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1	INVESTIGATION OF FOOD WASTE VALORIZATION THROUGH SEQUENTIAL
2	LACTIC ACID FERMENTATIVE PRODUCTION AND ANAEROBIC DIGESTION
3	OF FERMENTATION RESIDUES. PART I: TECHNICAL ASSESSMENT
4	
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20 Abstract

This work concerns the investigation of the sequential production of lactic acid (LA) and 21 22 biogas from food waste (FW). LA was produced from FW using a Streptococcus sp. strain via simultaneous saccharification and fermentation (SSF) and separate enzymatic hydrolysis and 23 fermentation (SHF). Via SHF a yield of 0.33 g_{LA}/g_{FW} (productivity 3.38 $g_{LA}/L^{-}h$) and via SSF 24 0.29 g_{LA}/g_{FW} (productivity 2.08 g_{LA}/L^{-h}) was obtained. Fermentation residues and FW 25 underwent anaerobic digestion (3 wt% TS). Biogas yields were 0.71, 0.74 and 0.90 Nm³/kg_{VS} 26 for FW and residues from SSF and SHF respectively. The innovation of the approach is 27 considering the conversion of FW into two different products through a biorefinery concept, 28 therefore making economically feasible LA production and valorising its fermentative 29 residues. Finally, a mass balance of three different outlines with the aim to assess the amount 30 of LA and biogas that may be generated within different scenarios is presented. 31

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34 Keywords: biorefinery, fermentation, lactic acid, enzymatic hydrolysis, biogas

36 **1. Introduction**

Biowaste generation in EU was estimated at 94 Mt for 2015 and current treatment options include landfilling, incineration, mechanical-biological treatment (MBT), composting and anaerobic digestion (EU, 2010). Food waste (FW) from households, restaurants, caterers and retail premises represents an important fraction of biowaste. FW is globally one of the most severe environmental, social and economic problems of developed and developing countries, accounting for over one billion tonnes produced every year (Gustavsson, 2011).

Currently the main environmental threat from organic waste is methane production from such 43 waste decomposing in landfills. Before the adoption of the Landfill Directive 1999/31/EC 44 methane emissions from landfills accounted for 30% of the global anthropogenic emissions of 45 methane into the atmosphere (COM, 1996). Landfill Directive 1999/31/EC obliges member 46 states to reduce the amount of landfilled biodegradable municipal waste to 35% of 1995 levels 47 48 by 2016, however it doesn't prescribe specific treatment options for the diverted waste. The response of EU member states since mid 1990s was the implementation of MBT, anaerobic 49 digestion and composting processes. However 20 years later, it is mandatory to improve the 50 management of biowaste by supporting technical solutions that are able to generate added 51 value products. 52

The composition of FW is heterogeneous, being made of (w/w) 30-60 % starch, 5-10 % proteins and 10-40 % lipids (Pleissner et al., 2013). Hence, it represents an interesting feedstock for biorefinery processes. FW biological valorisation is not only an environmentally friendly waste treatment option, but it is also a benefit to the bio-based economy since valuable waste material can be employed instead of expensive raw substrates.

Lactic acid (LA) has many applications in the food and beverage sector as well as in the pharmaceutical and chemical industries, and its polymerisation gives origin to the biodegradable polymer poly(lactic acid) (PLA) (Abdel-Rahman et al, 2016). The main paths

of fermentative production of LA from FW are separate enzymatic hydrolysis and 61 62 fermentation (SHF), and simultaneous saccharification and fermentation (SSF) (see Table 1). 63 While in SSF enzymatic hydrolysis and fermentation are performed in a single reactor (with uniform temperature and pH conditions), SHF foresees two separate phases, and thus allows 64 65 the application of optimal temperature and pH conditions for each process and the use of an acid or enzyme in the hydrolysis step, which may be highly effective for complex substrates. 66 SSF in comparison with SHF showed shorter processing time, reduced substrate/product 67 inhibition and lower energy and plant costs (Castillo Martinez et al., 2013; Abdel-Rahman et 68 al., 2013). SSF was either carried out by enzymes added with the inoculum (Wang et al., 69 2016), using a single microbial strain (Pleissner et al., 2017) or an indigenous microbial 70 consortium (Kim et al., 2016; Tang et al., 2016). 71

The aim of this study is to investigate the technical, economic and environmental feasibility 72 of the sequential fermentative production of LA and biogas from FW using either SHF or 73 SSF. LA and biogas production can be carried out using mature technologies, however, the 74 two options are usually considered separately. Fermentative production of LA from FW was 75 already proven to be feasible (Pleissner et al., 2017; Pleissner et al., 2015a and 2015b; Kwan 76 77 et al., 2015; Li et al., 2015); however its main drawbacks are the high process costs, necessary 78 to achieve a marketable LA, and a relevant amount of fermentative residues to be managed. LA production costs include: sterilization, fermentation and downstream processes. The most 79 expensive cost items are sterilization and downstream processes, which represent up to 41% 80 of the conventional fermentation process (Wang et al., 2015) and 1.57-1.62 €/kg_{LA} (Joglekar 81 et al, 2006). Sterilization and downstream processes are strictly recommended in order to 82 achieve a LA quality that is commercially competitive, since food grade purity and 83 84 pharmaceutical plastic grade purity are 80% and 90%, respectively (Vijayakumar, 2007). LA fermentation costs vary in the range of 0.72-1.13 \notin /kg_{LA} (Wang et al, 2015), while market value of LA is 1.36 \notin /kg_{LA} (ICIS, 2016).

AD of FW has been widely explored (see Table 2), and it has been implemented on full scale for the last decades, in agreement with waste management hierarchy and EU policy about organic waste management. Biogas yields observed for FW were 0.26-0.63 m³/kg_{VS} (Fantozzi et al., 2011; Pavi et al., 2017; Alibardi and Cossu, 2015; Kafle et al., 2013; Dinuccio et al., 2010; Gunaseelan et al., 2009), while methane yields were 0.15-0.25 m³/kg_{VS} for fruit pulp (Gali et al., 2009) and 0.26-1.4 m³_{methane}/kg_{VS} for mixed FW (Labatut et al., 2011; Elbeshbishy et al., 2012; Maya-Altamira et al., 2008).

Plant size that makes AD profitable ranks at 50-100 kWe and investment costs vary between
3000-5000 € and 6000-7000 € for plant sizes of around 500-1000 kWe and 50-100 kWe,
respectively (Insabato, 2013). Electric energy has a current value of 0.10 €/kWh and thermal
energy of 0.105 €/kWh (Eurostat, 2016).

The novelty of the approach consists of taking into account the conversion of FW into two different high value products through a biorefinery concept, therefore making at the same time economically feasible LA production and solving the issue of fermentative residues valorization. This approach is consistent with EU strategy about circular economy; moreover, industrial biotechnology belongs to the Key Enabling Technologies (KET), whose development, exploitation and implementation into the development of marketable goods and services are among priority action lines of European industrial policy.

The present research, concerning the overall investigation of the feasibility of the proposed
 biorefinery chain, is structured in three parts. Part I (this study) covers the technical issues,
 while the economic and environmental assessments will be respectively discussed in parts II
 and III. Considering LA fermentation, this study describes SHF while the details about SSF

process are given elsewhere (Pleissner et al., 2017). Nevertheless, AD tests performed on
SHF and SSF fermentation residues, as well as on FW, are here fully taken into account.

Finally, a mass balance was evaluated for three different process outlines, with the aim to assess the amount of LA and biogas that may be generated considering different scenarios. In detail, LA fermentation through SHF or SSF (Scenario 1), biogas production from FW anaerobic digestion (scenario 2) and sequential LA fermentation and AD of fermentation residues (scenario 3) were discussed and compared.

116

117 2. Material and Methods

118 **2.1 Food waste**

FW, made of noodles, potatoes, vegetables, rice, fruits, meat and sauce, was collected daily from the canteen of Leibniz Institute for Agricultural Engineering and Bioeconomy Potsdam for 15 days in July 2015 (141 kg in total). Immediately after collection, FW was blended through a kitchen blender and stored at -20°C. At the end of the sampling period, all FW blends were pooled and homogenised. FW amounts employed in all tests, as well as glassware, were autoclaved at 121°C for 15 minutes before use to exclude the presence of autochthon microorganisms competing with the ones specifically inoculated for the study.

126

127 **2.2 Microorganisms**

A mesophilic *Streptococcus* sp. strain A620 (internal label), isolated from tapioca starch, was employed in LA fermentations. The strain was classified by the German Collection of Microorganisms and Cell Cultures (DSMZ Braunschweig, Germany) and is available at the Leibniz Institute for Agricultural Engineering and Bioeconomy Potsdam. The strain was cultured in 300 mL flasks, containing 60 mL of MRS broth (Merck, Germany) and 0.67 g Everzit Dol (Evers, Germany) dolomite as buffer. Incubation occurred at 35°C for 24 hours. The initial pH in all flasks was equal to 6.0. Flasks were shaken at 100 rpm in an orbitalshaker.

The microbial consortium used as inoculum for anaerobic digestion tests was supplied from a
 mesophilic anaerobic digester at Leibniz Institute for Agricultural Engineering and
 Bioeconomy Potsdam. It consisted of 3.2 % (w/w) total solids (TS) and 54.4 % (w/w) volatile
 solids (VS). The pH was 7.8.

140

141 2.3 Enzymatic hydrolysis

Enzymatic hydrolysis tests were carried out without repetitions in presence of 1 L FW in a 2 142 L BIOSTAT bioreactor (Sartorius AG, Germany). Stargen and Fermgen (Genencor 143 International, The Netherlands) were employed to hydrolyze starch and proteins at 59°C and 144 pH 4.5 for one hour, respectively. Hydrolytic performance was investigated regarding 145 different solid-to-liquid ratios (11, 12.5, 20 and 25%, w/w) and enzyme loading (see section 146 3.2.2). Enzyme loading investigations were carried out at a solid-to-liquid ratio of 20% (w/w). 147 Mixing was set between 400 and 800 rpm depending on viscosity of the FW. Samples were 148 withdrawn, then inactivated at 95°C for 20 minutes, centrifuged at 5000 RPM for 10 minutes 149 and supernatant was stored at -20°C until used in analyses. 150

151 Yields of glucose and FAN per gram of dry food waste (Y, g/g) was calculated as follows:

152 Y = P / FW, where P [g] is the release in glucose or FAN and FW the amount of food waste 153 applied [g].

154

155 **2.4 Lactic acid fermentation**

LA fermentation was carried out in duplicate using a 2 L BIOSTAT bioreactor (Sartorius AG, Germany) containing 1 L of FW with a 20% (w/w) solid-to-liquid ratio. After enzymatic hydrolysis (see section 2.3), the reaction conditions were changed to 35°C and pH 6.0. A 6% (v/v) *Streptococcus* sp. strain A620 inoculum was used. Samples were analyzed for sugars (glucose, fructose and sucrose) and lactic acid concentrations. Results are presented as mean values of two replicates. After LA fermentation, solids and the oily phase were separated through centrifugation, and the supernatant was afterwards inactivated at 95°C for 20 minutes and stored at -20°C. The residual solids were mixed with the oily fraction floating on the supernatant and employed as feedstock for anaerobic digestion (AD) tests.

165

166 **2.5 Anaerobic digestion**

Three substrates (homogenized FW and fermentation residues from SHF and SSF processes) 167 underwent AD. AD batch tests were carried out at 37°C using 3% (w/w) total solids (TS) in 2 168 L (1.5 L working volume) SCHOTT glass bottles. Substrate-to-inoculum ratio was 2:1. 169 Digesters were manually shaken once a day. Each bottle was connected by 4/6 mm Teflon 170 171 tubes (PTFE, Germany) to 3 L sampling tubes containing a saturated saline solution acidified with some drops of concentrate sulphuric acid. Biogas volume and composition were daily 172 measured through water displacement and a gas analyzer (see section 2.6), respectively. Each 173 AD test was carried out in triplicate. Furthermore, controls using inoculum and cellulose, and 174 only inoculum (blanks) were carried out in triplicate. AD tests were finished when marginal 175 biogas production was below 1%. 176

Solubilization (made of disintegration and hydrolysis) is assumed as the rate-limiting step during AD of complex substrates rich in suspended solids (Van Lier et al., 2008). The disintegration constant (k_d) values were calculated as follows (Angelidaki et al., 2009). Assuming a first order kinetic model, the disintegration rate may be achieved through the first part of the cumulative biogas curve obtained from AD tests, according to:

$$B(t) = B)_{exp} \left(1 - e^{-k_{dis}t} \right)$$

182 where:

- 183 B(t) represents the cumulative biogas/methane production at a given time
- 184 B_{exp} is the ultimate biogas/methane potential yield of the substrate
- 185 k_{dis} is the first order disintegration rate [1/d]
- t is the time [day].
- 187

188 **2.6 Analytics**

Samples characterization was carried out in duplicate according to EPA reference methods (EPA, 2016) where not otherwise specified and mean values are presented. TS of FW and fermentation residues were analyzed after drying at 105°C until constant weight. Then dried FW and fermentative residues were weighted and combusted at 550°C for 5 hours in a muffle furnace for volatile solids (VS) analysis.

Fibers (ADF, NDF and lignin) were analyzed using an ANKOM²⁰⁰⁰ fiber analyser on FW
pre-dried at 60°C for 48 hours.

Sugars determination was carried out by cold water extraction. 3-5 g of dried FW and 50 mL of demineralized water were shaken for 30 minutes; afterwards 2 mL of a 30% (w/w) ZnSO₄ solution and 2 mL of a 15% (w/w) $C_6N_6FeK_4$ solution were added. After shaking, the mixture was filtrated and the clear filtrate analyzed by HPLC.

200 LA and sugars concentrations in fermentation samples were analyzed by HPLC (DIONEX, 201 USA): 10 μ L of sample was injected in a Eurokat H column (300 mm × 8 mm × 10 μ m, 202 Knauer, Germany) and eluted isocratically with 0.8 mL/min of 5 mM H₂SO₄. Detection was 203 carried out by a refractive index detector (RI-71, SHODEX, Japan). Each analysis was carried 204 out in duplicate.

Cat- and anion concentrations in fermentation samples were analyzed by ion chromatography
(DIONEX, USA). For quantification of cations, 25 μL of sample was injected in an IonPac

CS 16 column (250 mm × 4 μ m, DIONEX, USA) and eluted isocratically with 1.0 mL/min of 30 mM CH₃SO₃H at 40°C. For quantification of anions, 25 μ L of sample was injected in on an IonPac AS9-HC column (250 mm × 4 μ m, DIONEX, USA) and eluted isocratically with 1.2 mL/min of 9 mM Na₂CO₃ at room temperature. Detection of cat- and anions was carried out though a conductivity cell. Each analysis was carried in duplicate.

- Lipids analysis was performed by means of ANKOM Technology (USA) according to the ANKOM Technology Method 2, 01-30-09: Determination of Oil/Fat Utilizing High Temperature Solvent Extraction (ANKOM, 2014). Kjeldahl-nitrogen content in FW was determined according to DIN-EN-25663 standard method using a Kjeldahl System K-370/37. Protein content was calculated by multiplying the Kjeldahl-N content by 5.7 (Leung et al., 2012). Free amino nitrogen (FAN) concentration was measured using the ninhydrin reaction method (Lie, 1973), employing glycine as standard.
- Elemental analysis was performed with a VARIO EL III elemental analyzer according to themanufacturers' protocol (Elementar Analysensysteme GmbH, Germany).
- 221 Quantification of methane, carbon dioxide, oxygen and hydrogen sulfide produced during AD
- was carried out using a GA 2000 (Ansyco, Germany) gas analyzer.

223

224 **2.7 Statistical analysis**

One way analysis of variance was carried out in SigmaPlot and used to measure the statistical difference of LA formation between repetitions. Statistically significant difference in median values was accepted for P < 0.05.

228

229 3. Results and discussion

230 **3.1. Food Waste characterization**

FW consisted of (w/w): 18.1% TS, 93.2% VS/TS, 33.5% starch, 14.8% protein, 12.9% fat ,
8.5% free sugars, 8% NDF, 3.2% ADF and 0.1% lignin. Elemental analysis showed (values
referred to dry weight): 47.9% C, 7.67% H, 2.56% N and 0.09% S. FW composition is in
agreement with literature (Alibardi and Cossu, 2015; Campuzano and Gozalez-Martinez,
2016) and FW proved to be a suitable substrate for the proposed biorefinery concept.

236

237 **3.2. Enzymatic hydrolysis**

The efficient recovery of nutrients from FW strongly depends on the activity of enzymes added. Rosgaard et al. (2007) reported that the efficiency of an enzyme based hydrolysis of pretreated barley straw decreases when the viscosity of the slurry gets too high. To investigate this effect on food waste and to reduce the amount of enzyme needed to effectively hydrolysis food waste and to recover glucose and FAN different solid-to-liquid ratio and enzyme loadings were investigated.

244

245 3.2.1 Solid-to-liquid ratio

Glucose recovery was strongly dependent on the solid-to-liquid ratio (see Figure 1A). After 5-246 10 hours glucose concentration leveled off and 54.2 g/L was obtained when 11% (w/w) was 247 applied. Glucose concentration steadily increased to 80.9 g/L when 25% (w/w) was used. A 248 33.5% (w/w) starch content and a 25% (w/w) solid-to-liquid ratio accounts to a starch loading 249 of 83.8 g. The theoretical conversion of starch into glucose is 0.9 (Wymann et al., 2004), and 250 thus 94.4 g/L can be theoretically recovered. The obtained glucose concentration (80.9 g/L) 251 implies a recovery of 85%. Theoretically, 41.8 g/L of glucose can be obtained at a solid-to-252 liquid ratio of 11% (w/w). The obtained glucose concentration of 54.2 g/L, however, indicates 253 the presence of a remarkable amount of free glucose. Table 1 - 3 shows that the yield of 254 glucose per gram of FW decreases with increasing solid-to-liquid ratio. It is assumed that 255

better mixing conditions achieved at 11% (w/w) contributed to a better hydrolytic performance, and thus to a higher yield (0.49 g/g_{FW}), while at 25% (w/w) a yield equal to 0.33 g/g_{FW} was obtained.

Contrarily, even when the solid-to-liquid ratio was increased, the amount of recovered FAN 259 260 remained relatively constant (see Figure 1B). Even though the concentration increased from 0.23 g/L to 0.29 g/L within 24 hours with increasing solid-to-liquid ratio, this trend is not 261 comparable to the results shown in Figure 1A. The complete digestion of 14.3% (w/w) 262 proteins in FW would certainly have an effect on FAN concentration. However, it might be 263 concluded that proteases used are not appropriate for the digestion of proteins in FW. The 264 yield of FAN (see Table 1 3) decreased by increasing solid-to-liquid ratio. While 2.04 mg/g of 265 dry FW was obtained at 11% (w/w), only 1.15 mg/g was obtained at 25% (w/w). 266

267

268 **3.2.2 Enzyme concentration**

In order to determine the lowest specific enzyme loading for glucose and FAN recovery different specific enzyme loadings as shown in Table 2 4 were tested. Contrarily to the solidto-liquid ratio, the specific enzyme loading had no remarkable effect on glucose and FAN recovery (see Figures 1C and D). Yields were between 0.33 and 0.39 g glucose and between 1.82 and 1.92 g FAN per gram of dry FW (see Table 4).

274

275 **3.3 Lactic acid fermentation**

Due to the previously mentioned viscosity problems, 20% (w/w) solid-to-liquid ratio was chosen for LA fermentation. FW hydrolysis with Stargen was kept short for only one hour as it was found that the release of glucose occurs quickly (see Figure 1). After one hour 67.3 g/L of glucose was obtained which is in agreement with Figure 1. The hydrolyzed substrate was then inoculated with *Streptococcus* sp. strain A620 and the fermentation was carried out for 281 29 hours. Immediately after inoculation, LA concentration increased exponentially, reaching 282 39.2 g/L after 11 hours. Afterwards, it further increased linearly to 66.5 g/L until fermentation 283 was stopped (see Figure 2). Glucose was completely consumed, but traces of sucrose and 284 fructose, available as additional carbon sources, were still present. The first 11 hours was also 285 the period of time where most of the FAN was consumed (see Figure 2B). Fermentation was 286 carried out in duplicate and no statistical difference (P=0.637) was found for LA formation 287 between repetitions.

The overall yield obtained in this study using SHF, considering the LA concentration after 29 288 hours, was 0.33 g_{LA} per gram of dry FW with a productivity of 3.38 g_{LA}/L⁻h. SSF performed 289 290 on same FW resulted in a yield of 0.29 g_{LA}/g_{FW} and a productivity of 2.08 g_{LA}/L th after 28 hours (Pleissner et al., 2017), thus SHF resulted in higher yield and productivity. Higher 291 Yields (g_{LA}/g_{dry FW}) were found in literature are 0.27 (Kwan et al., 2016) and 0.99 292 293 (Kitpreechavanich et al., 2016) for SHF processes; 0.85 (Kim et al., 2016) and 0.46 (Tang et al., 2016) are accounted for SSF processes. (see Table 1), However the yield strongly depends 294 295 on substrate composition and on the strain. Productivity, defined as mass of LA generated per volume of fermentation broth in a time unit, is therefore a more reliable criterion to assess the 296 performance of a fermentation process. During exponential phase 3.38 g_{LA}/L⁺h was produced 297 298 in the present study, which is remarkably higher than productivity values in literature (see 299 Table 1). It is known that Streptococcus sp. strain A620 (Pleissner et al., 2017) is able to 300 degrade food waste, and thus this capability may additionally contribute to the release of 301 glucose. Lowest productivity of 0.28 g_{LA}/Lth was found when FW was converted with an indigenous microbial consortium (Tang et al., 2016). This is not surprising, as the microbial 302 consortium is not specialized to form only LA, but a mixture of different organic acids. The 303 study of Kim et al. (2016) is of particular relevance for FW utilization approaches as it 304 illustrates how FW can be utilized in repeated batch cultures over a long period of time. Even 305

though a higher productivity was obtained in the present study and by Kwan et al. (2016)
when FW was first enzymatically pretreated, the simplicity of processes presented by
Pleissner et al. (2017), Kim et al. (2016) and Tang et al. (2016) clearly shows that the process
steps can be reduced to a minimum.

310

311 **3.4 Anaerobic digestion**

AD tests lasted 20 days and resulted in following yields (see Figure 3): FW 0.710±0.02 312 Nm³/kg_{VS} biogas, 0.398±0.035 Nm³/kg_{VS} methane (56.35% v/v); fermentative residues from 313 SSF: 0.743 ± 0.01 Nm³/kg_{VS} biogas, 0.499 ± 0.008 Nm³/kg_{VS} methane (67.19% v/v); 314 fermentative residues from SHF: 0.90±0.016 Nm³/kg_{VS} biogas, 0.62±0.013 Nm³/kg_{VS} 315 316 methane (68.8% v/v). Biogas and methane yields obtained from fermentation residues are higher than the ones achieved from FW, mostly likely because of the differences among the 3 317 substrates in relative abundance of carbohydrates, proteins and lipids. In detail, fermentative 318 residues were rich in proteins and lipids, since their carbohydrate fraction was mostly already 319 exploited in LA fermentation. Hence biogas and methane yields of fermentative residues were 320 similar to pure proteins $(0.7 \text{ Nm}^3/\text{kg}_{\text{VS}} \text{ biogas}, \text{ with an average methane content equal to 70%},$ 321 v/v) and lipids (1.2 Nm³/kg_{VS} biogas with an average methane content equal to 68%, v/v) 322 (Weiland, 2010). FW was made of carbohydrates, proteins and lipids, but carbohydrates are 323 the most abundant fractions, and thus biogas and methane trends were comparable to 324 carbohydrates typical values (0.8 Nm³/kg_{VS} biogas, with an average methane content of 50%, 325 v/v) (Weiland, 2010). 326

Both SSF and SHF demonstrated two accomplishments: generation of a value added product (LA) and enhancement of biogas and methane yields. In a certain way, SSF and SHF had on AD the effect of a highly effective biological pre-treatment resulting in an improvement of methane production. In fact, the main purpose of AD pre-treatments is breaking the structure 331 of substrate particles and transforming them in easily biodegradable liquefied products 332 (Bracchitta, 2012). Considering the results achieved in the present research, it is possible to 333 affirm that LA fermentation exploited carbohydrate (mainly) and protein (partly) fractions, leaving the lipids almost unaltered for the consequently carried out AD process (see Table 3 334 335 $\frac{5}{5}$) and boosting the kinetics of methane production. This assumption was confirmed by the values of the disintegration constant (k_d), calculated according to Angelidaki (Angelidaki, 336 2009), which were equal to 0.43 1/d for FW, 0.35 1/d for SSF residues and 0.33 1/d for SHF 337 residues. These values are of the same order of magnitude of the ones obtained in other 338 studies (Fiore et al, 2016; Ruffino et al., 2015) using rice bran and husk (0.38 1/d), coffee dust 339 and peel (0.31 1/d), mixed vegetable waste (0.38 1/d) and pesto sauce waste (0.25 1/d). Other 340 Authors obtained 0.15-0.29 1/d for fruit pulp (Gali et al., 2009), 0.34 1/d and 0.26 1/d for 341 onion and potato respectively (Giuliano et al., 2013), and 0.14-0.35 1/d for mixed food waste 342 (Alibardi and Cossu, 2015). Moreover, However, the trend of k_d values obtained in this study 343 (FW>SSF>SHF) was expected because, as before mentioned, both fermentative residues were 344 deprived from the readily digestible carbohydrate fraction, with a higher efficiency of 345 enzymatic hydrolysis. 346

347

348 **3.5. Mass balance**

A mass balance was evaluated (see Figures 4-6) for three different process outlines with the aim to assess the amount of LA and biogas that may be generated considering different scenarios. In detail, LA production by means of SHF or SSF (Scenario 1); biogas generation through anaerobic digestion (Scenario 2); sequential production of LA from FW and of biogas from fermentative residues (Scenario 3). The mass balance starts with a theoretical amount of 1000 kg dry FW made of 335 kg of starch, 148 kg proteins 129 kg fat and 85 kg free sugars. About LA production, downstream processes are considered according to the process scheme usually adopted at Leibniz Institute for Agricultural Engineering and Bioeconomy in
Potsdam. In detail, a sequence of micro- and nanofiltration, softening, mono- and bipolar
electro dialysis, decolorisation, anion and cation exchange and distillation was taken into
account.

360 Considering Scenario 1, 148.2 kg of LA and 851.8 kg of wastes (residual solids plus LA lost in downstream process) and 149 kg of LA and 851.1 kg of wastes (residual solids plus LA 361 lost in downstream process) were produced respectively through SSF and SHF. Using 362 Scenario 2, 260.49 Nm³ of CH₄ and consequentially 2604.9 KWh of primary energy could be 363 produced. Taking into account Scenario 3, combined SSF and AD produced 148.2 kg LA and 364 236.5 Nm^3 of CH₄ and therefore 2365 KWh of primary energy and 417 kg of digestate; while 365 coupling SHF and AD produced 149 kg LA and 269.64 Nm³ of CH₄ and therefore 2696.4 366 KWh of primary energy and 408.52 kg of digestate. Wastes generated within the three 367 scenarios, residual solids generated by Scenario 1, as well as digestate deriving from 368 Scenarios 2 and 3 could be valorized in a composting process. 369

The mass balance of Scenario 1 (see Figure 4) underlines that the main bottleneck of LA fermentation is the huge amount of wastes produced after fermentation and downstream processes. In Scenario 3, this drawback is partially solved by the consecutive AD. Anyway, downstream processes are usually highly complex and expensive, and they require a careful optimization (Komesu et al, 2017).

375

376 4. Conclusions

This work investigated the technical feasibility of a sequential biorefinery process for the production of LA and biogas from FW via either SHF or SSF, which was proven. The main findings of the research are that SHF achieved higher yield and productivity than SSF, lasting one hour more than SSF. Sequential LA and biogas production moved forward from biomass conventional management and showed two profits: first, AD reduced and valorised the
fermentative residues generated from LA fermentation; second, SSF and SHF determined an
effective enhancement of biogas and methane yields with respect of FW.

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521

523 Figure captions

524 Figure 1. Solid-to-liquid ratio and enzyme loading. Recovery of glucose (A) and FAN (B) when enzymatic hydrolysis of blended food waste was carried out in presence of 350 µl 525 Stargen and 700 µL Fermgen at different solid-to-liquid ratios (w/w): 11.1% (open circle), 526 527 12.5% (closed circle), 20% (open triangle) or 25% (closed triangle). Recovery of glucose (C) and FAN (D) when enzymatic hydrolysis was carried out at a solid-to-liquid ratio of 20% 528 (w/w) at different specific enzyme loadings: 3.5 μ L/g Stargen and 5 μ L/g Fermgen (open 529 circle), 1.75 µL/g Stargen and 2.5 µL/g Fermgen (closed circle), 0.88 µL/g Stargen and 1.25 530 μ L/g Fermgen (open triangle), 0.44 μ L/g Stargen and 0.63 μ L/g Fermgen (closed triangle) or 531 0.11 µL/g Stargen and 0.32 µL/g Fermgen (open square). Results are based on single 532 measurements. 533

534

Figure 2. Lactic acid fermentation. Change of glucose (closed circle), fructose (open triangle), sucrose (open square), FAN (closed triangle) and lactic acid (open circle) concentrations during enzymatic pretreatment of food waste with 700 μ L Stargen and subsequently carried out lactic acid fermentation using *Streptococcus* sp. strain A620 (A and B). Fermentations were carried out in duplicate and mean values are shown. No statistical difference (P=0.637) was found between replicates.

541

Figure 3. Specific methane production from food waste (continuous line), SSF fermentative
residues (triangle-dot line) and SHF fermentative residues (dotted line) through anaerobic
digestion.

545

Figure 4. Mass balance from food waste to lactic acid: Scenario1 represents the L(+)-lactic
acid production through separate hydrolysis and fermentation (SHF) and simultaneous

- saccharification and fermentation (SSF). Mass balance is based on dry weight. OFMSW:organic fraction of municipal solids wastes
- **Figure 5**. Mass balance from food waste to biogas: Scenario 2 represents biogas and methane
- production through anaerobic digestion (AD). Mass balance is based on dry weight. OFMSW:
- 552 organic fraction of municipal solids wastes
- 553
- **Figure 6**: Mass balance from food waste to lactic acid and biogas Scenario 3 represents combined L(+)-lactic acid and biogas production. Mass balance is based on dry weight.
- 556 OFMSW: organic fraction of municipal solids wastes

558 **Table 1.** Lactic acid productivity (Pr) and yields of lactic acid per gram of dry food waste (Y_{FW}) when fermentation was carried out after

559 enzymatic hydrolysis or by simultaneous saccharification and fermentation.

		Pr	Y_{FW}	D.C.
Mode	Strain	[g/L⁻h]	[g/g]	Keterence
SHF	Streptococcus sp. strain A620	3.38	0.33	this study
SHF ¹	<i>L. casei</i> Shirota	2.61	0.27	(Kwan et al., 2016)
SHF ²	<i>Bacillus</i> sp. strain T27	0.44	0.99	(Kitpreechavanich et al., 2016)
SSF	Streptococcus sp. strain A620	2.08	0.29	(Pleissner et al. 2017)
SSF³	Indigenous microbial consortium	1.58	~0.85	(Kim et al., 2016)
SSF ⁴	L. casei	0.70	-	(Wang et al., 2016)
SSF	Indigenous microbial consortium	0.28	0.46	(Tang et al., 2016)

560 ⁴Food waste was pretreated with fungal enzymes

561 ²Studies were carried out with model kitchen refuse pretreated with glucoamylase

562 ³Fermentation was carried out as repeated batch culture

⁴Co-fermentation with sophoraflavescens residues in presence of cellulase and amylase

Substrate	Yield	$k_{dis} - (1/d)$	Reference 566
orange pulp	0.25 m ³ _{methane} /kg _{VS}	0.29	
pear pulp	0.15 m ³ methane / kg _{VS}	0.18	Gali et al., 2009 567
apple pulp	0.18 m^3 _{methane} /kg _{VS}	0.15	568
onion	0.92 m³biogas ∕kg _{VS}	0.34	Giuliano et al. 2013
potato	0.83 m3_{biogas}/kg_{VS}	0.26	<u>569</u>
tomato skins and seeds	0.42 m^3 biogas/kgvs		Dinuccio et al., 2010
fruit and vegetable waste	$\frac{0.32-0.63}{0.32-0.63}$ m ³ _{biogas} /kg _{VS}	nd	Gunaseelan, 2009 570
bread waste	0.58 m³biogas /kg _{VS}	nd	Kafle et al., 2013 571
vegetable waste	0.36 m ³ methane /kg _{COD}	nd	Maya-Altamira et al., 2008572
plain pasta	0.33 m^3 methane/kgvs	nd	573
cabbage	0.26 m^3 -methane/kgvs	nd	Labatut et al., 2011
potatoes	0.33 m^3 methane/kg _{VS}	nd	574
FW (50% bread, 20% vegetables, 10% fruit, 5% meat, 15 % nd)	0.43 m ³ biogas/kg _{VS}	0.35	575
FW (50% meat, 20% vegetables, 10% fruit, 5% bread, 15 % nd)	0.59 m ³ biogas/kg _{VS}	0.14	Alibardi and Cossu, 2015 576
FW (36% bread, 20% vegetables, 10% fruit 19% meat, 15 <u>% nd</u>)	$0.49 \text{ m}^3_{\text{biogas}}/\text{kg}_{\text{VS}}$	0.27	577
OEMSW	$\frac{1}{0.26 \text{ m}^3_{\text{biogas}}/\text{kg}_{\text{VS}}}$	nd	Fantozzi et al., 2011 578
	0.49 m ³ _{biogas} /kg _{VS}	nd	Pavi et al., 2017
FW	0.4-1.4 m ³ methane/kg _{VS}	nd	Elbeshbishy et al., 2012 579

Table 2. Biogas and methane yields and disintegration constants (k_{dis}) from food waste through AD in mesophilic conditions. nd= not defined

- **Table 1 3.** Yields of glucose (Y_{Glc/FW}) and FAN (Y_{FAN/FW}) per gram of dry food waste when enzymatic hydrolysis was carried out at different
- 582 solid-to-liquid ratios.

Solid-to-liquid ratio	$Y_{Glc/FW}$	Y _{FAN/FW}
[%, w/w]	[g/g]	[mg/g]
11.1	0.49	2.04
12.5	0.48	2.20
20	0.34	1.27
25	0.33	1.15

Table 2 4. Yields of glucose (Y_{Glc/FW}) and FAN (Y_{FAN/FW}) per gram of dry food waste when enzymatic hydrolysis was carried out at a solid-to-

- 592 liquid ratio of 20% (w/w) and different enzyme concentrations of Stargen and Fermgen per gram of dry food waste (n. a. = not analyzed).

Enzyme co	oncentration	$Y_{Glc/FW}$	Y _{FAN/FW}
[μ]	$[\mu L/g]$		[mg/g ⁻]
Stargen	Fermgen		
3.50	5.00	0.36	1.92
1.75	2.50	0.39	1.92
0.88	1.25	0.33	1.82
0.44	0.63	0.38	1.61
0.11	0.32	0.39	n. a.

Table 3 5. Evaluation of anaerobic digestion performances in terms of biogas and methane yields, of methane content and of disintegration

599 constant

Substrate	Biogas	CH ₄	CH ₄	k _{dis}
	$[Nm^3/kg_{vs}]$	[Nm ³ /kg _{vs}]	[%]	[1/d]
Food waste	0.71 ± 0.020	0.39 ± 0.035	56.35	0.43
Fermentative residues from SSF	0.74 ± 0.01	0.499 ± 0.008	67.19	0.35
Fermentative residues from SHF	0.90±0.016	0.62±0.013	68.80	0.33











Figure 4.



Figure 5.

Scenario 2



Figure 6.

