Biodegradability evaluation of waste-derived Polyhydroxyalkanoates

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Biobased and biodegradable plastics are currently object of high interest within the biorefinery and circular economy approach. In this view, the European Union implemented the current legislations in order to limit non-biodegradable plastic materials, such as petroleum-based single use plastic (SUP), and define a proper plastic-waste management. Waste valorization and energy recovery can be realized through composting and anaerobic digestion of these materials, since they are generally identified as biodegradable and compostable and then collected with the Organic Fraction of Municipal Solid Waste (OFMSW). Indeed, biodegradable bioplastics can be subjected to biological decomposition both in aerobic and anaerobic conditions, for the consequent production of carbon dioxide or methane. However, in order to be addressed to composting and anaerobic digestion treatments, either biodegradable and compostable plastics must meet the EU standards for SUP and packaging. Among biodegradable and biobased plastics, polyhydroxyalkanoates (PHA) are high added-value materials that can be recovered from waste activated sludge and different fermentable organic waste. They are biodegradable polyesters produced as intracellular carbon source by numerous microbes and have thermoplastic properties comparable to fossil-based plastics. The present study regards the evaluation of the Bio-Methane potential (BMP) of several PHA-based materials produced from mixed microbial culture and fermented mixture of OFMSW and sludge as substrate, within a pilot platform based in Treviso (Italy). PHA was produced from a feedstock composed by a mixture of the liquid slurry coming from squeezed OFMSW and the thickened sewage sludge from the treatment of municipal wastewater. The productive process is extensively described in previous study (Valentino et al. 2019; Moretto et al. 2020). Extraction with a mixture of aqueous-phase inorganic reagents was performed following a reserved protocol optimized by Biotrend S.A.. At the end of the extraction, the polymer was oven-dried obtaining a white powder (RU powder). In the framework of RES URBIS project, PHA purified by Biotrend was sent to SABIOMATERIALS for obtaining a melt-compounded pellet by mixing PHA with biodegradable additives (RU pellet). Finally, the pellet was sent to MiPlast, to produce blended PHA-PBS films by blown extrusion (Blend PHA-PBS). In this study, the powder and the pellet of RES URBIS PHA were melted and pressed (at a working T slightly below the melting point) to obtain films (RU film powder and RU film pellet, respectively). All these above-mentioned materials were tested for BMP in comparison with a commercial PHBV (HV 3%w/w) (Tianan powder) and thermal and chemical properties were determined before and after the biodegradation tests.

For the experimental set up, 14 serum bottles were filled with the 70 mL of mesophilic methanogenic sludge, 40 mL of mineral medium and 15mL of a 0.86 M NaHCO3 solution as buffer. In order to maintain the Inoculum/Substrate ratio = 2:1 (g VSS/g material), about 175 mg of PHA material was added. The tests were carried out in duplicate for each material and for the blanks. From the headspace, 50 µL of gas phase were sampled with a gas syringe and directly injected into the gas chromatograph equipped with heat conductivity detector (TCD) for the determination of methane. Liquid samples were taken to monitor the volatile fatty acids (VFAs) concentration and the pH. VFAs analysis of filtered sample was conducted on GC equipped with flame ionization detector (FID). The materials were analyzed for viscosity average molecular weight (Mn) determination, differential scanning calorimetry (DSC) and thermogravimetric analysis (TGA), following the procedures reported elsewhere (Lorini et al. 2021). After 140 days, tests were stopped and Blend PHA-PBS and RU pellet residues were recovered for further analysis (i.e. DSC and TGA).

VFA concentration monitored for each material is reported in figure 1A for the first 60 days of BMP test.

Figure 1. (A) VFA concentration; (B) specific methane production.
After an initial peak caused by the fermentation of PHA materials, the concentrations of acetic and propionic acid started to decrease, until zero from day 40 in all the bottles. This effect corresponded to the beginning of methane production, as it can be observed in Figure 1B, confirming the activity of the aceticlastic methanogens. Considering the kinetics of biodegradation and the produced methane, the highest conversion was obtained from both the PHBV powders (RU powder and Tianan) and from RU film powder, as a consequence of the high specific surface area. As regards RU pellet and Blend PHA-PBS, a significantly lower level of conversion into methane was obtained, as shown by the trends. Therefore, following the results obtained, it can be stated that the biodegradability is confirmed for those raw materials based on PHA, which have also demonstrated a rapid biodegradation kinetics, which has allowed to complete the conversion to methane in about forty days, when the trends reached the plateau. However, the kinetics of Blend PHA-PBS was probably slowed down because of a lower biodegradability of the PBS or for the presence of additives which may have inhibit methanogenesis. On the other hand, the lower kinetics of RU pellet conversion was due to the greater size of the pellet grains, which corresponds to a smaller specific surface. This result is even more evident considering that the RU film pellet was almost completely biodegraded with a significantly higher kinetics, as shown in figure 1B. Overall, considering the specific methane production (expressed as LCH4/kgTVS), a complete conversion to methane of PHA raw materials, including those produced from organic waste, can be confirmed (Battista et al. 2021). In Table 1, results of the chemical and thermal characterization are reported. Monomeric composition (i.e. HV content) of the whole RES URBIS set was in the range of 9 – 14 %w/w and the Mw of RU powder and pellet was lower than that one of Tianan. Melting temperatures (Tm) and temperatures at the highest rate of degradation (Tdmax) were determined by DSC and TGA analysis, respectively, for all the tested samples and also for RU pellet and Blend-PHBS residues. Indeed, since these latter were not easily biodegraded, it was possible to collect the residues at the end of the test. As an evaluation of the preliminary characterization of the recovered materials, it can be considered that they resulted macroscopically unchanged, as their properties, if compared to the original materials (158 – 167 °C and 263 – 288 °C, Tm and Tdmax ranges).

As a main result, the biodegradability of PHA raw materials has been confirmed in anaerobic conditions, suggesting the possibility to dispose PHA wastes together with the OFMSW for anaerobic digestion and compost applications. On the other hand, further investigation are required, since the presence of additives and plasticizers, necessary for plastic processing and depending from the application requested, affect the biodegradability. In this view, further characterization of the residual solid materials and tests conducted in different conditions (e.g. thermophilic anaerobic BMP tests) will be carried out.

Table 1. Chemical and thermal properties of the tested materials and two recovered materials (R) at the end of the tests.

<table>
<thead>
<tr>
<th>Sample</th>
<th>HV (%w/w)</th>
<th>Mv (kDa)</th>
<th>Tm (°C)</th>
<th>Tdmax (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RU powder</td>
<td>12</td>
<td>207.4</td>
<td>158</td>
<td>288</td>
</tr>
<tr>
<td>RU pellet</td>
<td>9</td>
<td>159.2</td>
<td>164</td>
<td>283</td>
</tr>
<tr>
<td>Tianan</td>
<td>3</td>
<td>322.4</td>
<td>175</td>
<td>261</td>
</tr>
<tr>
<td>Blend PHA-PBS</td>
<td>14</td>
<td></td>
<td>167</td>
<td>280</td>
</tr>
<tr>
<td>RU pellet (R)</td>
<td>9</td>
<td></td>
<td>165</td>
<td>282</td>
</tr>
<tr>
<td>Blend PHA-PBS (R)</td>
<td>14</td>
<td></td>
<td>164</td>
<td>263</td>
</tr>
</tbody>
</table>

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References


