according to [37], and
306 bacteria were analysed by PCR of the 16S rRNA gene, using the Y1 (5'-
307 TGGCTCAGAACGAACGCTGGCGGC-3') and Y2 (5'-CCCCTGCTGCCTCCCGTAGGAGT-
308 3') primers, according to [38]. Firstly, yeasts and bacteria were grouped by Restriction Fragment
309 Length Polymorphism (RFLP) of the 5.8S ITS rRNA region (HaeIII and HinfI restriction enzymes)
310 and Amplified Ribosomal DNA Restriction Analysis (ARDRA) of the 16S rRNA gene (HaeIII and
311 AluI restriction enzymes), respectively. Then, three samples for each PCR-RFLP and PCR-ARDRA
312 profile were chosen to sequence the 26S D1/D2 rRNA region (NL1 and NL4 primers) and 16S
313 rRNA regions (fD1 and rD1 primers) for yeasts and bacteria, respectively [39]. The obtained
314 amplicons were purified and sequenced by Sanger method (Eurofins Genomics, Germany). The
315 sequences were analyzed and compared with the sequences of the National Center for
316 Biotechnology Information (NCBI) using BLASTN [40]. To differentiate the genotypically closely
317 related *Lactobacillus plantarum*, *Lactobacillus pentosus*, and *Lactobacillus paraplantarum*, the
318 multiplex PCR of *recA* gene was carried out, according to [41].

319

320 **3. Results**

321

322 *3.1. Ensiling and subsequent treatments*

323

324 During ensiling the OPW lost weight (minimum $2.72\% \pm 0.81\%$ in DRY mode, maximum $3.15\% \pm$
325 0.27% for ENS mode) (Table 2). This weight loss was fast-until the 10th day and subsequently
326 slower (Figure 2).

327

328 **Table 2.** Main physico-chemical properties of the OPW (orange peel waste) subjected to different ensiling conditions and treatments.

329

OPW	Loss	TS		VS		
	(% on fresh weight)	(% on fresh weight) <i>initial</i>	(% on fresh weight) <i>final</i>	<i>initial</i> (% on TS)	<i>final</i> (% on raw biomass)	<i>final</i> (% on raw biomass)
RAW <i>(non-ensiled)</i>	-	17.8 ± 0.53	-	96.9 ± 0.01		
ENS	3.15 ± 0.27 a	17.8 ± 0.53	11.1 ± 0.28 ab	96.9 ± 0.01	94.4 ± 0.06 a	10.5 ± 0.26 a
WET	2.79 ± 0.09 a	14.2 ± 0.40*	9.2 ± 0.20 a	96.9 ± 0.01	94.6 ± 0.00 a	8.7 ± 0.19 b
DRY	2.72 ± 0.81 a	17.8 ± 0.53	13.1 ± 0.1 b	96.9 ± 0.01	94.6 ± 0.01 a	12.4 ± 0.09 c

330 Notes: different letters indicate significant differences according to t-test (at p < 0.05); TS = Total Solids; VS = Volatile Solids; ENS: naturally ensiled OPW; WET: OPW ensiled

331 in wet conditions; DRY: OPW ensiled in dry conditions; *water addition.

332 As expected, biogas ($> 95\% \text{ CO}_2$) was produced only in the first days of ensiling (Figure 1 - SI) In
333 fact, in this period, aerobic bacteria and yeasts were dominant, producing CO_2 through their
334 metabolism, mainly due to the air entrapped in the OPW pores, in accordance to [36,40].

335 The highest reduction in TS was measured for ENS samples ($-37.4\% \pm 3.44\%$) and the lowest for
336 DRY ($-26.2\% \pm 1.79\%$). VS reduced on average by only $2.4\% \pm 0.02\%$ (WET and DRY) – $2.6\% \pm$
337 0.05% (ENS) (Table 2). Also for TS and VS the parameters, the decrease was faster at the start of
338 ensiling and then tended to slow (Figure 2). This is in agreement with biogas production that was
339 quantitative only in the first days (Figure 2).

340 The initial COD of the OPW ($928 \pm 158 \text{ mg}\cdot\text{g}^{-1}$) did not noticeably change for the tested conditions
341 and treatments, with a maximum value ($994 \pm 95 \text{ mg}\cdot\text{g}^{-1}$) measured for ENS and a minimum of 936
342 $\pm 41 \text{ mg}\cdot\text{g}^{-1}$ for WET (Figure 2). The pH (initially 3.7 ± 0.0) was stable for ENS and WET and
343 lowered for DRY (3.3 ± 0.0) (Table 2). Generally, the pH evolution was not monotonic, but
344 fluctuated around the initial value with a slightly more noticeable variability detected for ENS
345 ensiling mode (Figure 2).

346 The liquid recovered by centrifugation from ENS and WET samples or collected at the reactor
347 bottom for DRY samples (passive drainage) contained amounts of lactic acid, while butyric, and
348 propionic acid have not been measured due to their low concentrations. Table 3 reports the lactic
349 acid concentrations in the liquid phase after 15 and 30 days of ensiling, respectively.

350

351

352

353

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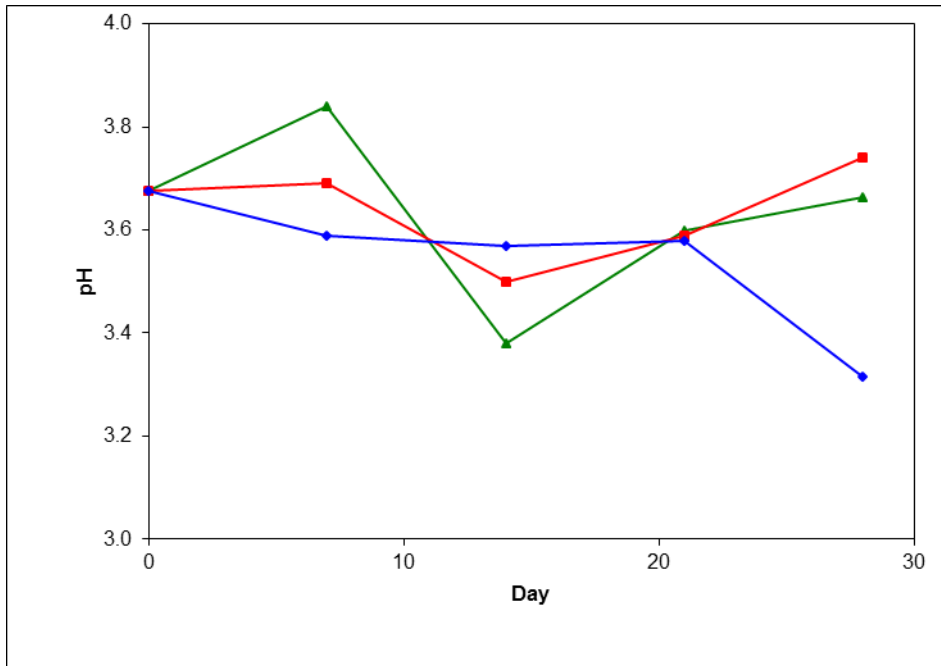
356 **Table 3.** Lactic acid concentration in the liquid phase separated by centrifugation (ENS and WET
357 ensiling modes) or by passive drainage (DRY ensiling mode).

358

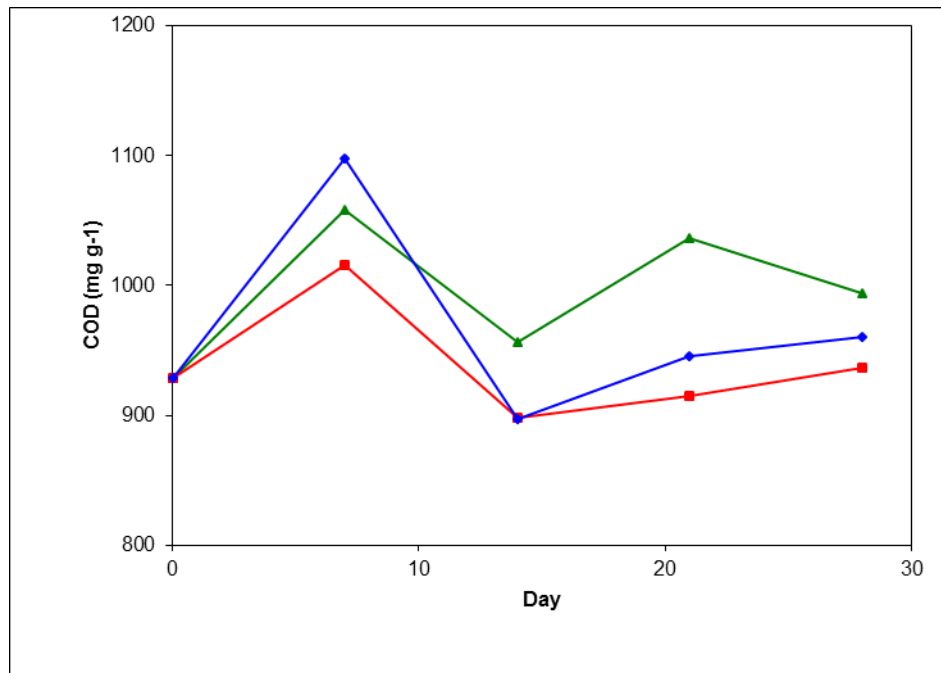
	Lactic acid (g/L)
ENS _{15 days}	1.8
WET _{15 days}	1.5
DRY _{15 days}	1.1
ENS _{30 days}	2.6
WET _{30 days}	1.8
DRY _{30 days}	1.6

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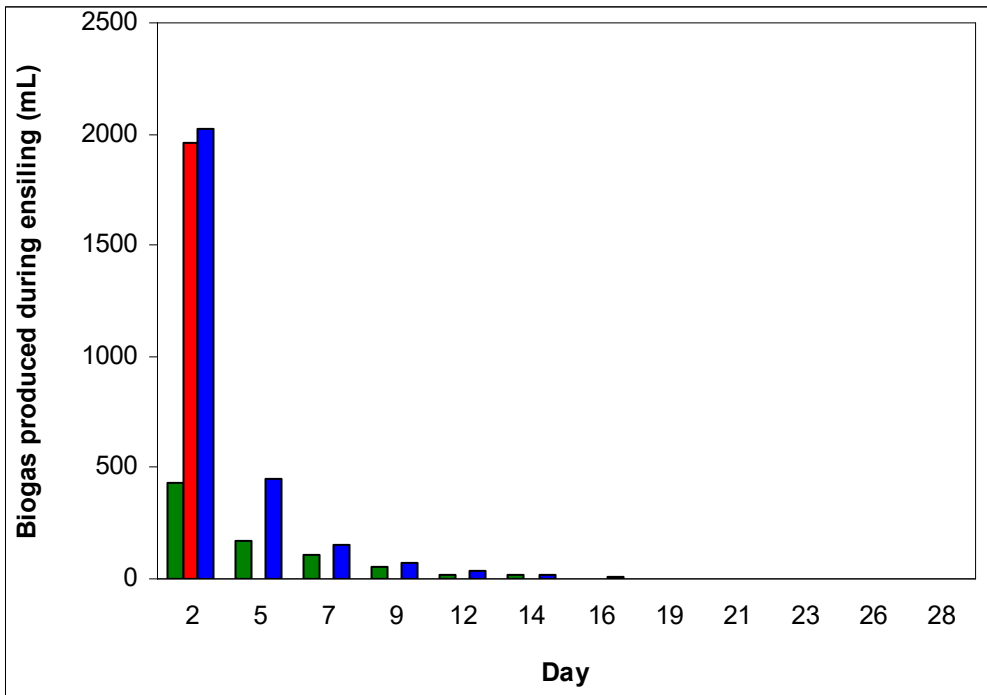
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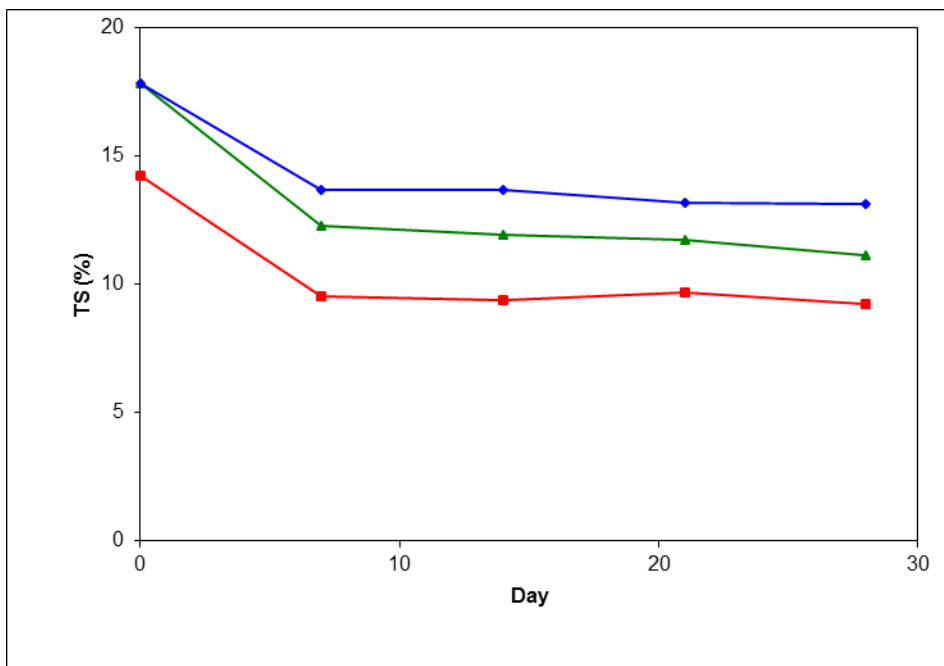
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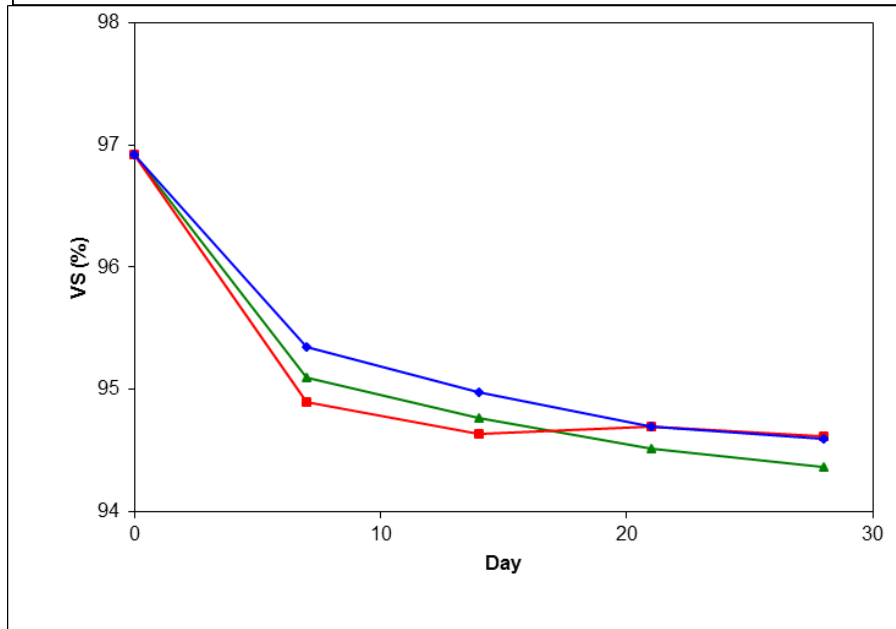
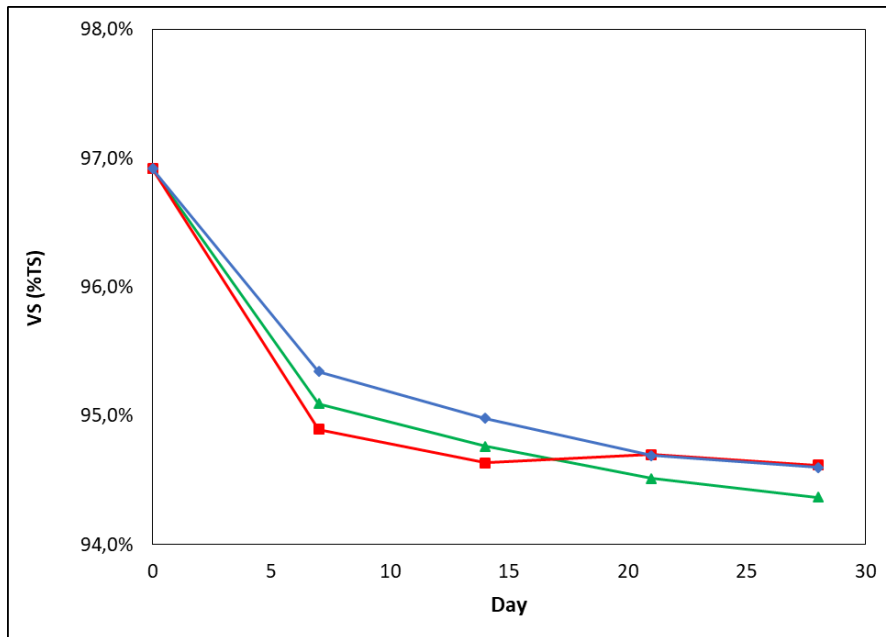


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371 **Figure 2.** Temporal evolution of the main physico-chemical properties of the OPW (orange peel
 372 waste) subjected to different ensiling conditions and treatments (ENS - naturally ensiled OPW;
 373 WET - OPW ensiled in wet conditions; DRY - OPW ensiled in dry conditions)

374 **Table 4.** Values of d-Limonene and residual content of VS in OPW subjected to subjected to
 375 different ensiling conditions and treatments

376

OPW	d-Limonene(mg/g)	VS/VS of raw OPW
RAW	0.55	1.00
ENS	0.14	0.59
WET	0.14	0.49
DRY	0.18	0.58
ENS+CEN	0.34	0.53
WET+CEN	0.25	0.41
DRY+CEN	0.28	0.54
ENS+CHEM	0.14	0.51
WET+CHEM	0.19	0.37
DRY+CHEM	0.16	0.50

377

378 Notes: VS = Volatile Solids; ENS: naturally ensiled OPW; WET: OPW ensiled in wet conditions; DRY: OPW ensiled
 379 in dry conditions; CEN = OPW subjected to centrifugation; CHEM = OPW subjected to centrifugation and chemical
 380 treatment with ethanol.

381

382 Compared to the raw biomass, ensiling reduced d-Limonene content of OPW by 67 (DRY) to 75%
 383 (WET and ENS conditions). The chemical treatment of ensiled OPW gave slightly lower d-
 384 Limonene contents only in for DRY+CHEM (-71% respect to -67% for DRY), while centrifugation
 385 of the ensiled OPW achieved the lowest decreases.

386 The reduction in VS content was in the range -41% (ENS) to -63% (WET and CHEM) with an
 387 average gradient CHEM (-54%) > CEN (-51%) > ENS/WET/DRY OPW (-45%) (Table 4).

388

389

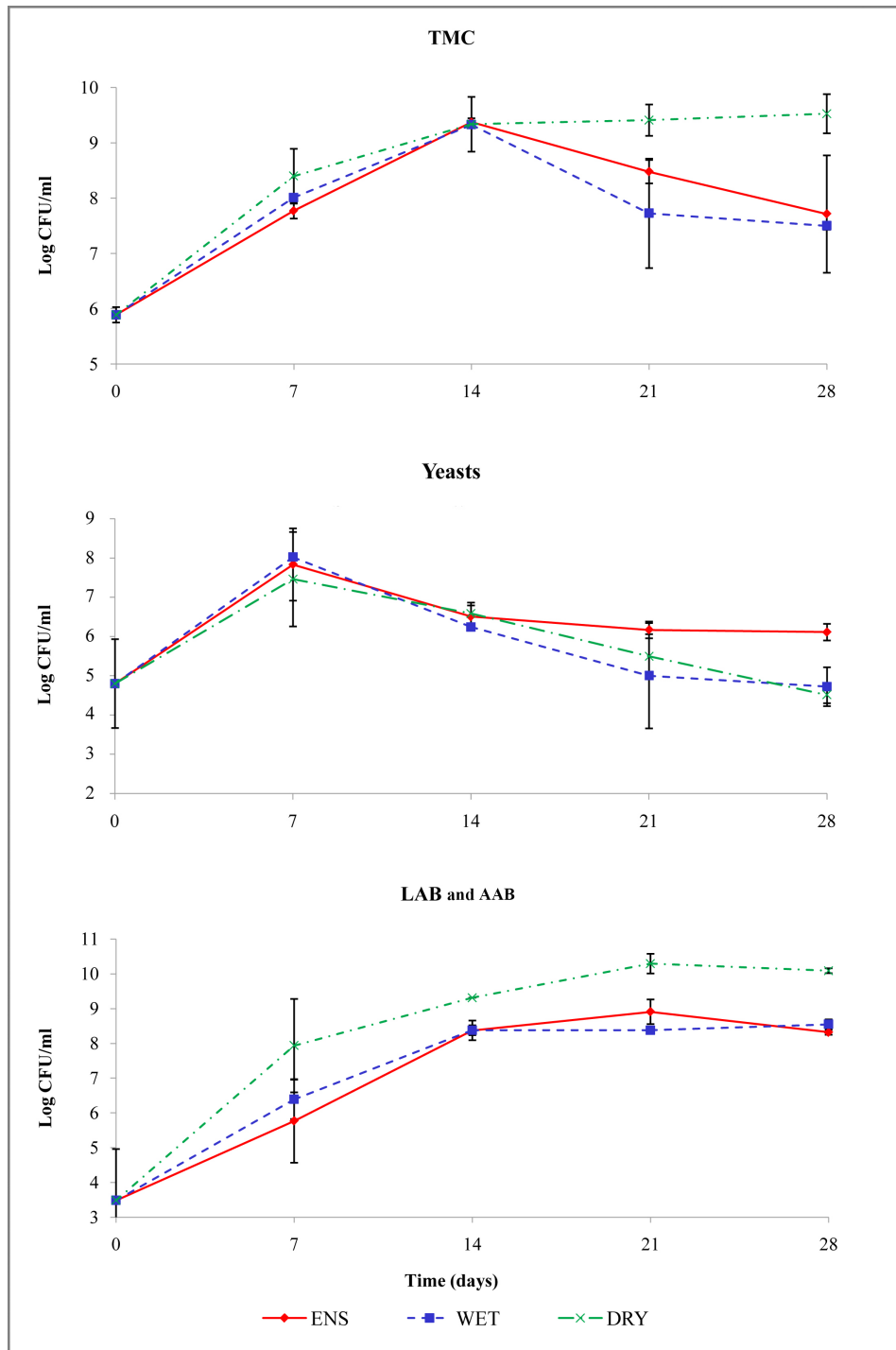
390 *3.2 Microbiological changes*

391

392 The microbial loads refer to raw OPW analysed before the start of ensiling and to OPW treated by
393 ENS, WET (solid material), and DRY methods (solid material and leachate). As regards the ENS
394 and WET OPW, the aerobic TMC loads gradually increased up to the maximum values at the 14th
395 day of ensilage. Then, the population decreased down to 7.50 and 7.71 Log CFU/mL, respectively,
396 after 28 days. The decrease was more marked in WET OPW compared to ENS samples. On the
397 contrary, the TMC loads of the leachate of DRY OPW always increased until 9.53 Log CFU/mL at
398 the end, but the rate of increase was higher until the 14th day and lower thereafter (Figure 3).

399 Acetic acid bacteria (AAB) were only counted in raw OPW, therefore at day 0. Then, the bacteria
400 detected were LAB. At the first stages of ensiling, 0 and 7 days, for all samples the load of yeasts
401 was higher compared to LAB. Subsequently, LAB were present in greater quantity than yeasts and
402 evolved by similar rates in all the tested ensiled OPWs (Figure 3). In WET OPW and leachate of
403 DRY OPW, yeasts evolved with similar trend throughout the process and, at the end (after 28 days)
404 yeast counts were lower (4.72 - 4.51 Log CFU/mL) than in ENS OPW (6.11 Log CFU/mL) while
405 LAB counts were higher in the leachate of DRY OPW and lower in ENS and WET samples (10.09
406 Log CFU/mL against 8.32 - 8.54 Log CFU/mL) (Figures 3).

407



408

409

410 **Figure 3.** Total microbial count (TMC), yeasts, as well as lactic (LAB) and acetic acid (AAB)
 411 bacteria counts of raw OPW (day 0) and OPW subjected to different ensiling conditions (ENS:
 412 naturally ensiled OPW; WET: OPW ensiled in wet conditions; DRY: leachage of OPW ensiled in
 413 dry conditions).

414

415 Concerning the solid fraction of the DRY ensiling, the microbial population observed was
416 negligible except for the yeasts at the 7th day (3.70 Log CFU/mL) (data not shown). This could be
417 due to a progressive loss of humidity as the liquid part flowed into the lower part of the fermenter.

418

419 3.3 Bacteria and yeast's identification

420

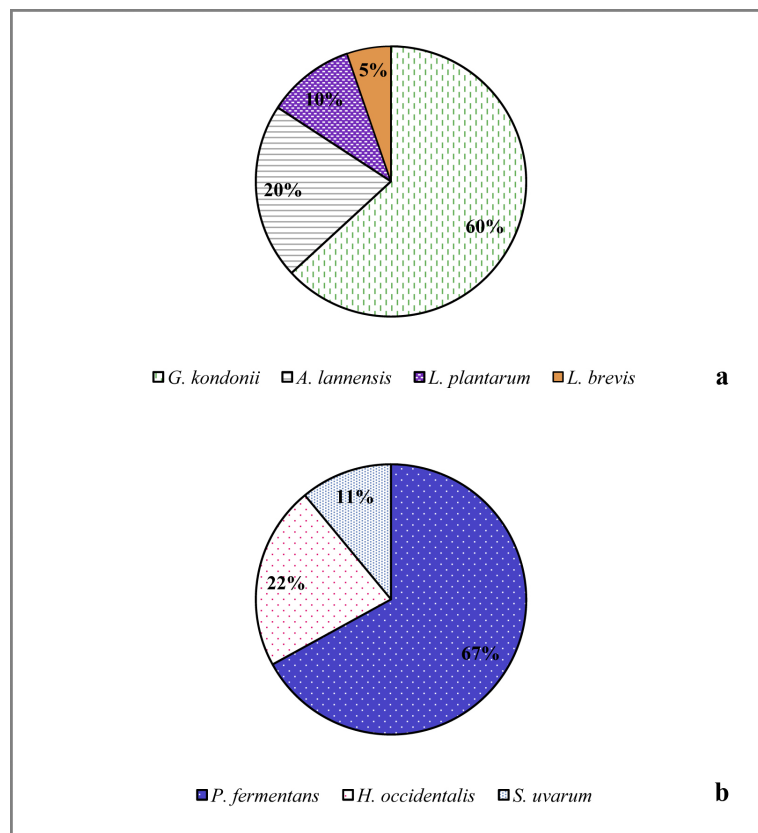
421 87% of the total population of the bacteria isolated was catalase-negative and Gram positive, while
422 the remaining 13% was catalase-positive and Gram negative. Eight patterns of ARDRA profiles
423 were observed. Bacteria were identified as *L. plantarum*, *Lactobacillus brevis*, *Gluconobacter*
424 *kondonii*, *Lactobacillus suebicus*, *Leuconostoc pseudomesenteroides*, *Lactobacillus*
425 *paracollinoides*, *Leuconostoc citreum*, and *Asaia lannensis*. LAB species of *L. plantarum* and *L.*
426 *brevis* were present in the OPW at day 0 together with AAB species of *G. kondonii* and *A. lannensis*
427 (Figure 4). These AAB dominated the matrix at this stage, consistent with the presence of oxygen.
428 The AAB were not recovered from all the samples throughout the ensilage.

429 Figure 5 reports the species distribution detected in the samples. At the 7th day, all the samples
430 harboured *L. plantarum*. The leachate of DRY OPW and the WET OPW favoured the growth of *L.*
431 *citreum*, *L. pseudomesenteroides* characterised the leachate of DRY OPW. After 14 days, *L. brevis*
432 was detected in all the samples, while *L. suebicus*, *L. pseudomesenteroides*, and *L. paracollinoides*
433 were present in ENS, leachate of DRY OPW and WET OPW, respectively. After 21 days, ENS and
434 WET samples of OPW were characterised by *L. plantarum*, *L. brevis*, and *L. suebicus*, while the
435 leachate of DRY OPW contained *L. brevis* and *L. plantarum*. At the end of the ensilage process,
436 ENS and WET OPW showed LAB composition similar to the population on the 21st day, while the
437 leachate of DRY OPW was dominated by *L. plantarum*.

438 Eleven patterns of RFLP profiles were observed. Yeasts were identified as *Pichia fermentans*,
439 *Saccharomyces cerevisiae*, *Kregervanrija fluxum*, *Saccharomyces uvarum*, *Pichia*
440 *membranifaciens*, *Hanseniaspora occidentalis*, *Pichia kudriavzevii*, *Pichia occidentalis*,

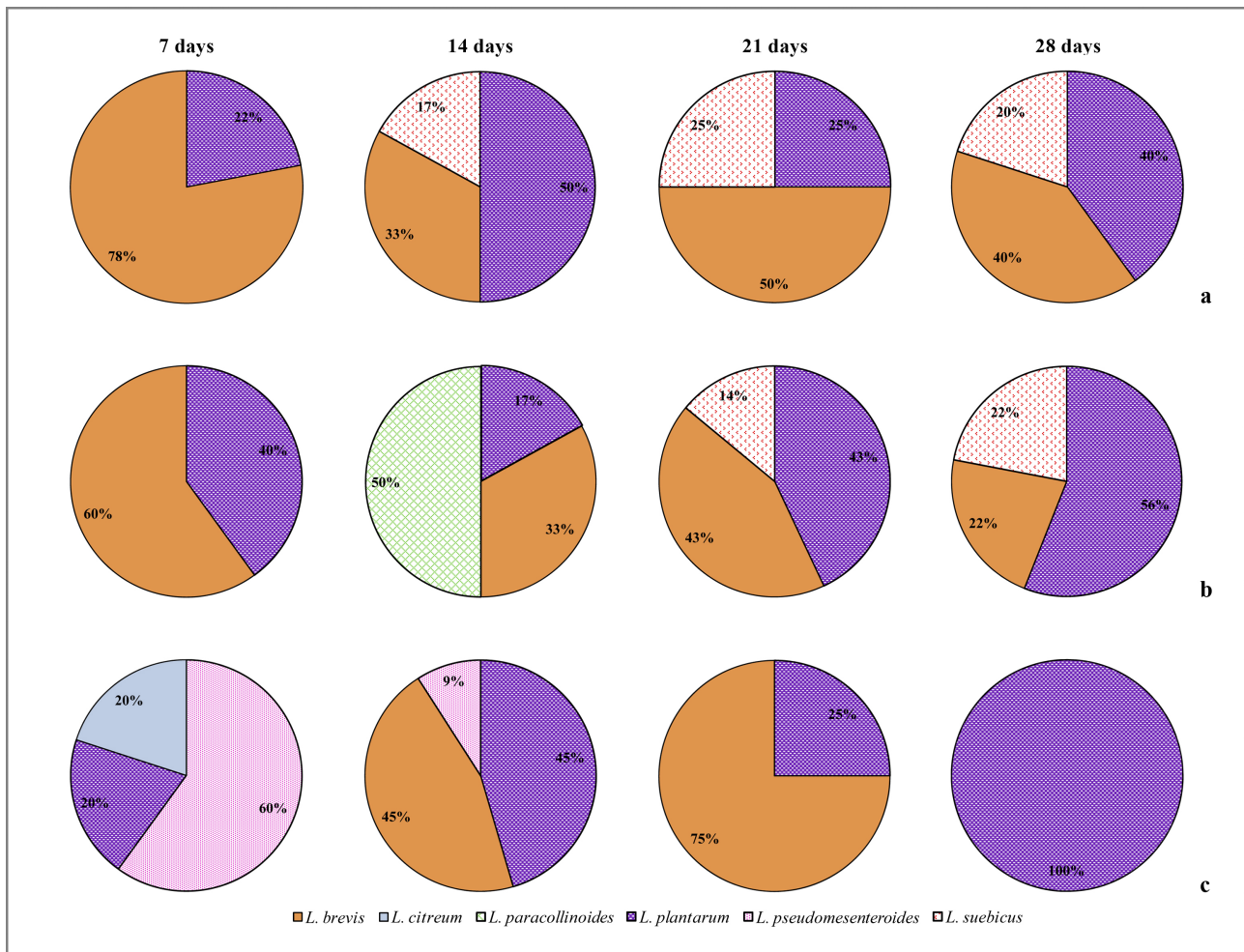
441 *Hanseniaspora nectarophila*, *Kazachstania barnettii*, and *Torulaspora delbrueckii*. *P. fermentans*,
 442 *H. occidentalis*, and *S. uvarum* were detected in OPW at day 0 (Figure 4). *P. fermentans* and *S.*
 443 *cerevisiae* were isolated from all the samples throughout the ensilage. On the 7th day, *H.*
 444 *occidentalis* and *H. nectarophila* were found in WET OPW and leachate of DRY OPW,
 445 respectively. In the middle stage of ensilage, ENS OPW was characterised also by *K. barnettii* and
 446 *K. fluxum*, while the leachate of DRY OPW by *Saccharomyces* sp. and *T. delbrueckii*. At the end of
 447 the ensilage, all the samples contained *K. fluxum* and *Pichia* spp. (Figure 6).
 448 As regards the representative strains of LABs and yeasts sequenced, the percentage of similarity,
 449 and the accession numbers of the closest relative by BLAST, reported in Table 5, the sequences
 450 with a percentage homology of 97% or higher were considered to belong to the same species,
 451 according to [42].

452



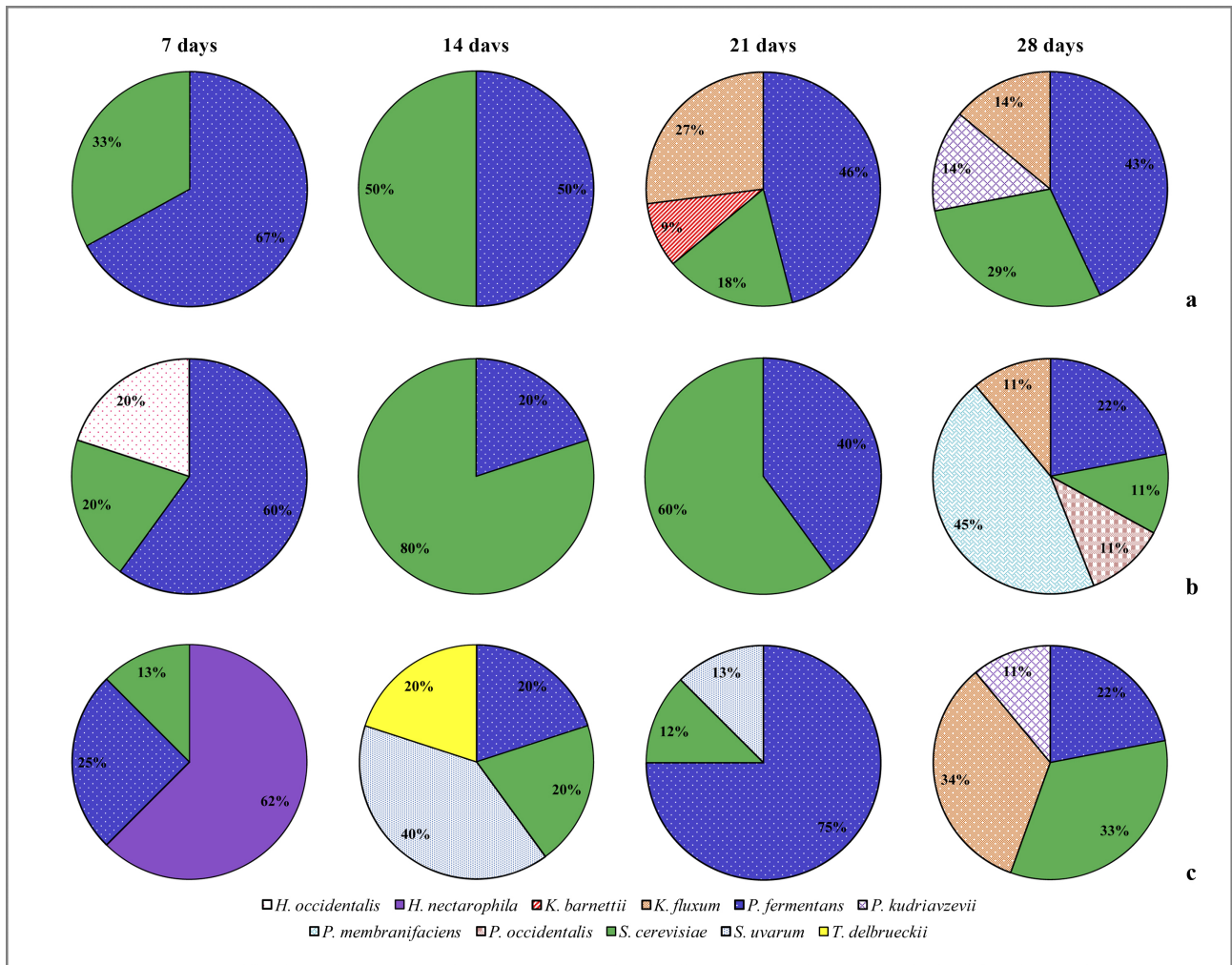
453

454 **Figure 4.** Number (in % on the total) of LAB and AAB (a) and yeast (b) species recovered from
 455 raw orange peel waste (OPW) (day 0).



457

458 **Figure 5.** Number (in % on the total) of bacteria recovered from ENS (a), WET (b), and leachate of
 459 DRY (c) orange peel waste (OPW) throughout ensilage.



460

461 **Figure 6.** Number (in % of the total) of yeasts recovered from ENS (a), WET (b), and leachate of
 462 DRY (c) orange peel waste (OPW) throughout ensilage.

463

464 *3.4 BMP test results*

465

466 In general, all the values of the net specific methane production for OPW under the tested
 467 conditions and treatments were significantly higher compared to the value measured for the raw
 468 biomass ($415 \pm 26.4 \text{ NmL} \cdot \text{gVS}^{-1}$). In more detail, the lowest increases in the methane yield was
 469 measured for the centrifuged OPW (on average +47%), while the highest production was detected
 470 for WET OPW subjected to the chemical treatment with ethanol and then centrifuged (+86%
 471 compared to raw OPW) (Figure 7). The differences were significant both for the conditions and the
 472 treatments and the same was for the interaction condition x treatment.

473 AD process regularly evolved in time in all the BMPs, as shown by the monotone cumulated
 474 methane production. The chemical treatments with ethanol - specifically the tests carried out on
 475 ENS+CHEM, WET+CHEM and DRY+CHEM OPW - were the exceptions. In these tests, the AD
 476 process were slower at the earlier stages (until the 10th-20th day) (Figure 8, h, i, l). Other slight
 477 evidences of AD inhibition were detected for one reactor fed with ENS OPW (Figure 8b) and for
 478 the reactors fed with ENS+CEN, WET+CEN and DRY+CEN OPW (Figure 8e, f, g).

479

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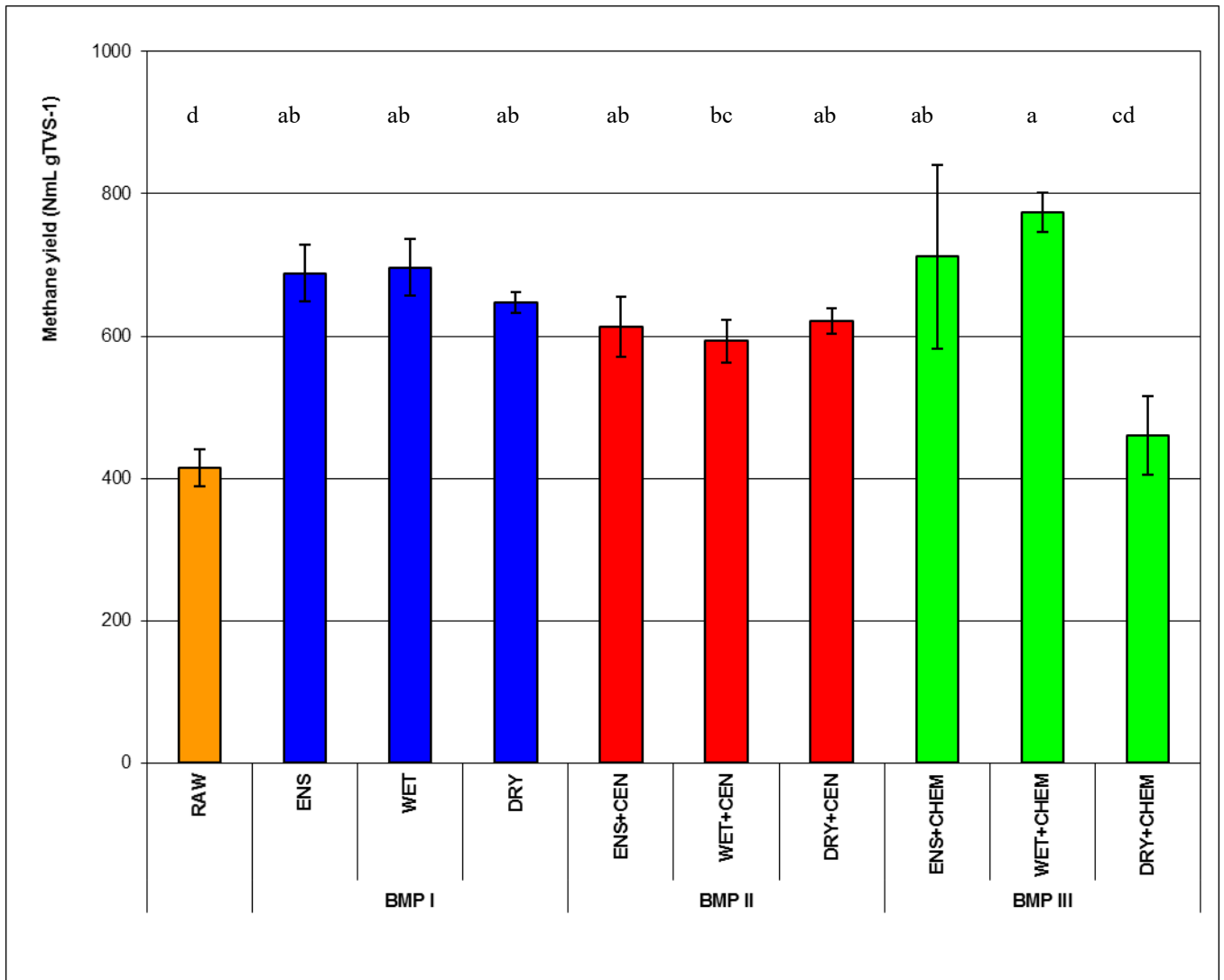
481 **Table 5.** Representative strains of LABs and yeasts sequenced together with the percentage of
 482 similarity and the accession numbers of the closest relative by BLAST.

483

	Strains	Species	Similarity (% , accession no. of the closest relative by Blast)
Yeasts	PY 1	<i>Saccharomyces uvarum</i>	100% - KY109468.1
	PY 12	<i>Saccharomyces cerevisiae</i>	100% - NG_042623.1
	PY 21	<i>Hanseniaspora nectarophila</i>	97% - NG_055397.1
	PY 2	<i>Pichia fermentans</i>	99.82% - KY108804.1
	PY 82	<i>Pichia occidentalis</i>	100% - KY108912.1
	PY 76	<i>Pichia kudriavzevii</i>	100% - KY108786.1
	PY 80	<i>Pichia membranifaciens</i>	100% - KY108889.1
	PY 24	<i>Kregervanrija fluxum</i>	100% - KY108172.1
	PY 4	<i>Hanseniaspora occidentalis</i>	100% - NG_055416.1
	PY 56	<i>Kazachstania barnetii</i>	100% - KY107903.1
	PY 44	<i>Torulaspota delbrueckii</i>	100% - NG_058413.1
Bacteria	PB 22	<i>Lactobacillus plantartum</i>	100% - NR_115605.1
	PB 31	<i>Lactobacillus brevis</i>	100% - NR_116238.1
	PB 47	<i>Lactobacillus suebicus</i>	100% - NR_114977.1
	PB 30	<i>Lactobacillus paracollinoides</i>	100% - NR_042322.1
	PB 12	<i>Leuconostoc citreum</i>	100% - NR_041727.1
	PB 24	<i>Leuconostoc pseudomesenteroides</i>	99.89% - NR_109004.1
	PB 4	<i>Asaia lannensis</i>	100% - NR_114144.1
	PB 5	<i>Gluconobacter kondonii</i>	100% - NR_104680.1

484

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487 Note: different lowercase letters indicate significant differences (at $p < 0.05$).

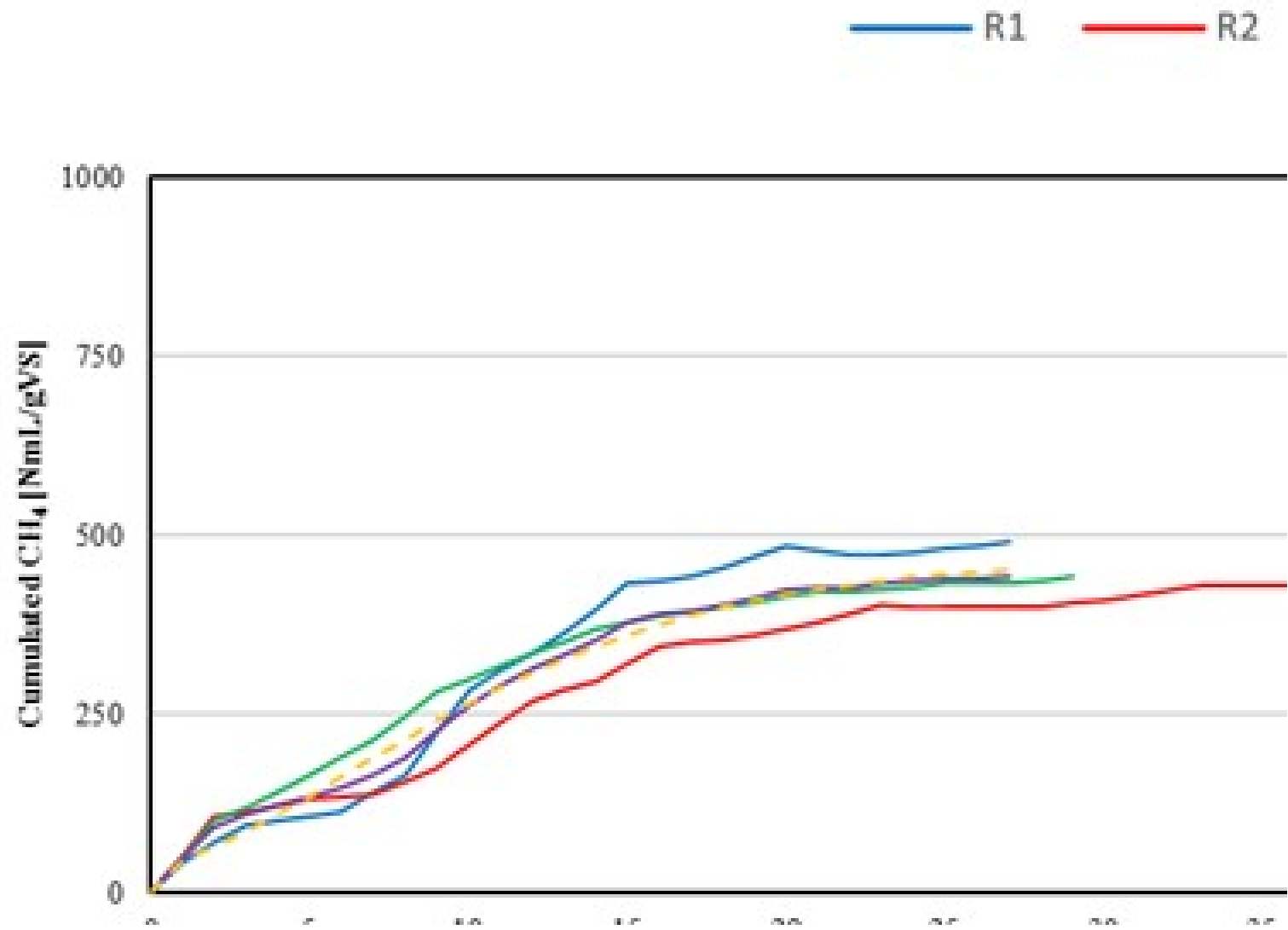
488 **Figure 7.** Cumulated net specific methane yields of the BMP I, II and III batch tests.

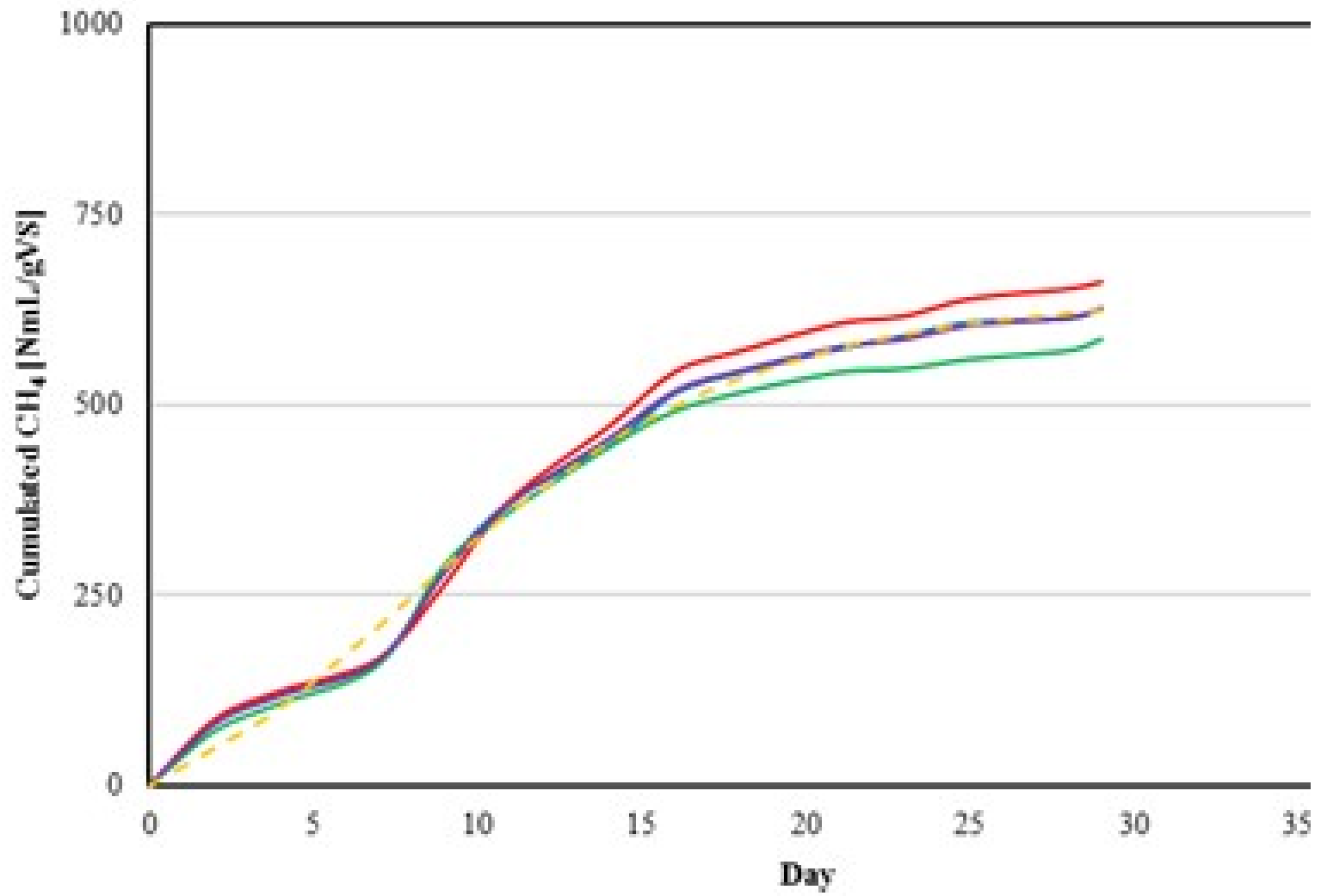
489

490 Gompertz equation fitted well the experimental data of all BMP tests ($r^2 > 0.99$) (Table 6).

491 Table 6 reports a comparison between two options: (i) the digestion of a given amount of raw OPW
 492 without pre-treatments; and (ii) ensiling (with or without other treatments, such as ethanol addition
 493 and centrifugation) before digestion. In more detail, the ratio CH_4/CH_{4raw} OPW, which considers the
 494 VS losses occurring during the different treatments before AD, shows that the VS losses are not
 495 balanced by a corresponding increase of the specific methane production combining all the
 496 processes (ensiling, chemical treatment and centrifugation); in fact, the methane production is
 497 between 55 and 89% of the methane production without any pre-treatment (that is, by the direct

498 digestion of OPW). This theoretical production can not be achieved because of the reasons already
499 explained.





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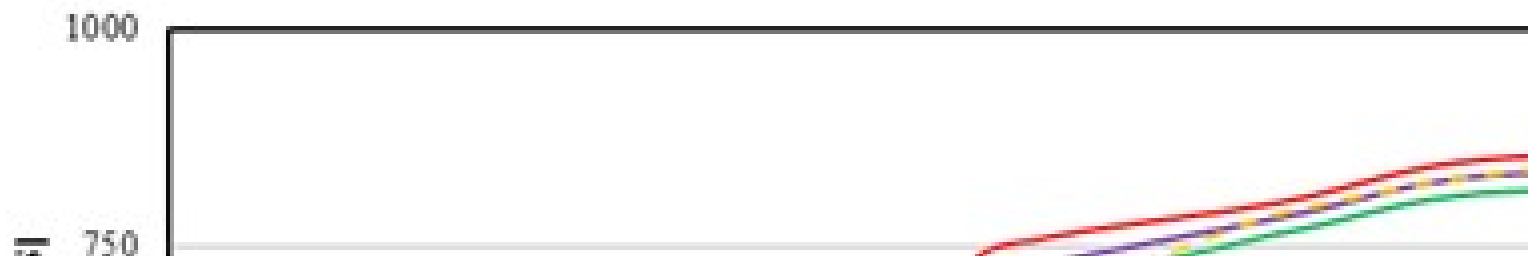


Figure 8. Cumulated net specific methane yields and interpolating Gompertz equation for the orange peel waste (OPW) under the tested conditions and treatments: a. raw OPW; b. ENS OPW; c. WET OPW; d. DRY OPW; e. ENS+CEN OPW; f. WET+CEN OPW; g. DRY+CEN OPW; h. ENS+CHEM OPW; i. WET+CHEM OPW; l. DRY+CHEM OPW).

507 **Table 6.** Methane net specific production compared to the theoretical value yielded by the anaerobic digestion of raw orange peel waste (OPW)
 508 together with the parameters of the interpolating Gompertz equation.

509

OPW	P (NmL·g _{VS} ⁻¹)	λ (d)	R_m (NmL·(g _{VS} ·d) ⁻¹)	r²	CH₄/CH_{4raw} OPW
RAW	0.47	0.00	0.027	0.996	1.00
ENS	0.69	0.00	0.051	0.997	0.89
WET	0.68	0.00	0.067	0.998	0.75
DRY	0.64	0.00	0.057	0.999	0.82
ENS+CEN	0.64	1.62	0.039	0.997	0.73
WET+CEN	0.61	0.83	0.041	0.999	0.55
DRY+CEN	0.63	0.74	0.041	0.999	0.76
ENS+CHEM	0.72	0.00	0.058	0.994	0.84
WET+CHEM	0.88	5.34	0.044	0.992	0.67
DRY+CHEM	0.93	12.57	0.032	0.990	0.80

510 Notes: P, R_m and λ = parameters of Gompertz equation.

511

512 **4. Discussions**

513 The changes in the OPW (i.e. weight loss, variations of TS and VS, and biogas production),
514 mainly occurring in the first week of the process, were quite similar among the three ensiling
515 conditions and the differences were not significant at all, with the exception of TS variations.
516 The latter was determined by the very different conditions between WET and DRY ensiling
517 mode in terms of water management. Weight loss and biogas production are coherent as
518 observed in previous literature [20,27]. Also the pH, noticeably acid as usually recorded for
519 raw OPW [1,19], was kept basically stable during the process.

520 The removal of d-Limonene was efficient under all the tested ensiling conditions, which
521 confirms the viability of its removal before OPW anaerobic digestion to increase the methane
522 yield of this process.

523 Presumably, during ensiling, OPW decomposition allows the d-Limonene-containing sacs in
524 the flavedo to rupture, and the simultaneous biogas production enhances its stripping. This
525 process is confirmed at a the sensorial level by the strong orange smell during biogas venting.

526 The treatment with ethanol did not increase the d-Limonene removal rate compared to the
527 untreated biomass (on average by 70% against 72%), whereas simple centrifugation reduced
528 this rate by only 47%. These results are obviously influenced by a number of factors (specific
529 cultivar and ripening stage of oranges, type of processing, ensiling conditions), in order to
530 confirm these results, experiments at a larger scale would be beneficial.

531 The low efficacy of the OPW chemical treatment may be explained by the scarce suitability of
532 ethanol for d-Limonene leaching (despite its biodegradability, which suggested its use for this
533 scope) for the chemical treatment and the low solubility of d-Limonene in water and the high
534 affinity of the solid compounds of OPW for centrifugation. With regards to the latter, in fact,
535 d-Limonene concentration was higher in the centrifuged OPW compared to the simply ensiled
536 biomass.

537 In terms of residual VS content after ensiling and treatments, natural ensiling, allowing the
538 minimum removal (on average 45% of the initial content against 51% of chemical treatments
539 and 54% of the centrifugation), assures the lowest loss of VS and thus, potentially, a more
540 efficient preservation of the bio-methane potential production of OPW.

541 Therefore, this study suggests using natural ensiling to decrease the d-Limonene loads in the
542 substrate without further treatments, since this choice maximises the removal and minimizes
543 complexity and cost of the processing. Wet conditions are not advised, because a higher
544 reduction of VS content is achieved, which may determine lower bio-methane yields. For dry
545 conditions the overall balance of the ensiling process would be more favourable if a
546 valorisation option (e.g. for bio-ethanol production or as an additive to wastewater treatment
547 plants for denitrification) is found for the leachate extracted from OPW.

548 Centrifugation is not advisable since it causes an additional loss of substrate (e.g. soluble
549 sugars, lactic acid) through the discarded liquid and does not improve the efficiency of d-
550 Limonene removal.

551 Under the microbiological approach, LAB population increases throughout the process, as
552 expected considering the type of fermentation characterizing the ensiling. This increase
553 corresponded to a decrease in yeast population, observed with a more noticeable trend in
554 leachate of ensiling under dry conditions than in the others. As facultative anaerobes, yeasts
555 were not suppressed during ensilage. Despite the presence of EO in the matrix, both yeasts
556 and LABs grew and persisted to the end of the ensilage. Most likely, the autochthonous
557 microorganisms are accustomed to the OPW environment confirming a certain adaptation as
558 reported for the treatment of citrus processing wastewater in aerated ponds [43,44].

559 The analysis of the organic acids confirmed that LAB population was dominant, since butyric
560 acid produced by Clostridia was absent [25]; the very low initial pH presumably helped to
561 prevent their presence in the reactors.

562 The study confirms that the d-Limonene removal, the particle size reduction, and the biomass
563 homogenization and fermentation during ensiling and/or the subsequent treatments
564 significantly improve the specific (that is, the methane production per unit VS added)
565 efficiency of the OPW energy conversion by AD. As a matter of fact, higher methane yields
566 were measured for ensiled OPW (close to upper limit of the literature range [13,47]),
567 compared to the raw substrate, which is close to the literature average [1,3,12]. In the case of
568 the OPW subjected to the chemical treatment (ENS+CHEM, WET+CHEM), the
569 biodegradation of residual ethanol [48] presumably enhanced the methane yield, since ethanol
570 can be an additional carbon source for microorganisms. In the other cases (ENS, WET, DRY,
571 ENS+CEN, WET+CEN, DRY+CEN), it is possible that the high methane yields can be
572 ascribed to the peculiar characteristics of the inoculum. The latter was taken from a full-scale
573 anaerobic digester, where fresh and ensiled OPW is routinely used as co-substrate during the
574 orange processing season. For this reason, the inoculum is adapted to the tested substrate,
575 increasing methane yields [28].

576 The increase in methane yield partially compensated for the reduction in VS during ensiling.
577 In general, the process regularly evolved, that is, no evidence of partial inhibition was
578 observed, except for the reactors fed with centrifuged OPW and, especially, for the reactors
579 with chemically treated OPW. In the first case the slight inhibition was presumably due to the
580 higher residual d-Limonene content (over the inhibition limit of the anaerobic process) of
581 centrifuged OPW compared to the other treatments, Table 6). However, this partial inhibition
582 played a lower effect on methane yields compared to other BMPs of literature [7,18]. For the
583 substrates treated with ethanol the inhibition was more evident; it was due to the adaptation of
584 the microbial consortium to this compound [48].

585 Among the tested BMPs, the treatment with ethanol gave the highest methane yields but also
586 caused an irregular AD process, while the simple centrifugation of OPW was not efficient

587 compared to the other techniques. . The analysis of the parameters estimated for Gompertz
588 equation confirms that ensiling significantly improves the net specific methane yield and in
589 many cases also the degradation rates, as shown by the highest R_m estimated for the naturally
590 ensiled OPW and the biomass subjected to chemical treatment after natural ensiling).
591 The best performing treatment is ENS, which reduces methane production by 11% compared
592 to to the AD of the raw OPW, ENS+CHEM (-16%) and DRY (-18%).
593 The methane production of OPW digested after ensiling (natural or subjected to the treatments
594 of centrifugation or solvent extraction) is 55 – 89% of the production of the same quantity of
595 raw OPW in AD under the same process conditions (Table 6).

596

597

598 **5. Conclusions**

599

600 The possibility to increase the viability of the anaerobic digestion of OPW through ensiling
601 and subsequent treatments has been explored in this study. The laboratory tests have
602 confirmed that biomass storage allows a high (over 70%) d-Limonene removal but with heavy
603 significant reductions (41 – 63% compared to the raw OPW) of the content in volatile solids
604 (to be degraded during the energy conversion process). ENS and DRY ensiling modes without
605 subsequent treatments appear to be the most suitable techniques since they minimize the
606 reduction in CH_4 production of the overall process.

607 LAB and yeast species associated with ensiled OPW were assessed for the first time. The
608 microbiological population showed high biodiversity that can be further explored with the aim
609 of applying specific microbial strains as ensiling inocula to try to further accelerate the
610 process with a subsequent better preservation of the methane potential.

611 Further research is needed to select more efficient biodegradable solvents for improving d-
612 Limonene removal from ensiled OPW and to suggest additional valorisation opportunities for
613 the leachate released from DRY ensiling.

614

615 **Acknowledgement**

616

617 We thank the journal Editor and three anonymous Reviewers, whose comments helped
618 improve this manuscript.

619

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SUPPLEMENTARY INFORMATION

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Figure 1 - SI – Images of Orange Peel Waste (OPW) ensiled in different conditions

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in the laboratory tests.