

Polystyrene shaping effect on the enriched bacterial community from the plastic-eating *Alphitobius diaperinus* (Insecta: Coleoptera)

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Abstract

Plastic pollution has become a serious issue of global concern, and biodegradation of plastic wastes is representing one attractive environment-friendly alternative to traditional disposal paths. It is known that insects are involved in the plastic polymer degradation process, with reported evidence of tenebrionid beetle larvae capable to degrade polystyrene (PS), one of the most used plastics worldwide. Recently, a ribosomal RNA based survey on the insect gut microbiota of the lesser mealworm *Alphitobius diaperinus* has revealed differentially abundant microbial taxa between PS-fed larvae and control group. Following these findings, an enrichment bacterial culture was set up in liquid carbon-free basal medium with PS film as sole carbon source using PS-fed larvae of *A. diaperinus* as inoculum. After two-months the culture was analysed both by molecular and culture-based methods. Isolated bacteria which had become prevalent under the selective enrichment conditions resulted ascribable to three taxonomic groups: *Klebsiella, Pseudomonas*, and *Stenothrophomonas*. The predominance of these groups in PS-fed larvae was confirmed by using bacterial 16S rRNA gene amplicon sequencing, and it was consistent with the results of previous reports. Isolated bacteria were able to attach to PS surfaces and SEM observations showed the presence of thin fibrillar structures connecting the bacterial cells to the abiotic surface.

Keywords Larval microbiome · Polystyrene · Enrichment culture · 16S rRNA · Alphitobius diaperinus

1 Introduction

Plastics are represented by a large family of polymeric materials deriving from hydrocarbons that, since 1950s, offer innovative solutions to human-society requirements. In 2019 global plastic production reached 368 million tons with Europe accounting for 16% of totality (Plastics Europe 2020). Among the most popular plastics in the world, polystyrene (PS) is still representing the fifth most used thermoplastic in Europe due to its excellent processability, low cost, resistance and insulating properties (Plastics Europe 2020); consequently, its end-of-life disposal is a source of particular concern. Nowadays, PS recycling rates are still low at global level (Takada and Bell 2021), and a consistent portion of

PS waste ends up in the environment where its degradation is a complex and slow process which depends on chemical structure of the polymer and environmental conditions (Ho et al. 2018; Ali et al. 2021). The high durability of PS in the environment and its progressive fragmentation into submillimetre-sized particles can have a negative impact on different levels of biological complexity (Hollóczki and Gehrke 2019). Recently, a great interest has been directed towards some insect species involved in PS biodegradation with lepidopteran species, such as the greater wax moth Gal*leria mellonella* (Jiang et al. 2021), and coleopteran species, such as the superworm Zophobas atratus (Peng et al. 2020), the dark mealworm Tenebrio obscurus (Peng et al. 2019), the yellow mealworm Tenebrio molitor (Yang et al. 2015a, 2015b) and the red flour beetle Tribolium castaneum (Wang et al. 2020), as candidate taxa to be tested for their ability to process plastics. Mealworms and superworms are provided with basic mandibulate mouthparts that enable them to chew and ingest the polymer. Moreover, suppression of their gut microbes with the antibiotic gentamicin indicated a gut-microbial dependence for PS biodegradation (Peng et al. 2019, 2020; Yang et al. 2021). A recent study carried

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out on another coleopteran species, Alphitobius diaperinus, concluded that also this beetle is capable of eating and degrading PS. In this species, comparative analyses of the gut microbiome have revealed several microbial taxa to be differentially abundant between PS-fed larvae and control groups, thus providing an initial list of microbial candidates involved in the degradation process (Cucini et al. 2020). The lesser mealworm A. diaperinus is a nocturnal cosmopolitan Tenebrionidae, commonly found in litter, manure and chicken coops. In chicken farms, the beetle feeds on manure, spilled feed, cracked eggs, and bird carcasses. A. diaperinus complete life-cycle lasts from 40 to 100 days. Adults, 5-8 mm long, usually lay their eggs (1.5 mm in length) in litter, droppings and grain hulls; larvae can reach 7-11 mm in length and, during the last larval instar, they leave the litter and make galleries into thermal insulation materials, frequently including PS, in order to construct pupal cells (Dunford and Kaufman 2006).

The aim of this study was to further expand the work started by Cucini et al. (2020) focusing on the gut-associated bacterial communities of PS-fed larvae of *A. diaperinus*. An enrichment procedure in liquid carbon-free basal medium with PS films as sole carbon source starting from PS-fed larvae of *A. diaperinus* was performed and analysed in order to (i) compare the composition of enriched communities by culture-dependent and culture-independent molecular methods and to (ii) obtain a collection of bacteria potentially exploitable for synthetic microbiota application.

2 Materials and methods

2.1 Insect material

Last instar larvae of *A. diaperinus* were purchased online from Agripet Garden (https://www.agripetgarden.it/; Padua, IT). Three-hundred grams (approximately 1700 individuals) were reared under controlled conditions $(20 \pm 2 \,^{\circ}\text{C}; 12 \text{ L}:12D)$ photo-period; 50–70% RH) and fed with 20 g of PS as unique source of carbon. The PS foam (0.01 g/cm³), known as Extir® (CAS 9003-53-6), was purchased from Toscoespansi s.r.l. (http://www.toscoespansi.it/). According to the manufacturer information, no extra additives or catalysts were present. After 30 days, 50 larvae were collected, immersed 1 min in 75% ethanol and washed in Saline Water (SW) [0.85% NaCl] for the same time, then intestines were sterilely dissected under a Leica Wild M3C stereomicroscope.

2.2 Enrichment culture and bacteria isolation

Enrichment culture was established as essentially reported by Yang et al. (2014). Dissected intestines were harvested by centrifugation ($7000 \times g$, 20 min), and washed twice in SW. The pellet was resuspended in 40 ml of Liquid Carbon-Free Basal medium (LCFB) [0.7 g of KH₂PO₄, 0.7 g of K₂HPO₄, 1.0 g of NH₄NO₃, 0.7 g of MgSO₄·7H₂O, 0.005 g of NaCl, 0.002 g of FeSO₄·7H₂O, 0.002 g of ZnSO₄·7H₂O, and 0.001 g of MnSO₄·H₂O in 1000 ml of distilled water, pH 7.2] containing PS films 0.0125 g/ml. PS films were prepared dissolving 3.5 g of PS foam in 116 ml of trichloromethane (0.03 g/ml), then the solution was poured on glass plates and dried overnight under chemical fume hood (Yang et al. 2015b; Sekhar et al. 2016). Films were cut $(10 \text{ mm} \times 10 \text{ mm}; 9-10 \text{ mg/each})$, washed once with 40 ml of ethanol (>99%) and three times with sterile water then sterilized, both sides, under UV light. Serial dilutions of the suspension were performed in SW and spread-plated on Luria Bertani Agar (LBA) [10 g of bacto tryptone, 5 g of yeast extract, 5 g of NaCl and 16 g of agar to 1000 ml of distilled water, pH 7.2]. The plates were incubated under aerobic conditions at 27 °C for 24-48 h, and the total number of viable cells was determined. The enrichment culture was incubated on a shaker with gentle agitation (90 rpm) under aerobic conditions at room temperature (20-22 °C) for 60 days under natural light conditions. From the two-months old culture aliquots were prepared and stored at -80 °C for molecular analysis; at the same time, serial dilutions were performed in SW and spread-plated on LBA for bacterial enumeration and strain isolation. Plates were then incubated as reported above. Individual colonies representing isolates with different morphologies were selected according to differences in colour, size and appearance. They were picked from the different dilutions LBA plates and sub-cultured at least three times before identification. Bacterial isolates were then stored in LB broth with 30% glycerol (v/v) at -80 °C.

2.3 Phenotypic and phylogenetic characterization of bacterial isolates

Gram determination was carried out either by staining (Gram stain kit, Carlo Erba Reagents) or chemical method (Powers 1995). Catalase and oxidase activity were determined according to Smibert and Krieg (1981). The level of salt tolerance of the bacterial isolates was determined in LB broth supplemented with various concentrations of NaCl (5, 7.5, and 10%) inoculated with overnight grown bacteria and incubated at 27 °C for 48 h. LB broth without extra addition of NaCl was maintained as control. Optical density at 600 nm was measured using UV-Vis spectrophotometer (Novaspec, Amersham Biosciences, Amersham, UK). Sensitivity of the isolates to common antibiotics (all purchased from Sigma-Aldrich) was tested by spotting 4 µl drops of overnight cultures on LBA plates supplemented with carbenicillin (100 µg/ml), gentamicin (30 µg/ml), kanamycin (50 µg/ml), and antibiotic-free (control). Results of bacterial growth after incubation at 27 °C for 24-48 h were evaluated in comparison to positive (antibiotic-free) control. The ability of the isolates to produce lipolytic activity was tested on Tributyrin agar (Oxoid) [5 g of peptone, 3 g of yeast extract, 10 g of glyceryl tributyrate and 15 g of agar to 1000 ml of distilled water, pH 7.5]. Production of urease, lysine decarboxylase, ornithine decarboxylase, tryptophanase, hydrogen sulphide and acetoin, acid production from glucose, adonitol, lactose, arabinose, sorbitol and dulcitol, and assimilation of citrate were determined by the EnteroPluri-Test (Liofilchem®) according to the manufacturer's instructions. All assays were repeated at least twice.

From bacterial cells grown overnight in LB, total DNA was extracted using the kit Wizard® SV Genomic DNA Purification System (Promega Corporation, Madison, WI, US). For each DNA sample the amplification of 16S rRNA gene was performed using universal primers: 16S_V3V4-341F (5'-CCTACGGGNGGCWGCAG-3') and 16S_V3V4-785R (5'-GACTACHVGGGTATCTAATCC-3') [Klindworth et al. 2013]. PCRs were executed in 25 μ l reaction volume with 2.5 µl of DNA from each sample, 1.25 µl of both forward and reverse primers (10 mM), 2.5 µl of MgCl₂ (25 mM), 2.5 µl of deoxynucleotides (10 mM), 5 µl of Green GoTag Flexi Buffer (Promega, US), 0.125 µl of GoTaq Flexi DNA polymerase (Promega, US), and 9.875 µl of ddH₂O. Amplifications were obtained in a GeneAmp® 2700 Thermal Cycler (Applied Biosystem, Foster City, CA, US), using the following conditions: (i) 95 °C for 3 min; (ii) 35 cycles of 95 °C for 1 min, 50 °C for 1 min, 72 °C for 90 sec; (iii) 72 °C for 5 min. The kit Wizard® SV Gel and PCR Clean-Up System (Promega Corporation) was employed for PCR products purification and their DNA concentration was assessed through Nanodrop ND1000UV vis device (NanoDrop Technologies, Wilmington, DE, US). Sanger sequencing was executed on a DNA Analyzer ABI 3730 at the BioFab Research core facility (Rome, IT). Electropherograms were manually checked with the Sequencher 4.4.2 software (Gene Codes, Ann Arbor, MI, US). The resulting sequences were then aligned against GenBank (NCBI) through the nucleotide Basic Local Alignment Search Tool - BLASTn (Altschul et al. 1990).

The sequence data were submitted to the DDBJ/EMBL/ GenBank databases under accession numbers: OL470964, OL470965, and OL470969.

2.4 Sequencing of 16S rRNA amplicons

Total DNA extractions were carried out on three technical replicates from aliquots of the enrichment culture, using the Wizard® Genomic DNA Purification Kit according to manufacturer's protocol. Final DNA concentrations were determined by NanoDrop ND1000V and Qubit® fluorometer (Life Technologies Corporation, Carlsbad, CA, US), while DNA quality was evaluated through 1% agarose gel electrophoresis. NGS metabarcoding was conducted with the same set of primers (V3V4: 341F-785R) used for Sanger sequencing at the BioFab Research core facility (Rome, IT). PCR reactions were performed in a 25 µl volume with 2.5 µl DNA, 5 µl 5x PCRBIO Reaction Buffer, 1 µl of both primers (10 µM), 0.25 µl PCRBIO HiFi Polymerase, 15.25 µl ddH2O with the following conditions: (i) 95 °C for 3 min; (ii) 25 cycles of 95 °C for 30 sec, 55 °C for 30 sec, 72 °C for 30 sec; (iii) 72 °C for 5 min. PCR products were purified with AMPure XP Beads (Beckman Coulter, High Wycombe, UK) according to the manufacturer's specifications and each purification product was indexed using the Nextera XT Index Kit (Illumina) for library production. The latter reaction was carried out using the following conditions: (i) 95 °C for 3 min; (ii) 8 cycles of 95 °C for 30 sec, 55 °C for 30 sec, 72 °C for 30 sec; (iii) 72 °C for 5 min. Indexed samples were purified with Ampure XP Beads according to the PCR Clean-Up protocol suggested by Illumina; resulting products were quantified using a Bioanalyzer DNA 1000 (Agilent Technologies, Santa Clara, CA, US) and pooled in equal amounts for sequencing. A MiSeq Illumina sequencer was used to perform paired-end sequencing. Raw data, which amounted to a total of 207,858 reads (ranging from 56,899 to 83,230 per replicate) were filtered on a quality basis and primers trimmed using trimmomatic v.0.39 tool (Bolger et al. 2014). To investigate enriched bacterial community composition, sequences were processed using Quantitative Insights Into Microbial Ecology 2 (QIIME2 v.2021.4) (Boylen et al. 2019). In accordance with the software pipeline, through DADA2 plugin, already demultiplexed sequences were joined, filtered and dereplicated (Callahan et al. 2016). The vsearch plugin was employed both for chimera removal and clustering in the different Operational Taxonomic Units (OTUs) at 97% using the de novo strategy (Rognes et al. 2016). Furthermore, singletons and contaminants (i.e. mitochondrial and chloroplast sequences) were discarded. To train the classifier artifacts for taxonomic attribution of 16S sequences, the SILVA 132 database was employed (Quast et al. 2013), using feature-classifier plugin (Bokulich et al. 2018) [Protocol S1]. Resulting sequences were matched with those of Cucini et al. (2020) dataset though the NCBI BLAST+ tools (Camacho et al. 2009). Taxonomic distribution results were visualized QIIME2 view and also compared with those of the first 16S rRNA survey on PS-fed A. diaperinus larvae (Cucini et al. 2020). Raw sequencing reads were deposited in the Sequence Read Archive (SRA, NCBI) with the Bio-Project ID PRJNA781815.

2.5 Analysis of adhesion and growth of bacterial isolates in LCFB with PS

For each bacterial strain, cells were collected from overnight cultures in LB by centrifugation $(10,000 \times g, 5 \text{ min})$, washed three times and suspended in sterile SW. A bacterial inoculum to give a final concentration of 5×10^5 Colony Forming Unit (CFU) per millilitre (ml) was added to sterile tubes containing 4 ml each of LCFB or Phosphate buffered saline (PBS) with 0.0125 g/ml PS films (10 mm × 10 mm). Bacterial suspensions were statically incubated under aerobic conditions at room temperature (20–22 °C) for 28 days.

For SEM observation, one PS film was collected from each culture and washed 1 h in 8 ml PBS on a shaker (90 rpm) to remove unattached cells. Films were fixed in 2.5% glutaraldehyde for 2 h and post-fixed in 1% osmium tetroxide for 1 h (Dallai et al. 2016). They were later washed in PBS and then dehydrated in a graded series of ethanol (50–100%), dried after terbutilic alcohol sublimation, and gold-coated with a Balzers MED 010 sputtering device. Samples were examined and photographed through a FEI Quanta 400 Scanning electron microscope operating at 20 kV.

For mechanic and enzymatic assay, the procedure reported by Banar et al. (2016) was modified as follows: three PS films from bacterial suspensions from both LCFB and PBS (control) were collected after 3 h and 28 days, respectively. Each film was individually transferred in a tube containing 8 ml of sterile PBS and left 1 h on a shaker (90 rpm). PBS was then removed and replaced with 1 ml of PBS containing 1 mg of trypsin (Sigma-Aldrich), vortexed 1 min, incubated 1 h at 37 °C, vortexed again for 1 min. Viable counts were determined through serial dilutions that were spread-plated on LBA. Plates were incubated at 27 °C for 24–48 h.

3 Results and discussion

3.1 Phenotypic and phylogenetic characterization of bacterial isolates

The liquid enrichment from PS-fed larvae of A. diaperinus in a carbon-free basal medium with PS film as sole carbon source was examined by the viable plate method at the beginning and after two months of incubation at the reported conditions. The estimated number of CFU/ml at the end of the trial showed that viable cells concentration increased from $\sim 6.0 \times 10^6$ to $\sim 1.5 \times 10^8$ CFU/ml (Table S1), most likely indicating that few bacteria became "enriched" by asserting their dominance through growth in such minimal medium over the considered period. In total three different morphologies were observed among colonies grown after 48 h in LBA from the two-months PS-enriched culture. Isolates representative of each morphotype were named 2 m/a, 2 m/b, and 2 m/c (Table S2). All three isolates resulted Gram-negative rod-shaped bacteria, and their phenotypic characterization, based on various biochemical and physiological parameters,

is shown in Table 1. Within the set of whole determined phenotypic characteristics, some distinctive phenetic traits were considered by referring to a simplified dichotomic scheme for the identification of Gram-negative rods based on Bergey's Manual of Determinative Bacteriology (Holt 1994; Fig. S1). Following this scheme, strain 2 m/a was included among lactose-fermenting Enterobacteriaceae; 2 m/b and 2 m/c, both unable to ferment glucose and lactose, but being different with respect to cytochrome oxidase production, were included within oxidase-positive (Pseudomonas), and oxidase-negative group (Acinetobacter, Burkholderia, Stenotrophomonas), respectively. Furthermore, the three isolates showed different level of resistance to a beta-lactam and aminoglycosides, a common trait among Klebsiella pneumoniae, Pseudomonas aeruginosa, and Stenotrophomonas maltophilia species which were included in the top ten list of multidrug-resistant bacteria (Rello et al. 2019). Crippen and Poole (2009) reported that the gastrointestinal tract of lesser mealworm larvae is capable of supporting the occurrence of

 Table 1 Differential physiological and biochemical characteristics of bacterial strains isolated from PS-fed larvae of *Alphitobius diaperinus* after two-months enrichment on carbon-free medium with PS as sole carbon source

Characteristics	Bacterial isolates		
	2 m/a	2 m/b	2 m/c
Growth with:			
NaCl (7.5%, w/v)	+	-	-
Carbenicillin (100 µg/ml)	W	+	w
Gentamicin (30 µg/ml)	W	-	+
Kanamycin (50 µg/ml)	-	+	_
Production of:			
Catalase	+	+	+
Cytochrome oxidase	-	+	-
Urease	-	+	+
Lysine decarboxylase	+	-	_
Ornithine decarboxylase	+	-	-
Tryptophanase	-	-	-
Hydrogen sulphide	-	-	-
Acetoin	+	-	_
Lipase	W	+	+
Acid production from:			
Glucose	+	-	_
Adonitol	+	-	_
Lactose	+	-	_
Arabinose	+	+	_
Sorbitol	+	-	_
Dulcitol	-	-	-
Assimilation of:			
Citrate	+	+	+

+, positive reaction; w, weakly positive; -, negative reaction

horizontal transfer of antibiotic resistance genes between different species of bacteria under laboratory conditions, therefore *A. diaperinus*, as well as other edible and household insects, could constitute a reservoir of transferable resistance genes of human health concern (Gwenzi et al. 2021).

A partial sequence of 16S rRNA gene (464–475 bp) was determined for each of three bacterial isolates. A BLAST search with the 16S rDNA sequences against NCBI nucleotide databases, indicated that 2 m/a and 2 m/b shared 99.65% identity with *Klebsiella aerogenes* ATCC 13048 (GenBank: NR_118556.1; Zhang et al. 2014) and *Pseudomonas aeruginosa* DSM 50071 (GenBank: NR_117678.1; Swiderski 2012), respectively; while 2 m/c showed 97.59% identity with *Stenotrophomonas maltophilia* ATCC 13637 (Gen-Bank: MT354120.1; Reid et al. 2021).

Members belonging to genus *Klebsiella, Pseudomonas* and *Stenotrophomonas* have been already found associated with plastic compounds and proposed as plastic eaters (Brandon et al. 2021), and, more specifically, in association to *Z. atratus* (Kim et al. 2020), and *T. molitor* (Urbanek et al.

2020; Brandon et al. 2021). The repeated finding of similar bacteria associated to PS-fed larvae strengthens then the hypothesis of their involvement in PS degradation process in *A. diaperinus* as well as in the other plastic-eater insects so far investigated.

3.2 Sequencing of 16S rRNA amplicons

From the Miseq Illumina sequencing of the 16S molecular marker (V3-V4 regions) a total amount of 207,858 reads, ranging from 56,899 to 83,230 per library, were obtained. After filtering, dereplication, clustering, singletons and chimeras' removal, the three technical replicates had respectively 26,580, 24,920 and 36,302 sequences, for a total of 87,802, and 57 OTUs. Sequences shared an average length of 399 nucleotides. At family level, Xanthomonadaceae, Pseudomonadaceae and Enterobacteriaceae were largely dominant in the sample with regard to other taxa (Fig. 1), with the former owning the highest relative abundance. At genus level, two taxa predominated





Fig. 1 16S rRNA-based analysis of the enrichment culture from PSfed *Alphitobius diaperinus*' gut. Taxa bar plots exhibit OTUs diversity of bacteria at (**a**) Family level: left, present study; right, based on data from Cucini et al. (2020); (**b**) Genus level, showing their relative

abundance in the three technical replicates (Meta1, Meta2, Meta3). Taxa are colour coded. In (a), OTUs with a relative abundance smaller than 1% were grouped and displayed as 'Other'

over the others, namely *Stenotrophomonas* and *Pseudomonas* (Fig. 1), with an average relative abundance of above 41% and 35.5%, respectively.

Partial 16S rRNA gene sequences from strains 2 m/a, 2 m/b and 2 m/c were aligned on the 16S rRNA gene amplicons dataset, and all three found a meaningful correspondence (>99% identity). However, taxonomical association of the strain 2 m/a, namely Klebsiella sp., was assessed only up to OTU-family level - i.e. Enterobacteriaceae - probably due to a limitation of SILVA database used in the QIIME2 pipeline. This observation also explains the absence of Klebsiella genus in the taxonomic barplot (Fig. 1). From the initial 16S rRNA survey on A. diaperinus larval microbiome reported by Cucini et al. (2020), a total amount of 295 different OTUs were retrieved with only six as differentially abundant in PS group, namely: two Pseudomonas sp., Cronobacter sp., Kocuria sp., Pseudogracilibacillus sp. and Virgibacillus sp.. BLAST analysis of 16S rRNA gene sequences from strains 2 m/a, 2 m/b and 2 m/c against libraries from Cucini et al. (2020) retrieved significant matches (> 99% identity). Sequence of 16S rRNA gene from the three bacteria isolated in this study matched with OTUs that had a higher relative abundance in the previously reported PS libraries, meaning that they were also present with higher abundance in PS-fed A. diaperinus larvae (Fig. 1).

The alignment of 16S rRNA sequences obtained in this study against sequences differentially abundant in *A. diaperinus*'s PS-fed libraries from Cucini et al. (2020) provided confirmation of the presence of *Lactococcus*, *Pseudomonas* spp., and a group of Enterobacteriaceae (i.e. *Klebsiella* family). Other Enterobacteriaceae and *Stenotrophomonas* were also present in PS libraries, but their higher relative abundance with respect to control libraries was not significant by statistical analysis, (p-adj > 0.05), and then not indicative of

differentially abundant bacterial taxa in the insect PS-fed group (Fig. 1).

3.3 Analysis of adhesion and growth of bacterial isolates in LCFB with PS

The ability of isolates *Klebsiella* sp. 2 m/a, *Pseudomonas* sp. 2 m/b, and *Stenotrophomonas* sp. 2 m/c to grow on PS films was evaluated by SEM observation and enzymatic/ mechanical assay.

SEM observations were carried out on PS films from 28 days-old liquid cultures in carbon-free mineral medium. Bacterial cells on PS films appeared healthy, rod-shaped with smooth intact surface. Moreover, the used magnification showed thin fibrillar structures connecting the bacterial cells to the abiotic surface (Fig. 2). For *Stenotrophomonas* sp., few bacterial aggregates were also observed (Fig. 2).

The number of attached viable cells on PS films after 3 h and 28 days incubation in mineral medium and phosphatebuffer was recorded for all strains. After 3 h no significative difference was observed between LCFB and PBS by considering the number of detached viable cells with enzymatic/mechanical treatment. After 28 days no viable cells of Klebsiella sp. 2 m/a and Stenotrophomonas sp. 2 m/c were recovered from PS films kept in PBS, on the contrary Pseudomonas sp. 2 m/b did not show any decline in viability accordingly with what was already stated for other members of the same genus which were found able to survive in PBS for at least 30 weeks (Liao and Shollenberger 2003). An increase of the number of viable cells detached from PS films maintained in LCFB was observed for all strains after 28 days when compared to the number of viable detached after 3 h, but the only significant difference (p value < 0.05) was that exhibited by Stenotrophomonas sp. 2 m/c (Fig. 3a; Table S3).



Klebsiella sp. 2m/a

Pseudomonas sp. 2m/b

Stenotrophomonas sp. 2m/c

Fig. 2 SEM analysis of bacterial adhesion to PS films. White arrows show the presence of thin fibrillar structures connecting bacterial cells to abiotic surface after 28 days incubation in LCFB medium



Fig. 3 Analysis of bacterial adhesion to PS film after enzymatic/ mechanic assay. For each strain, histograms show viable counts after 3 h and 28 days (dd) incubation in LCFB medium and PBS, respectively. **a:** Cells detached from PS films; **b:** Planktonic cells. Error bars represent standard deviation. Effect sizes are presented in Table S3 and in Table S4

The viability of cells in planktonic form was also determined. The patterns of population changes in planktonic state were not significantly different in LCFB after 3 h and 28 days of incubation (Fig. 3b; Table S4). As already observed for the attached state, no viable planktonic cells were recovered from PBS for Klebsiella sp. 2 m/a and Stenotrophomonas sp. 2 m/c after 28 days; nonetheless, Klebsiella sp. 2 m/a, even if not increased significantly its concentration, survived well in mineral medium both in planktonic and attached state until the end of the observation period (Fig. 3b; Table S4). Results indicate that a pure culture made up of individuals of one species only, with the possible exception of Stenotrophomonas sp. 2 m/c (Fig. 3a; Table S3), does not show any significant change in LCFB with PS after an incubation period of four weeks. It must be highlighted that the approach followed in this study ignores the possibility that PS biodegradation may result from the cooperative action of different microbial species under natural conditions; in addition, the metabolic process of a single species may lead to the accumulation of intermediates or dead-end products with potentially toxic effects on bacterial growth.

4 Conclusion

This is the first study where culture-dependent and molecular analyses have been combined to the analysis of the bacterial population associated with plastic-eating A. diaperinus (Insecta: Coleoptera) after an enrichment phase in a medium with limited chemical composition and PS as sole carbon source. The results obtained indicate the predominance of three taxonomic groups, Klebsiella, Pseudomonas, and Stenothrophomonas, either by culture-dependent and -independent analysis of the enriched communities. Nonetheless, bacterial isolates of each taxon, when grown as monoculture in a synthetic medium with PS, revealed a weak metabolic activity. In nature, PS degradation process could be influenced by the interactions between members of a complex microbial community, the higher genotypic and phenotypic diversity present in microbial communities compared with a single strain population may mean that mixed communities are more efficient degraders of the pollutant. Therefore, the lack of cooperative microbial interactions at the applied experimental conditions could have been constraining. Notwithstanding these limitations, the study suggests that bacterial isolates could be representative members of microbial taxa involved in PS degradation as already reported (Brandon et al. 2021). Those findings could represent a valid starting point for the generation of synthetic bacterial communities to be used for microbiome transfer to gnotobiotic insects in order to better understand the web of interactions involving both heterospecific symbionts and the animal host during the process of PS degradation.

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Data availability All data generated or analysed during this study are included in this published article.

Declarations

Competing interests The authors have no competing interests to declare that are relevant to the content of this article.

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