Microbial Enzyme Applied to Plastic Depolymerization

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The accumulation of synthetic plastic waste in the environment has become a global concern. Microbial enzymes (purified or as whole-cell biocatalysts) represent emerging biotechnological tools for waste circularity; they can depolymerize materials into reusable building blocks, but their contribution must be considered within the context of present waste management practices.

Keywords: biotechnology ; biodegradation ; plastic ; waste managment ; circular economy ; recycling

1. Introduction

Biotechnology tools have been employed for more than 20 years to depolymerize complex biomass waste into organic building blocks that, in turn, can be converted into useful intermediate feedstocks. These processes are typical applications of the circular (bio)economy concept. One of the most interesting examples is the conversion of lignocellulosic materials contained in plant waste from farming activities into bioethanol ^{[1][2]}. More recently, biotechnology strategies based on the use of microbial enzymes have been applied also to the degradation of petroleum-based plastics with the aim of providing greener alternatives to the traditional methods of recycling.

Various enzymes are used by bacterial or fungal organisms to fragment polymers into units of about 10–50 carbon atoms, which are used as carbon sources ^[3]. The catalytic capacity of highly purified stocks of enzymes (produced at an industrial level) can be exploited in biochemical processes performed under mild conditions (e.g., water solution in the 20–80 °C range) because they are biocatalysts evolved to act under conditions compatible with life. This allows for lowering of the economic and environmental costs of traditional processes, which usually require high temperatures and pressures ^[4]. The wide diversity in the metabolism of living organisms allowed for the discovery of biocatalysts suitable for many chemical reactions, including those responsible for plastic degradation, although consortia of whole organisms may also be required in some cases ^[5]. The enzymatic or microbial recycling of plastic materials is divided into three main steps: (i) the production of the biocatalysts, (ii) the depolymerization reaction of the polymer into monomers, and (iii) the purification and reuse of the monomers for resynthesis of the original material (closed-loop processes) or their bioconversion into different products (open-loop processes) ^[6].

A plethora of options is available for open-loop routes leveraging biotechnologies, as enzymes can catalyze multistep reactions to transform monomers into completely different molecules, which possibly have a value (per mass unit) superior to that of the original polymeric material ^[Z]. The processes for open-loop plastic recycling can be performed in different bioreactors (employing purified enzymatic cocktails or whole organisms) or in a one-pot system ^[6]. The main classes of enzymes involved in plastic degradation and their products are reported in **Figure 1**. **Table 1** reports the best-performing bio-based plastic depolymerization systems, which are predominantly implemented at laboratory scale.



Figure 1. The main classes of enzymes involved in the degradation of different types of traditional petroleum-based plastics in order of their biodegradability to summarize what was reported in the text. Degradation products are exemplified. PURs—polyurethanes, AA—adipic acid, BDO—1,4-butanediol, TDA—toluene diamine, PE—polyethylene, PP—polypropylene, PAs—polyamides, PVC—polyvinylchloride, PS—polystyrene, PET—polyethylene terephthalate, EF— ethylene glycol, TPA—terephthalic acid, MHET—2-hydroxyethyl terephthalate, BHET—bis(2-hydroxyethyl) terephthalate.

Plastic Polymer	Material	Source	Reaction Condition	Degradation Rate	Reference
PET	Ultrathin PET film (2.5 to 7 nm) obtained from amorphous PET sheets of 2 mm thickness	ISPETaseTM ^[8] and ISMHETaseSM ^[9] , two variants of <i>I. sakaiensis</i> PETase and METase	PBS buffer, pH 7.4, 50 °C	≈70% in 1 h	[10]
PET	Amorphized and micronized PET (≈200–250 μm) from post-consumer bottle-grade PET	Cutinase from leaf– branch compost (ICCG variant)	100 mM potassium phosphate buffer, pH 8.0, 72 °C	≈90% in 10 h	[11]
PET	2 × 1 cm ² amorphous Goodfellow PET film and low-crystallinity (13%) PET powder	PHL7/PES-H1, a cutinase from a compost site (L92F/Q94Y variant)	1 M potassium phosphate buffer, pH 8.0, 72 ℃, shaking at 1000 rpm	≈100% in 24 h	[12]
PET	3 × 0.5 cm ² flakes of an amorphous PET clamshell container	PHL7/PES-H1, a cutinase from a compost site	1 M potassium phosphate buffer, pH 8.0, 70 ℃, shaking at 1000 rpm	>95% in 24 h	[13]
PET	Low-crystallinity (1.2 to 6.2%) discs (6 mm) from 51 different post-consumer PET products	FAST-PETase, a variant of <i>I. sakaiensis</i> PETase	100 mM potassium phosphate buffer, pH 8.0, 50 °C, shaking at 180 rpm	≈100% in 1 to 7 days	[14]
PET	PET/PE composite packaging tray lid (4 mg, thickness of 325 μm PET and 40 μm PE)	HotPETase, a directed- evolved variant of <i>I.</i> <i>sakaiensis</i> PETase	pH 9.2, 50 mM gly- OH buffer with 4% BugBuster	≈20% in 24 h	[15]

Table 1. List of the most performant biological systems for traditional petroleum-based plastic degradation known to date. They are ordered by plastic polymer type and listed by decreasing degradation rate.

Plastic Polymer	Material	Source	Reaction Condition	Degradation Rate	Reference
PET	≈37% crystallinity PET microplastics (≈300 μm)	TS-∆IsPET, a variant of PETase from Ideonella sakaiensis	100 mM potassium phosphate buffer, pH 8.0, 40 °C, shaking	≈26% in 2 days	[<u>16</u>]
Polyester- type PUR	Lab-prepared 10 µm thin PUR and segmented PUR urea films based on lysine diisocyanate	Papain, Bromelain, Ficin, Chymotrypsin, Proteinase K	PBS, pH 7.0, 37 °C	Up to ≈50% in 7 days	<u>[17]</u>
Polyester- type PUR	Impranil-DLN®	Pseudomonas putida A12	рН 8.0, 25 °С	92% in 4 days	[18]
Polyester- type PUR	Impranil-DLN [®]	Pestalotiopsis Microspora E2712A	25 °C in a rotary incubator	99% in 2 weeks	[19]
Polyether- type PUR	PUR foam	Tenebrio molitor	Guts of the larvae (probably microbiota-assisted)	67% after 30 days	[20]
Polyester- type PUR	Lab-prepared 0.3 mm thin polycaprolactone thermoplastic PUR film (Capa 2302)	Amidase E4143 and esterase E3576	PBS, pH 7.0, 37 °C	33% in 51 days	[21]
Polyester- type PUR	Impranil-DLN®	Cladosporium pseudocladospo-rioides, Cladosporium tenuissimum, Cladosporium asperulatum, Cladosporium montecillanum, Aspergillus fumigatus, Penicillium chrysogenum	25 °C, no shaking	40–87% in 2 weeks	[22]
Polyester- type PUR	Commercial 1 mm thin PUR film	Aspergillus flavus ITCC 6051	28 °C under 120 rpm shaking	60.6% in 30 days	[23]
Polyester- type PUR	Lab-prepared ~0.2 mm thin PUR	Bacillus subtilis MZA-75 and Pseudomonas aeruginosa MZA-85	37 °C under 150 rpm shaking	40% in 30 days	[24]
Polyether- type PUR	PUR foam used for commercial production of mattress cushioning	Cladosporium pseudocladospo-rioides, Cladosporium tenuissimum, Cladosporium asperulatum, Cladosporium montecillanum, Aspergillus fumigatus, Penicillium chrysogenum	25–30 °C, no shaking	10–65% in 21 days	[<u>22]</u>
Polyether- type PUR	Cubical ether–PUR (1 cm ³)	Alternaria sp. PURDK2	30 °C, no shaking	27.5% in 70 days	[25]
LDPE	LDPE bag	Galleria mellonella	-	13% in 14 h	[26][27][28]
LDPE	LDPE film	Pseudomonas citronellolis	37 °C under 15 rpm shaking	17.8% in 4 days	[29]
LDPE	LDPE powder	Cupriavidus necator H16	30 °C in a rotary incubator	33.7% in 21 days	[30]
LDPE	LDPE foam	Tenebrio molitor	Gut of the larvae (probably microbiota-assisted)	49.0% in 32 days	[31]
LDPE	20 μm thin PE film	Oscillatoria subbrevis	-	30% in 42 days	[32]
PS	PS pyrolysate oil	Pseudomonas putida CA-3	Four consecutive treatments at 30 °C	10% in 8 days	[33]

Plastic Polymer	Material	Source	Reaction Condition	Degradation Rate	Reference
PS	Brominated high-impact PS	Bacillus sp.	30 °C, no shaking	23.7% in 30 days	[34]
PS	Brominated high-impact PS	Exiguobacterium sp. strain YT2	30 °C under 150 rpm shaking	12.4% in 30 days	[35]
PP	Max 250 µm microplastic PP	Bacillus cereus	33 °C under 150 rpm shaking	12% in 40 days	[36]
PP	Isotactic PP strips	Pseudomonas sp., Vibrio sp., Aspergillus niger	рН 7.0, 30 °С	60% in 175 days	[37]

2. Enzymatic Polyethylene Terephthalate Recycling: Potential Industrial-Scale Application

2.1. Enzymatic Degradation of Polyethylene Terephthalate

PET is a semicrystalline material made up of polymeric chains of alternating terephthalic acid (TPA) and ethylene glycol (EG) joined by ester bonds. In crystalline regions, the polymer chains are tightly packed in parallel, while in amorphous regions, the chains assume a disordered conformation. The degree of crystallinity depends on the production process and affects the chemophysical properties of the polymer which, in turn, are related to the final product performance. For instance, PET used in the production of bottles and textile fibers has a high crystallinity (between 30–40%), while PET is mainly amorphous in food packaging (crystallinity < 10%).

Research on biotechnological degradation of PET started in 2005 ^[38]. Since then, several microbial PET-hydrolyzing enzymes (PHEs) have been discovered for their ability to break down PET into mono-(2-hydroxyethyl)-terephthalate (MHET), bis-(2-hydroxyethyl)-terephthalate (BHET), TPA, and EG by hydrolyzing the ester bond ^[39] (**Figure 1**). Examples of the most active enzymes are reported in **Table 1**. These enzymes descend from the same ancestral carboxyl ester hydrolase of the GxSxG family ^[40]. They share the same catalytic mechanism and span different enzymatic classes specialized on different natural substrates: cutinase (EC 3.1.1.74), lipase (EC 3.1.1.3), and PETase (or PET hydrolase) (EC 3.1.1.101) ^{[41][42][43]}. Remarkably, the PETase class was created in 2016 after the discovery of a PHE from *Ideonella sakaiensis* (ISPETase), a bacterium isolated near a WWTP, with optimal PET degradation activity at 40 °C ^[44]. ISPETase and other *I. sakaiensis* enzymes were identified and shown to allow for the depolymerization and metabolic assimilation of PET degradation products as a carbon source ^[45]. The relevance of such a PET waste assimilation process in natural environments was questioned because the depolymerization of highly crystalline PET (such as that used to produce bottles) under natural conditions for the growth of the microorganism (30 to 40 °C) is highly inefficient ^{[46][42]]}. Indeed, highly crystalline PET regions and low temperatures (<40 °C) greatly reduce the accessibility of PET chains to PHEs. Accordingly, the accessibility is maximal between 70 and 74 °C, near the glass transition temperature of bulk PET ^{[48][49]}.

Consequently, known PHEs seem unsuitable for efficient degradation of untreated highly crystalline PET at room temperature ^{[49][50]}. This prompted a search for thermostable PHEs produced by protein engineering ^{[8][11][12][14][51][52]} or enzyme discovery ^{[13][53][54][55]}. Among such attempts, the company Carbios, in a public–private partnership with the Toulouse Biotechnology Institute and other global partners, was the first to produce a thermostabilized engineered variant of the leaf–branch compost cutinase (LC-cutinase) suitable for an industrial-scale recycling process of PET in 2020 ^[56]. Carbios built a demonstration plant in September 2021 with a business plan to extend the system to other companies under a paid license agreement by 2023 and revenues expected by 2025 ^[57].

The Carbios technology and other promising enzymatic PET recycling processes (**Table 1**) act on low-crystallinity submillimeter PET films ^{[12][13][14][58]} or amorphized/micronized post-consumer PET materials (C-ZYME[®], based on the enzyme variants reported by Tournier et al. ^[11] and various patents (i.e., WO2017198786A1, EP3517608A1, and WO2020021118A1). In practice, post-consumer PET degradation processes were operated after sorting, washing, grinding, and amorphization. These pretreatment steps were already established for non-enzymatic recycling processes ^[59]. The step involving the use of enzymes was designed to be performed in <1 day in a buffered water suspension (pH between 7 and 9) inside a bioreactor containing a preprocessed polymer and close to the PET glass transition temperature. However, enzymatic incubation above 70 °C does not permit complete degradation of the amorphized PET due to an ageing process that results into polymer recrystallization after <10 h ^{[60][61]}. Recent studies ^{[61][62][63]} revealed relatively high degradation rates, even when performing the reaction at lower temperatures (50 to 60 °C), despite requiring more time (≥1 day). These milder conditions were also tested on untreated and non-micronized low-crystallinity post-

consumer PET materials $\frac{[13][14]}{2}$. Moreover, Tarazona et al. recently demonstrated that a reduced PET thickness and an increased accessible surface area reduce the glass transition temperature by up to ~40 °C (for nanometric film) $\frac{[10]}{2}$; while this can speed up PET degradation at lower temperatures and inhibit recrystallization, it does not block degradation-induced thermal inactivation, which consists of the accumulation of inactivated protein aggregates on the PET surface, causing the inhibition of further polymer degradation (a process called *passivation*).

Overall, promising industrial-scale PET degradation technologies require a buffered water suspension at ≥50 °C and PHEs engineered to be stable, efficient, and with low product inhibition (eventually by combining multiple enzymes) allowing for complete depolymerization as fast as possible in order to prevent PET recrystallization and passivation [10]. A recent work showed significant condition-dependent differences in the depolymerization process (buffer, pH, enzyme, and substrate loading) catalyzed by similar engineered PHE variants [62]. In addition, the enzymatic capacity to adsorb and turn-over during PET chain hydrolysis were recently observed to follow a tradeoff [64]. For these reasons, the biochemical kinetics and the mechanism of enzymatic PET degradation should be studied in detail to select the best enzymes and reaction conditions under which both the PHE activity is optimal and the PET polymer is most degradable, prior to scaling up the process [65]. Moreover, it is critical to set up a standard assay condition to compare different PHEs (different conditions are used for the enzymes reported in Table 1). As an example, a recently thermostabilized IsPETase (HotPETase) was shown to have initial degradation rates at 60 °C, much higher than the ICCG LC-cutinase used by Carbios on low- and high-crystalline materials (including a composite PET-PE film); however, it mainly produced MHET (TPA is the main product of ICCG LC-cutinase) and was inactivated after few hours on highly crystalline substrates, before the degradation was completed [15]. Furthermore, L92F/Q94Y PHL7 cutinase was more efficient than ICCG LC-cutinase on low-crystalline materials, while it was less active on highly crystalline PET powder [12]. Finally, different relative degradation rates on PET of different sizes and crystallinity were also measured when comparing many recently discovered PHEs [66], motivating the possible need for a cocktail of multiple PHEs for the degradation of PET materials of different crystallinity and subjected to different pretreatments.

2.2. Closed-Loop Polyethylene Terephthalate Recycling by Enzymatic Depolymerization

Carbios company demonstrated the feasibility of a closed-loop recycling process by showing that it is possible to use the monomers (TPA and EG) produced from PET depolymerization of post-consumer PET flakes to produce a PET bottle possessing the same properties as a bottle produced from virgin PET ^[11]. This process may also be fostered by monomers recovered from PET contained in textile polyester waste; an enzymatic cocktail of PHEs and other enzymes could be added to chemical treatments (in concert or in a sequential manner) to allow for separation and to reach a superior depolymerization yield of polymers without causing the high material losses associated with traditional methods. This is because the PET contained in fibers of commercialized textile products is often blended with other natural (i.e., cellulose and cotton) or synthetic (polyesters, nylons, elastane, etc.) polymers ^{[67][68]}.

A recent work used Carbios data to conduct a technoeconomic analysis (TEA) of a closed-loop PET recycling process from bottle-derived post-consumer flakes ^[59]; a minimum production cost (MPC) for TPA of USD 1.93 kg⁻¹ was estimated, most of which (>50%) was due to the price of petroleum-based PET flakes (feedstock PET) acquired from MRFs, the plastic pretreatment, and the recovery of the degradation products and co-products. In contrast, the step involving enzymes accounted only for 20% of the cost. Nevertheless, the model shows that besides the feedstock PET price, which is an external factor, the load of PET in the degradation solution and the rate of enzymatic conversion are the other dominant factors affecting MPC, enabling a reduction in cost of up to USD 1.60 kg⁻¹ [59], which is less than double the average petroleum-based PET price in 2021 (up to USD 0.85 kg⁻¹) ^[69]. It must be pointed out that this TEA estimated that the environmental impact of TPA obtained from the enzymatically based recycling process is nearly halved with respect to virgin TPA production. A recently published life cycle assessment (LCA) based on the USA economy confirmed the previous estimation of the production costs [59] but indicated that current enzymatic PET recycling technology exhibits a 1.2 to 17 times superior environmental impact relative to virgin polymer production ^[70]. NaOH usage, PET waste collection/shredding/amorphization, and electricity were estimated to be the highest-impact contributors. Sensitivity analysis indicated that, through interventions during all stages of the recycling process, from feedstock preparation to product recovery, it is possible to achieve an environmental impact similar or even lower than virgin polymer production. However, a superior enzymatic depolymerization efficiency without changing any other condition was estimated to contribute only up to 14%. Main contributions were estimated to be provided by minimization of PET losses during sorting, elimination of PET flake pretreatments, enzyme immobilization, and reaction buffer reuse. A major environmental benefit should originate from the reduction in NaOH consumption by using completely different buffers and PET loadings to facilitate TPA precipitation or by using more environment friendly (e.g., ammonia) or recoverable bases. Overall, this research suggested that further process optimization is required to minimize waste during feedstock preparation and product recovery.

Novel biocatalysts should allow for effective PET depolymerization under different buffer conditions (including organic solvents) of less pretreated PET-containing materials rather than showing a superior degradation rate under a broad range of pH and temperature conditions. Moreover, this process should be evaluated on different sources of PET waste, including those from commonly mixed products (i.e., textiles) for which traditional technologies are less effective and for which both the high crystallinity and the material substructure are expected to pose accessibility challenges to degradative enzymes ^[67]. The goal is the setup of a closed-loop enzyme-based process with a halved environmental impact compared to virgin polymers, compensating for the superior costs of PET produced from the enzymatic recycling process with an increased environmental gain.

2.3. Open-Loop Polyethylene Terephthalate Bio-Recycling

The use of biotechnological tools into open-loop processes could represent an opportunity to overcome the overpricing problem of closed-loop enzymatic PET recycling; enzymes and/or microorganisms can be used to assimilate and metabolically convert PET building blocks (TPA and EG) into chemically different intermediate bulk compounds or polymers with high added value ^[71]. This could also ensure the economic sustainability of treating low-quality plastics collected from contaminated environments. Several processes have been tested at the laboratory scale (**Table 2**).

Kim et al. [72] presented a two-organism system for the biological conversion of chemically depolymerized PET; TPA was converted into protocatechuic acid (PCA) and other PCA-derived compounds (i.e., gallic acid, pyrogallol, catechol, muconic acid, and vanillic acid) by an engineered E. coli strain, while G. oxydans KCCM 40109 was used to produce glycolic acid from EG. These compounds are important bulk chemicals to produce inks, bioplastics, flavorings, and fragrances. The same authors also demonstrated the possibility of using TPA produced by the enzymatic depolymerization of PET to produce catechol for the production of multifunctional coating materials [73]. Kang et al. [74] proposed the use of an engineered E. coli strain to convert TPA monomers from post-consumer PET into 2-pyrone-4,6dicarboxylic acid, a valuable monomer for the synthesis of next-generation biodegradable plastics. A Pseudomonas putida KT2440 strain was engineered to use BHET (a different PET biodegradation product) to produce β-ketoadipic acid, a building block used to produce performance-advantaged nylons [75]. A one-pot system was proposed by Tiso et al. for the sequential enzymatic depolymerization of PET into EG and TPA ^[76], followed by *Pseudomonas umsongensis* GO16 conversion into intracellular polyhydroxyalkanoates (PHAs) and extracellular hydroxyalkanoyloxy alkanoate, which is used in a chemical copolymerization reaction to produce a novel bio-based poly(amide urethane) PUR. A similar system based on the use of an *E. coli* strain was proposed by Sadler et al. to produce vanillin flavor [77]. Liu et al. [78] proposed the first one-pot microbial recycling system from PET using two co-cultivated engineered organisms: a Yarrowia lipolytica Po1f strain expressing a PHE to depolymerize PET at low temperature and a Pseudomonas stutzeri strain to convert TPA into the bioplastic polyhydroxybutyrate. The same group showed that muconic acid can be produced with a single engineered Pseudomonas putida strain ^[79].

Table 2. List of recent open-loop microbial conversion attempts starting from PET and polyester-PUR degradation products. The different approaches are distinguished with respect to the processing strategies proposed in ^[6]. Abbreviations: EG—ethylene glycol, AA—adipic acid, BDO—1,4-butanediol, TPA—terephthalic acid, BHET—bis(2-hydroxyethyl) terephthalate, PHAs—polyhydroxyalkanoates, HAA—hydroxyalkanoyloxy alkanoate, PHBS—polyhydroxybutyrate.

Approach	Enzymes or Microbial Strain(s)	Plastic Monomer	Conversion Products (Yields %)	Reference
	Pseudomonas putida strains JM37 and KT2440	EG	Glyoxylic acid (≈20%)	[80]
	Three derived Pseudomonas putida KT2440 strains	AA, BDO, EG	Monorhamnolipids (<1%)	[81]
Separate depolymerization and conversion	Escherichia coli strain	TPA, EG	Gallic acid (≈93%), pyrogallol (≈40%), muconic acid (≈85%), vanillic acid (≈40%), glycolic acid (≈98.6%)	[72]
	Escherichia coli strain	TPA	2-pyrone-4,6-dicarboxylic acid (96.08%)	[74]
	Pseudomonas putida KT2440 strain	BHET	β-ketoadipic acid (≈76%)	[75]
	Escherichia coli strain	TPA	Catechol (≈67%)	[73]

Approach	Enzymes or Microbial Strain(s)	Plastic Monomer	Conversion Products (Yields %)	Reference
Combined	LCC ¹ , PseudomonasTPA, EGPHAs and HAACombinedumsongensis GO16 strain(<2%)	PHAs and HAA (<2%)	[76]	
conversion	LCC ² , <i>Escherichia coli</i> MG1655 RARE strain	TPA, EG	Vanillin (≈79%)	[77]
Consolidated	Yarrowia lipolytica Po1f expressing PETase ³ , TPA, EG F Pseudomonas stutzeri TPA3	PHB (≈2.5%)	[78]	
bioprocessing	Pseudomonas putida KT2440-tacRDL expressing LCC ⁴	TPA, EG	Muconic acid (≈50%)	[<u>79</u>]

1 Wild-type enzyme from ^[56]. ² WCCG variant from ^[11]. ³ Variant from ^[82]. ⁴ ICCG variant from ^[11].

These examples of open-loop PET recycling employing enzymes and microorganisms represent proof-of-principle experiments demonstrating the potential of microbial enzyme-driven upcycling processes.

3. Polyurethane Biodegradation Is Restricted to Polyester-Based Polymers

3.1. Enzymatic Degradation of Polyurethanes

PURs represent the largest fraction of plastic waste that is unrecyclable using traditional technologies. PUR polymers are formed by many different subtypes of two building blocks belonging to polyol (e.g., 1,4-butanediol, BDO) and isocyanate (e.g., toluene diamine, TDA) classes linked by urethane bonds (**Figure 2**). Moreover, isocyanates usually contain aromatic rings that increase the rigidity of the polymer and are covalently crosslinked to polyols of other chains by the formation of ester bonds in polyester-type PURs or by the formation of ether bonds in polyether-type PURs ^[83]. PURs are among the most diversified polymers, with different compositions in bonds, building blocks, and crosslinking distributions. This permits the generation of PUR materials with tailored physical properties ^[84]; however, the consequence is that a material-specific toolbox of enzymes with different activities is required for their degradation ^[85].

The enzymes identified to date to cause some degradation or weight loss of PURs are mainly promiscuous esterase (EC 3.1), urease (EC 3.5.1.5), protease (EC 3.4.21), and amidase (EC 3.5.1.4) enzymes ^[84]. Some esterases were shown to potentially degrade liquid dispersion (Impranil-DLN[®]) or bulk polyester-type PURs, but there is no evidence of their capacity to cleave urethane bonds ^[84]. Proteases and amidases have been shown to hydrolyze both urethane and ester bonds, but very low degradation rates were reported on commercial elastomer polyether-type PUR films or polyester-type PUR pellets ^[86]. A combined system involving an esterase and an amidase showed improved synergistic degradation (33%) after 51 days on a model PCL-based thermoplastic PUR ^[21]. A laccase-mediated (EC 1.10.3.2) system previously reported to degrade PE caused a reduction of a few percentage points in weight of lab synthesized polyether-type PURs in 18 days ^[87]. Urethanases (EC 3.5.1.75), which are enzymes capable of specifically hydrolyzing the urethane bond between polyols and isocyanate units, were recently discovered ^[88]; although their use for full depolymerization of PURs within a few days is very promising, these enzymes were used and demonstrated to be effective only on the urethane bonds in low-molecular-weight dicarbamate aromatic diamines obtained after the chemical depolymerization (by glycolysis) of non-urethane bonds in a polyether PUR foam. Therefore, this process is bio-based only in the last step.

Overall, even if enzymes capable of degrading the urethane and ester bonds in PURs have been discovered, processes that make exclusive use of enzymes have relatively high degradation rates only on synthetic or oligomeric PURs designed to be more biodegradable ^{[89][90]}. Therefore, the best options for PUR biodegradation are the use of environmental microbial strains or consortia, the design of more degradable novel PUR materials, or synergy with chemical methods of depolymerization ^[91].

3.2. Microbial Degradation of Polyurethanes

Some promising microbial degradation systems acting on polyester-type PURs have been tested at lab scale in the last 10 years. Impranil-DLN[®] was used as the sole carbon source by *Pseudomonas putida* A12, achieving 92% degradation within four days under mild conditions ^[18]. *Bacillus subtilis* MZA-75 and *Pseudomonas aeruginosa* MZA-85 were able to co-act on a polyester-type PUR film, resulting in 40% weight loss after 30 days ^[24]. Examples of polyester-type PUR biodegradation were also reported for fungal strains, both on Impranil-DLN[®] ^[19], a liquid varnish ^[22], and a commercial

film ^[23]. Oligomeric fragments are usually reported as degradation products, which means that complete hydrolysis to single building blocks was not achieved. Moreover, microorganisms usually require other carbon sources to grow on PURs and may be susceptible to the accumulation of metabolically toxic plastic degradation products or additives, as observed for a *Pseudomonas* strain that released diamines from degraded PURs ^[81].

Polyether-type PURs show higher recalcitrance to microbial biodegradation, in particular when rigid aromatic polyisocyanate units are present, since they are less accessible and not susceptible to degradation by esterase activity ^[83]. Until now, only three studies have reported a significant biodegradation of polyether-type PUR materials: *Tenebrio molitor* larvae were shown to ingest polyether PUR foams efficiently, resulting in a significant mass loss of nearly 67% after 35 days. However, polyether PUR fragments were found in the frass of the larvae, indicating partial degradation ^[20]. *Alternaria* sp. strain PURDK2 caused 27.5% degradation of polyether-type PUR cubes after 10 weeks ^[25], and some *Cladosporium* fungal strains were reported to cause about 65% weight loss of a polyether-type PUR foam in 21 days ^[22].

Considering the reported degradation rates, the use of simple ad hoc synthesized or non-waste PUR products, and the lack of knowledge on the complex enzymatic machinery produced by microbial consortia ^[21], it is not currently possible to depict a practical industrial-scale biodegradation process, even for less recalcitrant polyester-type PUR waste. Moreover, solid polyester-type PURs and polyether-type PURs were only partially biodegraded (up to 50% at most), requiring a few weeks or even a few months. Therefore, for more recalcitrant PURs, a manly bio-based depolymerization process is still unfeasible.

In the next few years, it will be crucial to isolate the microbial enzymatic machinery involved in the degradation of less recalcitrant polyester PURs by using model PUR substrates and developing new specific assays (as presented by Liu et al. ^[92]). This will allow for mimicking of the whole-organism biodegradation action using an enzymatic cocktail or a stepwise addition of enzymes. Future investigations should also be focused on specific commercial PUR materials ^[84].

3.3. Closed-Loop and Open-Loop Polyurethane Bio-Recycling

The complex structure of PURs leads to a variety of degradation products, including organic acids, organic alcohols (e.g., EG), and diamines (**Figure 1**). Carboxylic acids and alcohols can be used for the synthesis of virgin PURs and other poly(ester–ether–urethane) materials ^[91]. Amines such as toluene diamine (TDA) can be recovered and used to directly synthesize polyamides or toluene isocyanate (TDI) to subsequently make virgin PURs (bio-based closed-loop recycling). Although the full depolymerization of a polyether-PUR foam was demonstrated with a chemoenzymatic approach ^[88], its application on a real waste PUR material and the use of the degradation products (polyether-polyols, diethylene glycol, and aromatic diamines) for the resynthesis of a waste-like PUR material was never attempted.

Some PUR degradation products have also been tested as microbial carbon sources (**Table 2**). EG can be used as a substrate to produce PHAs, which can be used to manufacture biodegradable medical devices, as well as glyoxylic acid, which is usually used for buffering solutions and other products to treat human skin and hair ^[80]. Recently, polyols (adipic acid (AA), 1,4-butanediol (BDO), and EG) were used as a carbon source for a mixed culture of three recombinant *Pseudomonas putida* KT2440 bacterial strains ^[81]. These bacteria can produce rhamnolipids, which are specialty biosurfactants applied in many industrial sectors. Coupling this process with an efficient extraction of toxic aromatic diamines contained in PURs (i.e., TDA), as proposed by Chen et al. ^[93], can be used to realize a green route for the open-loop bio-recycling of PUR degradation products.

4. Prospective for the Biological Degradation of Other Petroleum-Based Plastics

The biodegradation of other recalcitrant petroleum-based plastics (PAs and plastics with only C–C covalent bonds between their building blocks, i.e., PE, PS, PP, and PVC) is challenging; no degradative enzyme acting on the hydrolytic cleavage of the C-C bond is known. Moreover, some of these plastics (PE, PP) are completely crystalline, preventing enzymes from accessing the potential attack sites. Therefore, a multistep reaction catalyzed by a combination of abiotic pretreatments and different types of oxidative enzymes has been proposed ^[94].

There are many different types of PAs (polymers made of repeating units of aliphatic, semi-aromatic, or aromatic molecules linked via amide bonds), the most popular being nylon and Kevlar ^[95]. Since natural silk is also a PA from a chemical point of view, it was expected that enzymes able to degrade synthetic PAs do exist in nature. Unfortunately, to date, no known microorganism able to effectively degrade polymeric PAs has been identified, while it was demonstrated that several bacteria can act on short oligomers of linear or cyclic nylon and are possibly useful for recycling the byproducts of PA production ^{[5][96]}. A more recent study by Biundo et al. ^[97] demonstrated that it is also possible to

engineer PHEs to abolish PET degrading activity and simultaneously increase the catalytic efficiency towards soluble PA oligomers. A manganese-dependent peroxidase from white rot fungus IZU-154 is the only enzyme reported to act on high-molecular-weight nylon fibers ^[98].

It was proposed that PE biodegradation only happens after abiotic (ultraviolet exposure) or enzymatic oxidation by different classes of oxidoreductase (alkane hydroxylase EC 1.14.15.3, laccase or multi-copper oxidase EC 1.10.3.2, peroxygenases EC 1.11.2, or Mn-peroxidase EC 1.11.1.13). This results in the introduction of aldehydic, ketonic, or alcoholic groups, which allows for subsequent processing by other enzymes. The latter can produce intermediate acid oligomers that are intracellularly taken-up by microorganisms and further metabolized ^{[99][100]}. Several microbial strains capable of degrading PE with this proposed system have been reported, including the gut microbiota of invertebrates ^[22] ^[94]; animal phenol oxidase (EC 1.14.15.3) enzymes contained in the saliva of wax worm (*Galleria mellonella*) larvae were also demonstrated to be involved in the oxidative degradation of PE ^[28]. No system studied to date can achieve complete biodegradation without abiotic intervention. Photo-oxidative and thermal pretreatments, although expensive and technically demanding, were shown to oxidize and release oligomers from PE, improving their biodegradation rates ^[101]. In some cases, PE degradation of up to 30% was reported in one month ^{[30][32]}, with even nearly complete bio-assimilation over >8 months ^[102]. In the absence of such pretreatments, the highest biodegradation rates, determined as polymer weight loss, were observed for LDPE films or bags (up to 10% within a few days to one month) ^{[26][29][94]}. These degradation rates were reported only for LDPE materials, while HDPE is less accessible to microorganism attack and much more recalcitrant ^[103].

Although PS is amorphous, its C–C backbone is more resistant to enzymatic cleavage than PE ^[6]. No enzyme has been identified with the ability to efficiently degrade high-molecular-weight PS polymers, despite some studies linking PS degradation to the same laccase and oxidoreductase enzymes associated with PE degradation ^[104]. Some organisms ^[34] ^[35], including invertebrates assisted by their gut microbiome ^[105], were reported to cause significant weight loss in pretreated PS material, with rates of around 10% over one month. Another study reported that *Pseudomonas putida* CA-3 cultures can reach this degradation level in 8 days and produce PHAs. In this case PS was thermally pretreated at an elevated temperature (pyrolysis) to convert it to a styrene oil; this two-step process is very energetically demanding ^[33].

PP polymers can be present in three stereoisomeric forms, namely atactic, isotactic, and syndiotactic [106]. All PP forms are more resistant than PE to heat and to chemical attack, including the action of enzymes [101]. Some studies of PP biodegradation reported weight loss rates of about 10% per month. In all cases, these processes required a strong pretreatment with either y- or UV-irradiation or heat or the use of polymer blends mixing PP and carbohydrates [36][37][107].

Finally, PVC, a synthetic polymer that contains chloride, is a form of waste that poses a serious pollution problem ^[108]. Very few studies have reported the biodegradation of low-molecular-weight PVC oligomers or film ^{[101][109]}. The capacity to degrade PVC was associated with the activity of catalase-peroxidase (EC 1.11.1.21) enzymes in white rot fungi ^[110] or from bacteria found in the gut of insects larvae ^[109]. Nevertheless, the use of microorganisms to remove PVC waste is still controversial because of the possible release of organochlorine compounds that may be highly toxic for the organisms themselves and the environment ^[101].

The fatty acids released by the oxidative bio-depolymerization of C-C plastics cannot be used in a closed-loop process of recycling, but they could be used as a carbon source for microbial growth to produce high-added-value products. While an open-loop fully bio-based process has not been established, in a recent proof-of-principle study, an open-loop recycling process was designed to produce β -ketoadipate or PHAs using microbial strains growing on organic acids obtained from chemically oxidative depolymerization of mixed resins or post-consumer waste (made with PS, HDPE, and PET) ^[111]; although promising, an LCA analysis is required to assess the industrial-scale feasibility of the proposed systems. A recent LCA evaluation focused on the use of fatty acids from chemically depolymerized and oxidized PE as a carbon source for growing a *Pseudomonas putida* KT2440 strain to target PHA production: the proposed approaches were more expensive or had a greater impact on the environment than waste-to-energy approaches ^[112].

5. Perspectives

European targets for plastic waste recycling rates are set by Directive 94/62/EC to 50% and 55% by 2025 and 2030, respectively. These targets are stricter for PPW, requiring that 65% and 70% of PPW be recycled by 2025 and 2030, respectively ^[113]. These targets can be achieved by summing the already recycled PCPW (\approx 20%) with the fraction (18,514 kt y⁻¹, \approx 50% PCPW) that researchers estimated to be potentially included in future recycling flows (**Figure 2**a). The polymer composition of this unrecycled waste plastic flow is reported in **Figure 2**b.



Figure 2. Unrecycled PCPW flows potentially included in future recyclable flows in the EU (Total 18,514 kt y^{-1}): (a) divided by source; (b) divided by polymer composition. Elaborated from data collection and research conducted for this review article. The main contribution (57%) of this fraction is thermoset plastics (29.4% PURs and 27.9% other thermosets) (**Figure 2**a). Therefore, the most urgent interventions should focus on finding alternative recycling options to allow for the production of virgin thermoset materials from waste.

Thermoplastic rejects from MRFs or RECs account for 25% of this unrecycled PCPW flow (**Figure 2**a). Exported plastic waste (10%) was included, as the European Commission has stated that all PPW should be recycled in the EU market by 2030 after the Chinese government's ban (April 2017) on imported recyclable solid waste ^[114]. An improvement in the amount of recycled plastic should be derived from improved sorting and preprocessing to reach higher purity of PCPW fractions at RECs; this is achievable by coupling traditional sorting technologies (i.e., manual sorting and NIR sensors) with machine-learning-based recognition of plastic waste from images ^{[115][116]}. This is expected to only increase the recycling rate in RECs, while thermoplastics rejects generated by mixed and small-particle plastics would remain an issue. This suggests that a fundamental change in the plastic value chain is required for marketable plastic materials to account for end-of-life circularity options in waste treatment processes, especially during the sorting step.

Although environmentally dispersed plastics are reported to be the smallest flow of unrecycled PCPW (18%, **Figure 2**a), they also represent the most complicated fraction to be recaptured (microplastics) and valorized (the quality of macroplastics recovered from the environment is poor). This flow should be lowered as much as possible by improving plastic collection rates through a focus on public behavior and municipal policies aimed at decreasing environmental leakages and progressively increasing the amount of collected plastics to target recycling.

The remining PCPW mixed with non-plastic materials and disposed of as non-plastic unrecyclable waste (targeted to landfilling, incineration, or recovery) was not considered as a potentially recyclable flow.

Considering the necessary interventions, microbial enzyme-based biotechnology could be integrated in the waste management sector and contribute to the introduction of new depolymerization methods for thermosets and rejected thermoplastics, allowing for resynthesis of virgin polymers or the open-loop upcycling of degradation products. Therefore, it is necessary to make biotechnological solutions available that are suitable to treat plastic waste of different quality, including materials made of (or blended with) many different polymers (not necessarily synthetic) that are difficult to separate with traditional methods prior to recycling; moreover, such systems should retain market competitiveness and an environmental impact equal to or lower than that of existing technologies. In this regard, the actual contribution of biotechnology to plastic degradation is limited, as only the bio-based depolymerization systems described for amorphous aliphatic polyesters (such as pretreated PET) and liquid polyester-type PURs have the potential to be exploited at the industrial scale, although not competitive with traditional approaches at present. It will be crucial to improve such systems by further enzyme discovery or engineering studies to increase the degradation efficiency (to overcome the *passivation* issue) under high PET load or for less pretreated materials.

Even if PET is, by far, the most consumed polyester, PET waste represents only a minor fraction (\approx 7%) of unrecycled flows with respect to PURs, thermosets, and other non-polyester plastics (**Figure 2**b). A synergy with promising methods of abiotic secondary and tertiary PUR depolymerization approaches could allow for depolymerization of recalcitrant PUR waste, but no method has been tested on real PUR waste to date [88][91][117][118][119]. PAs and C-C backbone plastics, such as PE and PS, are currently much less efficiently biodegraded. Moreover, some of these plastics (i.e., PVC) may release toxic components after their degradation. Although biodegradation could be enhanced by new methods of abiotic preprocessing in the presence of organometallic catalysts, these treatments require a lot of energy and relatively high temperatures (>150 °C), thus contributing to higher costs and environmental impact ^{[6][120]}. Moreover, as most biodegradation processes have only been conducted at laboratory scale ^{[99][101]}, without biochemical information on the kinetics of the involved reactions, with scarce data on degradation products, and often observing that mainly additives

were cleaved [121], a fully bio-based depolymerization of non-polyester petroleum-based plastics is predicted not to be economically and environmentally competitive with traditional approaches within a 10-year time frame. Instead, it is necessary to look at new approaches of tertiary recycling [122][123][124] and to use them in combination with microbial enzymes to convert degradation products into high-added-value chemicals [111][125][126].

The difficulties to apply enzymes for depolymerizing non-polyester plastic waste fractions indicate that particular attention should be taken on aliphatic polyesters (i.e., PEF, PBAT, and PBS) to replace traditional recalcitrant petroleum-based plastic waste. Regardless of whether these plastics are biodegradable or bio-based, researchers argue that the most important aspect is the creation of suitable technologies to allow for their bio-based degradation and circular production from waste within an industrial setup, while avoiding environmental leakages.

In conclusion, the use of enzymatic microbial biotechnologies for plastic degradation and valorization requires a rethink of the whole plastic value chain, as well as the development and marketing of alternative plastic materials (made with non-food biomass derivatives or aliphatic polyesters that have a biodegradability similar or superior to that of PET), the redesign of plastic packaging and products, the improvement of collection and sorting methods, and their tight integration with new, eventually chemo-assisted bio-based systems of degradation and recycling (competitive from an economical point of view and with a lower environmental impact compared to available technologies). This, in turn, will require stronger co-operation between biotechnologists and stakeholders involved with feedstock resources, material production, and waste management to find novel effective industrial technologies and regulations for a transition to a circular (bio)economy in plastic waste management.

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