Thermophilic Carboxylesterases from Hydrothermal Vents of the Volcanic Island of Ischia Active on Synthetic and Biobased Polymers and Mycotoxins

Distaso, Marco; Chernikova, Tatyana; Bargiela, Rafael; Coscolín, Cristina; Stogios, Peter J.; Gonzalez-Alfonso, Jose L.; Lemak, Sofia; Khusnutdinova, Anna; Plou, Francisco J.; Evdokimova, Elena; Savchenko, Alexei; Lunev, Evgenii; Yakimov, Michail M; Golyshina, Olga; Ferrer, Manuel; Yakunin, Alexander; Golyshin, Peter

Applied and Environmental Microbiology

DOI: 10.1128/aem.01704-22

Published: 28/02/2023

Peer reviewed version

Cyswllt i'r cyhoeddiad / Link to publication

Dyfyniad o'r fersiwn a gyhoeddwyd / Citation for published version (APA): Distaso, M., Chernikova, T., Bargiela, R., Coscolín, C., Stogios, P. J., Gonzalez-Alfonso, J. L., Lemak, S., Khusnutdinova, A., Plou, F. J., Evdokimova, E., Savchenko, A., Lunev, E., Yakimov, M. M., Golyshina, O., Ferrer, M., Yakunin, A., & Golyshin, P. (2023). Thermophilic Carboxylesterases from Hydrothermal Vents of the Volcanic Island of Ischia Active on Synthetic and Biobased Polymers and Mycotoxins. *Applied and Environmental Microbiology*, *89*(2), e0170422. https://doi.org/10.1128/aem.01704-22

Hawliau Cyffredinol / General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal ?

Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

1 Thermophilic carboxylesterases from hydrothermal vents of the volcanic island of

Marco A. Distaso^a, Tatyana N. Chernikova^a, Rafael Bargiela^a, Cristina Coscolín^b, Peter

Stogios^c, Jose L. Gonzalez-Alfonso^b, Sofia Lemak^c, Anna N. Khusnutdinova^a, Francisco J.

Plou^b, Elena Evdokimova^c, Alexei Savchenko^{c,d}, Evgenii A. Lunev^{a,e}, Michail M. Yakimov^f,

- 2 Ischia active on synthetic and biobased polymers and mycotoxins
- 3

4

5

6

- Olga V. Golyshina^a, Manuel Ferrer^{b,‡}, Alexander F. Yakunin^{a,c,‡}, and Peter N. Golyshin^{a,*,‡} 7 8 ¹Centre for Environmental Biotechnology, School of Natural Sciences, Bangor University, 9 Bangor, UK 10 ^bDepartment of Applied Biocatalysis, ICP, CSIC, Madrid, Spain 11 12 ^cDepartment of Chemical Engineering and Applied Chemistry, University of Toronto, Toronto, Canada 13 ^dDepartment of Microbiology Immunology and Infectious Diseases. University of Calgary, 14 Calgary, Canada 15 ^eInstitute of Gene Biology, Russian Academy of Sciences, Moscow, Russia 16 ^fInstitute of Polar Sciences (ISP), CNR, Messina, Italy 17 18 19 **‡Equal contributions.** 20 *Corresponding authors at: the Centre for Environmental Biotechnology, Bangor University, LL57 2UW Bangor, UK (P.N. Golyshin); ICP, CSIC, Marie Curie 2, 28049 Madrid, Spain (M. Ferrer). 21 22 23 E-mail addresses: mferrer@icp.csic.es (M. Ferrer), p.golyshin@bangor.ac.uk (P.N. Golyshin) 24 Phone numbers: +34915854872 (M. Ferrer), +441248383629 (P.N. Golyshin) 25 26 27 28 29
- KEYWORDS: Thermophilic bacteria, hydrothermal vents, Ischia, metagenome screening,
 carboxylesterase, polyesterase, 3PET, PLA, biochemical characterisation, crystal structure
- 32
- 33

34 ABSTRACT

35 Hydrothermal vents are geographically widespread and host microorganisms with robust 36 enzymes useful in various industrial applications. We examined microbial communities and carboxylesterases of two terrestrial hydrothermal vents of the volcanic island of Ischia (Italy) 37 predominantly composed of Firmicutes, Proteobacteria and Bacteroidota. High-temperature 38 39 enrichment cultures with the polyester plastics polyhydroxybutyrate (PHB) and polylactic acid (PLA) resulted in an increase of Thermus and Geobacillus spp., and to some extent, 40 Fontimonas and Schleiferia spp. The screening at 37-70°C of metagenomic fosmid libraries 41 from above enrichment cultures identified three hydrolases (IS10, IS11 and IS12), all derived 42 from yet uncultured Chloroflexota and showing low sequence identity (33-56%) to 43 44 characterized enzymes. Enzymes expressed in Escherichia coli exhibited maximal esterase 45 activity at 70-90°C, with IS11 showing the highest thermostability (90% activity after 20 min 46 incubation at 80°C). IS10 and IS12 were highly substrate-promiscuous and hydrolysed all 51 monoester substrates tested. Enzymes were active with PLA, polyethylene terephthalate 47 48 model substrate, 3PET, and mycotoxin T-2 (IS12). IS10 and IS12 had a classical α/β 49 hydrolase core domain with a serine hydrolase catalytic triad (Ser155, His280, Asp250) in their hydrophobic active sites. The crystal structure of IS11 resolved at 2.92 Å revealed the 50 51 presence of the N-terminal β -lactamase-like domain and C-terminal lipocalin domain. The 52 catalytic cleft of IS11 included catalytic Ser68, Lys71, Tyr160, and Asn162, whereas the lipocalin domain enclosed the catalytic cleft like a lid contributing to substrate binding. Our 53 54 study has identified novel thermotolerant carboxylesterases with a broad substrate range, 55 including polyesters and mycotoxins, for potential applications in biotechnology.

56

57 IMPORTANCE

High-temperature-active microbial enzymes are important biocatalysts for many industrial 58 applications including recycling of synthetic and biobased polyesters increasingly used in 59 60 textiles, fibres, coatings and adhesives. Here, we have discovered three novel thermotolerant 61 carboxylesterases (IS10, IS11 and IS12) from high-temperature enrichment cultures from 62 Ischia hydrothermal vents incubated with biobased polymers. The identified metagenomic 63 enzymes originated from uncultured Chloroflexota and showed low sequence similarity to known carboxylesterases. Active sites of IS10 and IS12 had the largest "effective volumes" 64 among the characterized prokaryotic carboxylesterases and exhibited high substrate 65 promiscuity, including hydrolysis of polyesters and mycotoxin T-2 (IS12). Though less 66 promiscuous compared to IS10 and IS12, IS11 had a higher thermostability with high 67 temperature optimum (80-90 °C) for activity, hydrolysed polyesters, and its crystal structure 68 revealed an unusual lipocalin domain likely involved in substrate binding. The polyesterase 69 70 activity of these enzymes makes them attractive candidates for further optimisation and 71 potential application in plastics recycling.

72

73 INTRODUCTION

74 Environmental microbial communities and microorganisms represent an enormous reserve of 75 biochemical diversity and enzymes for fundamental research and applications in 76 biotechnology (1,2). However, the vast majority of environmental microbes have never been 77 grown and characterised in the laboratory (3,4). The metagenomic approach has emerged as a 78 strategic way to study unculturable microorganisms and their enzymes using various 79 computational and experimental methods (5-7). Metagenomics includes shotgun sequencing 80 of microbial DNA purified from a selected environment, high-throughput screening of 81 metagenomic expression libraries (functional metagenomics), profiling of RNAs and proteins 82 produced by a microbial community (meta-transcriptomics and meta-proteomics), and identification of metabolites and metabolic networks of a microbial community (meta-83 metabolomics) (8). Global DNA sequencing efforts and several large-scale metagenome 84 sampling projects revealed the vast sequence diversity in environmental metagenomes and 85 microbial genomes, as well as the presence of numerous unknown or poorly characterised 86 genes (9-12). For example, a high-throughput project focused on carbohydrate-active 87 enzymes has identified over 27,000 related genes and demonstrated the presence of glycoside 88 hydrolase activity in 51 out of 90 tested proteins (13). Other large scale metagenomic projects 89 include the Sargasso Sea sampling (over one million new genes discovered), the Global 90 Ocean Survey (over six million genes), and human gut microbiome (over three million genes) 91 92 (9-12). Thus, through the advent of metagenomics, we are starting to generate insights into 93 the rich microbial worlds thriving in different environments. Nevertheless, a recent analysis 94 of metagenome screening studies suggested that all representative types of environmental habitats (terrestrial, marine, and freshwater) are under-sampled and under-investigated (14). 95 It is estimated that total number of microbial cells is 10^{30} , whereas the natural protein 96 universe exceeds 10¹² proteins indicating that our knowledge of proteins and biochemical 97 diversity on Earth is very limited (15-17). Therefore, the determination of protein function or 98 enzyme activity for millions of genes of unknown function and biochemically 99 100 uncharacterised proteins represents one of the main challenges of the postgenomic biology.

The approaches of experimental metagenomics include meta-transcriptomics, meta-101 proteomics, metabolomics, and enzyme screening (6,7,17-19). Activity-based screening of 102 metagenome gene libraries represents a direct way for tapping into the metagenomic resource 103 of novel enzymes. This approach involves expressing genes from metagenomic DNA 104 105 fragments in heterologous hosts, commonly Escherichia coli, and assaying libraries of clones 106 on agar plates for enzymatic activities using chromogenic or insoluble substrates (18). Importantly, this approach offers the possibility to identify novel families of enzymes with no 107 sequence similarity to known enzymes. Screening of metagenome gene libraries from 108 different terrestrial, marine, and freshwater environments has already expanded the number 109 110 of new enzymes including novel nitrilases, glycoside hydrolases, carboxyl esterases, and laccases (14,20,21). 111

112 Carboxylesterases (EC 3.1.1.1) are a diverse group of hydrolytic enzymes catalysing the 113 cleavage and formation of ester bonds, which represent the third largest group of industrial 114 biocatalysts (after amylases and proteases). Many esterases show a wide substrate range and 115 high regio- and stereo-selectivity making them attractive biocatalysts for applications in 116 pharmaceutical, cosmetic, detergent, food, textile, paper and biodiesel industries (22,23). 117 Most of known carboxylesterases belong to the large protein superfamilies of α/β hydrolases 118 and β -lactamases and have been classified into 46 subfamilies (including 11 true lipase subfamilies)based on sequence analysis (22,24,25). A significant number of these enzymes 119 120 have been characterised both biochemically and structurally, because they are of high interest 121 for biotechnological applications (22,23,26). Screening of metagenome gene libraries and genome mining has greatly expanded the number of novel carboxylesterases including 122 123 enzymes active against aryl esters or polymeric esters (polyesterases) (21-23,26,27). However, the increasing demand for environmentally friendly industrial processes has 124 125 stimulated research on the discovery of new enzymes and their application as biocatalysts to 126 meet the challenges of a circular bioeconomy (28,29). The global enzyme market is expected to grow from \$8.18 billion in 2015 to \$17.50 billion by 2024 (28). However, the majority of 127 known enzymes are originated from mesophilic organisms, which have limited stability under 128 129 harsh industrial conditions including high temperatures, extreme pH, solvents, and salts 130 (30,31). Thus, the discovery of robust enzymes including carboxylesterases and engineering 131 of more active variants represent the key challenges for the development of future 132 biocatalytic processes. Extremophilic microorganisms are an attractive source of industrial biocatalysts, because they evolved robust enzymes that function under extreme conditions 133 134 (high/low temperatures, high/low pH, salts) (14,26,30,32). In addition, extremophilic enzymes found in one environment are typically also tolerant to other extreme conditions 135 making them attractive biocatalysts for various applications including depolymerization of 136 137 synthetic polymers and inactivation of mycotoxins (32-36).

Hydrothermal vents are extreme environments located in tectonically active sites, which 138 are classified as terrestrial and marine (deep-sea and shallow-sea) systems (37). 139 140 Hydrothermal vents are characterised by harsh physico-chemical conditions (high 141 temperature and low pH) and are known as source of thermophilic microbes and enzymes 142 with biotechnological importance. Although terrestrial hydrothermal vents have relatively 143 easy access, they remain under-investigated compared to (sub)marine hydrothermal vents. To 144 provide insights into microbial diversity of terrestrial hydrothermal vents, we analysed the natural microbial communities of two thermophilic hydrothermal vents located on the 145 146 volcanic island of Ischia (Italy), as well as the effect of polyester plastic addition on these microbial communities using barcoded DNA sequencing of extracted DNA. Using activity-147 based metagenomic approach, we screened fosmid libraries for carboxylesterase activity 148 using tributyrin agar plates, identified 14 unique fosmids encoding putative hydrolases, from 149 which three soluble carboxylesterases (IS10, IS11, and IS12) were recombinantly produced 150 151 in E. coli and biochemically characterised including substrate range and stability using both 152 monoester and polyester substrates. The crystal structure of IS11 was resolved to reveal the 153 N-terminal β -lactamase-like serine hydrolase domain connected to the C-terminal lipocalin 154 domain. The active site of IS11 accommodated the conserved catalytic residues Ser68, 155 Lys71, Tyr160, and Asn162, as well as numerous hydrophobic residues potentially involved 156 in substrate binding. Structural models of IS10 and IS12 revealed classical α/β hydrolase domains with a catalytic serine hydrolase triad (Ser155, His280, Asp250), multiple 157 hydrophobic residues in their active sites with the largest "effective volumes" reported for 158 159 prokaryotic carboxylesterases.

160

161 MATERIALS AND METHODS

162 **Environmental sampling sites and enrichment cultures.** Sediment samples with water 163 were collected in September 2018 from the geothermal areas of the volcanic island of Ischia 164 (the Gulf of Naples, Italy). The samples were taken from the Cavascura hydrothermal springs (40.70403 13.90502): IS1 (pH 8.5, 45 °C) and IS2 (pH 7.0, 55 °C); and from the sandy 165 166 fumaroles of Maronti beach near St Angelo (40.70101 13.89837): IS3 (pH 4.5, 75 °C) and 167 IS4 (pH 5.0, 75 °C). For each sample, triplicate enrichment cultures were established 168 containing different polymers or plastics as substrates, polylactic acid film (PLA, poly-D,L-169 lactide, M_w 10,000-18,000 Da), polyhydroxybutyrate (PHB) and a commercial compostable polyester blend (P3, Blend), kindly provided by the Biocomposites Centre, Bangor 170 University, UK. Plastic films cut (3 mm x 20 mm), washed in 70 % ethanol and air-dried 171 were added to samples. For IS1 and IS2 cultures, modified DSMZ medium 1374 172 (https://bacmedia.dsmz.de/medium/1374) was used, which contained (g L⁻¹): NaCl, 1; 173 MgCl₂6H₂O, 0.4; KCl, 0.1; NH₄Cl, 0.25; KH₂PO₄, 0.2; Na₂SO₄, 4; NaHCO₃, 0.1; 174 CaCl₂2H₂O, 0.5. The medium was adjusted to pH 7.5 with 10N NaOH. For IS3 and IS4 175 cultures, modified DSMZ medium 88 (https://bacmedia.dsmz.de/medium/88) was used, 176 which contained (g L⁻¹): (NH₄)₂SO₄, 1.3; KH₂PO₄, 0.28; MgSO₄ 7H₂O, 0.25; CaCl₂ 2H₂O, 177 0.07. The medium was adjusted to pH 4.5 with 10N H₂SO₄. Additionally, the trace element 178 179 solution SL-10 (from DSMZ medium 320 https://bacmedia.dsmz.de/medium/320) was added 180 at 1:1000 (vol/vol) to both media. Enrichment cultures contained 0.5 g of sample sediment 181 and 0.25 g of a polymer in 10 mL of growth medium. The cultures were incubated at 50 °C (IS1-IS2) or 75 °C (IS3-IS4) with slow agitation (30 rpm) for 4 days, then culture aliquots 182 183 (20% of the volume of enrichment cultures) were transferred to a fresh medium and incubated for 11 days under the same conditions (Table S1). 184

DNA extraction and 16S rRNA amplicon sequencing. Prior to DNA extraction, the 185 enrichment cultures (9 mL each) were vortexed and biomass was collected by centrifugation 186 at 10,000 rcf for 10 min at 4 °C. The pellets were resuspended in 250 µL of sterile phosphate-187 188 buffered saline (PBS, pH 7.5) and transferred to 1.5 mL tubes. High molecular weight DNA 189 was obtained using the ZymoBIOMICS DNA Miniprep Kit (Zymo Research, Irvine, Ca, USA) in accordance with manufacturer's instructions. Finally, DNA was eluted with 50 µL 190 191 of nuclease free water. The quality of extracted DNA was assessed by gel electrophoresis, and DNA concentration was estimated using Qubit[™] 4.0 Fluorometer dsDNA BR Assay Kit 192 (Life Technologies, USA). The Illumina-compatible libraries of hypervariable V4 region of 193 194 16S rRNA gene were prepared by single PCR with dual-indexing primer system with heterogeneity spacer as described previously (38). Modified forward primer F515 (5'-195 GTGBCAGCMGCCGCGGTAA-3') and reverse R806 prokaryotic primer 196 (5'-GGACTACHVGGGTWTCTAAT-3') were used. PCR reactions were performed using 197 MyTaq[™] Red DNA Polymerase (Bioline) in a Bio-Rad[®] thermocycler with the following 198 program: 95 °C for 2 min for denaturation followed by 30 cycles at 95 °C for 45 s, 50 °C for 199 60 s, 72 °C for 30 s, with a final elongation at 72 °C for 3 min. PCR products of 200 201 approximately 440 bp were visualised by gel electrophoresis and gel-purified using the

QIAEX II Gel Extraction Kit[®] (QIAGEN). The purified barcoded amplicons were quantified
by Qubit[™] dsDNA BR Assay Kit (Life Technologies, USA), pooled in equimolar amounts
and sequenced on Illumina MiSeq[™] platform (Illumina Inc., San Diego, CA, USA) using
paired-end 250 bp reads at the Centre for Environmental Biotechnology (Bangor, UK).
Sequencing reads were processed and analysed as previously described (39). All statistical
analysis was conducted using R programming environment (40) *prcomp* function and inhouse scripts for graphical design.

209

Preparation of the Ischia metagenome library from polyester enrichment cultures. 210 High molecular weight DNA extracted from all enrichment cultures was combined in 211 equimolar amounts and used to prepare two metagenomic fosmid libraries 'IS Lib1' 212 213 (Cavascura enrichments) and 'IS Lib2' (Maronti enrichments) using the CopyControlTM 214 Fosmid Library pCC2FOS Production Kit (Epicentre Technologies, Madison, USA). DNA 215 was end-repaired to generate blunt-ended 5'-phosphorylated fragments according to 216 manufacturer's instructions. Subsequently, DNA fragments in the range of 30-40 kbp were resolved by gel electrophoresis (2 V cm⁻¹ overnight at 4 °C) and recovered from 1% low 217 melting point agarose gel using GELase 50X buffer and GELase enzyme (Epicentre). Nucleic 218 219 acid fragments were then ligated to the linearized CopyControl pCC2FOS vector following the manufacturer's instructions. After the in vitro packaging into the phage lambda 220 (MaxPlaxTM Lambda Packaging Extract, Epicentre), the transfected phage T1-resistant 221 EPI300TM-T1^R E. coli cells were spread on Luria-Bertani (LB) agar medium containing 12.5 222 µg mL⁻¹ chloramphenicol and incubated at 37 °C overnight to determine the titre of the phage 223 particles. The resulting library had estimated titre of 14×10^4 and 1×10^4 non-redundant fosmid 224 clones in IS Lib1 and IS Lib2 libraries, correspondingly. For long-term storage, E. coli 225 colonies were washed off from the agar surface using liquid LB medium containing 20 % 226 227 (v/v) sterile glycerol and the aliquots were stored at -80 °C.

Activity-based screening of the polyester enrichment metagenome library for esterase 228 activity. The metagenomic libraries were screened for carboxylesterase/lipase activity as 229 follows. The fosmid library was grown on LB agar plates containing 12.5 µg mL⁻¹ 230 chloramphenicol at 37 °C overnight to yield single colonies. Then, 3,456 clones were arrayed 231 in 9 x 384-well microtitre plates and cultivated at 37 °C in LB medium supplemented with 232 12.5 µg mL⁻¹ chloramphenicol. Those original microtitre plates were stored at -80 °C after 233 the addition of glycerol, at final concentration of 20 % (vol/vol). For screening clone 234 235 libraries, 384-pin replicators were used to print clones onto the surface of large LB agar square plates (245 mm x 245 mm) containing 12.5 µg mL⁻¹ chloramphenicol, 2 mL L⁻¹ 236 fosmid autoinduction solution (Epicentre), each plate contained 0.3 % (v/v) tributyrin 237 238 (Sigma-Aldrich, Gillingham, UK) as described earlier (27). After an initial overnight growth 239 at 37 °C, the LB agar plates were incubated for 48 hours at 37, 50 or 70 °C. Positive hits were 240 confirmed by re-testing of the corresponding fosmid clones taken from the original microtitre 241 plate.

242 Sequencing and analysis of metagenomic fragments. Positive fosmid clones were 243 cultivated in 100 mL LB medium containing 12.5 μ g mL⁻¹ chloramphenicol and 2 mL L⁻¹ 244 fosmid autoinduction solution (Epicentre) at 37 °C overnight. Biomass was collected by centrifuging at 3,200 g for 30 min and fosmid DNA was extracted from the pellet using the 245 QIAGEN Plasmid Midi Kit (QIAGEN) following the manufacturer's instructions. 246 247 Approximate size of the cloned fragments was assessed on agarose gel electrophoresis after 248 double endonuclease digestion with XbaI and XhoI (New England Biolabs, Ipswich, MA, USA). The Sanger sequencing of the termini of inserted metagenomic fragments of each 249 250 purified fosmid was done at Macrogen Ltd. (Amsterdam, The Netherlands) using standard pCC2FOS sequencing primers (Epicentre). Non-redundant fosmids were selected, their DNA 251 concentrations were quantified by Qubit[™] 4.0 Fluorometer dsDNA BR Assay Kit 252 253 (Invitrogen), pooled in equimolar amounts and prepared for Illumina MiSeq® sequencing. Pooled DNA was fragmented using the Bioruptor Pico Sonicator (Diagenode, Denville, NJ, 254 USA) with parameters adjusted to obtain 400-600 bp fragments. The fragment library was 255 256 prepared using the NebNext Ultra II DNA Library preparation kit (New England Biolabs, 257 Ipswich, MA, USA) according to the manufacturer's instructions. The obtained library was 258 sequenced on MiSeq® platform (Illumina, San Diego, USA) using a microflow cell 300-259 cycles V2 sequencing kit. Obtained paired end reads were subjected to quality filtering, trimming and assembly as previously described (41). Gene prediction and primary functional 260 the 261 annotation were performed using MetaGeneMark annotation software (http://opal.biology.gatech.edu) (42). Translated protein sequences were annotated using 262 263 BLAST searches of UniProt and the non-redundant GenBank databases (43). Multiple 264 sequence alignments were generated using MUSCLE application (44) and visualised on Geneious v.9 (Biomatters, New Zealand). The Neighbour-Joining and maximum likelihood 265 trees were constructed in MEGA X (45) using the settings for the Poisson model and 266 homogenous patterning between lineages. The bootstrapping was performed with 1,000 267 pseudoreplicates. 268

269 Gene cloning, expression and purification of selected proteins. Selected gene candidates 270 were amplified by PCR in a T100 Thermal Cycler (Bio-Rad) using Herculase II Fusion Enzyme (Agilent, Cheadle, UK) with oligonucleotide primer pairs incorporating pET-46 271 272 Ek/LIC vector adapters (Merck, Darmstadt, Germany). PCR products were then purified and 273 cloned into the above pET-46 Ek/LIC vector harbouring an N-terminal 6xHis tag, as 274 described by the manufacturer. The DNA inserts in the resulting plasmids were verified by 275 Sanger sequencing at Macrogen Ltd. (Amsterdam, The Netherlands) and then transformed 276 into E. coli BL21(DE3) for recombinant protein expression. E. coli BL21(DE3) harbouring pET-46 Ek/LIC plasmid were grown on LB medium to mid-log growth phase (OD₆₀₀ 0.7-277 0.8), induced with isopropyl- β -d-thiogalactopyranoside (IPTG, 0.5 mM) and incubated at 20 278 279 °C overnight. Cells were disrupted by sonication as reported earlier (46) and recombinant 280 proteins were purified using metal-chelate affinity chromatography on Ni-NTA His-bind 281 columns. Protein size and purity were assessed using denaturing gel electrophoresis (SDS-282 PAGE), and protein concentration was measured by Bradford assay (Merck, Gillingham, 283 UK).

Enzyme assays. Carboxyl esterase activity of purified proteins against *p*-nitrophenyl (*p*NP) or α -naphthyl (α N) esters was determined by measuring the amount of α -naphthol released by esterase-catalysed hydrolysis essentially as described previously (27,46). Under standard 287 assay conditions the reaction mixture contained 50 mM potassium phosphate buffer (pH 7.0), 1 mM pNP-butyrate as substrate, and 0.2-1.8 μ g of enzyme in a final volume of 200 μ L. 288 Reactions were incubated at 30 °C for 3-5 min and monitored at 410 nm (for pNP esters) or 289 290 310 nm (for αN esters). Non enzymatic hydrolysis of ester substrates was subtracted using a 291 blank reaction with denatured enzyme. The effect of pH on esterase activity was evaluated 292 using the following buffers: sodium citrate (pH 4.0 and 5.0), potassium phosphate (pH 6.0 293 and 7.0), Tris-HCl (pH 8.0 and 9.0). The activity was monitored at 348 nm (the pH-294 independent isosbestic wavelength of α -naphthol). The effect of temperature on esterase 295 activity was studied using a range of temperatures (from 20 °C to 90 °C). In order to assess 296 the thermal stability of purified esterases, the enzymes were dissolved in potassium 297 phosphate buffer (pH 7.0) and preincubated at the indicated temperatures (from 30 to 95 °C) for 20 min. The enzyme solutions were then cooled down on ice and the residual activity was 298 299 measured under standard conditions (at 30 °C). Substrate specificity of purified enzymes was 300 analysed using model pNP- and α N-esters with different chain lengths: pNP-acetate (C2), 301 α N-propionate (C3), pNP-butyrate (C4), α N-butyrate (C4), pNP-hexanoate (C6), pNP-302 dodecanoate (C12), and pNP-palmitate (C16), obtained from Sigma-Aldrich and Tokyo 303 Chemical Industry TCI. Kinetic parameters for these substrates were determined over a range 304 of substrate concentrations (0.012-4 mM; 30 °C) and calculated by non-linear regression 305 analysis of raw data fit to Michaelis-Menten function using GraphPad Prism software v.6. 306 Hydrolysis of 44 soluble non-chromogenic monoester substrates (Table S2) and T-2 307 mycotoxin (Merck Life Science S.L.U., Madrid, Spain) was assayed at 37 °C using a pH 308 indicator assay with Phenol Red and monitored at 550 nm (47). The reaction products of 309 enzymatic degradation of T-2 mycotoxin were analysed using reversed phase chromatography on a Waters 600 HPLC system, coupled to an autosampler (Waters, model 310 311 717plus), and equipped with a Zorbax Eclipse Plus C18 column (Agilent, 4.6 x 100 mm, 3.5 μm, 40 °C) and an Evaporative Light Scattering Detector (ELSD, Sedere Sedex model 55). 312 313 The reaction products were separated using gradient elution (1.0 ml/min) with acetonitrile and water (5%: 1 min, 5%-95%: 9 min, 95 %: 3 min, 5 %: 7 min). Polyester 314 315 depolymerization activity of purified proteins against 3PET (bis(benzoyloxyethyl) terephthalate) was measured using 1.5 % agarose plates containing 0.2 % of emulsified 316 polyesters. 3PET was purchased from CanSyn Chem. Corp. (Toronto, Canada). Agarose 317 plates with emulsified 3PET were prepared as described previously (48). After protein 318 loading, the plates were sealed and incubated at 37 °C for 1-5 days. The presence of 319 polyesterase activity was indicated by the formation of a clear zone around the wells with 320 proteins. Apart from plate assays, activity assays of IS10, IS11 and IS12 for 3PET suspension 321 hydrolysis were performed in 50 mM Tris-HCl buffer, pH 8.0, at 30 °C, in a shaker at 600 322 323 rpm, the final reaction volume for each experiment was 0.2 mL, and the final protein amount 324 50 μ g. The reactions were terminated after 13 h by filtering reaction mixture on a 10 kDa 325 spin filter. 10 µL of filtrate was analysed using the high-performance liquid chromatography 326 system (HPLC), Schimadzu, Prominence-I (Milton Keynes, UK) equipped with a Schimadzu C18 Shim-pack column (4.6 \times 150 mm, 5 μm). The mobile phase was 25 % (vol/vol) 327 methanol with 0.1 % (vol/vol) H₃PO₄ in HPLC-grade water at a flow rate of 0.7 mL min⁻¹ for 328 329 2 min, following increase to 55 % of methanol to 118 min, followed by 25 % methanol at 22 330 min; the effluent was monitored at the wavelength of 240 nm, the column was conditioned at 40 °C. The hydrolytic products of mono(2-hydroxyethyl)terephthalic acid (MHET), bis(2-hydroxyethyl)terephthalate (BHET) and terephthalic acid (TPA) were identified by
comparing the retention times with their standards, and reactions without enzyme were
served as negative controls. All samples of each experiment were analysed in triplicate.

335 **Enzymatic activity against PLA.** Hydrolysis of PLA was assayed by measurement of lactic 336 acid production as follows: 5 mg of each PLA (all, acid-terminated and purchased from 337 PolySciTech (W. Lafayette, USA)), P(D)LA 10-15,000 Da, P(D,L,)LA (Resomer R202H, 10-18,000 Da) or P(L)LA 15-25,000 Da) suspended in 0.5 mL of 0.4 M Tris-HCl (pH 8.0) were 338 mixed with 50 µg of purified enzyme and incubated for 48 h at 37 °C with shaking (1000 339 rpm). Samples were then centrifuged at 12,000 g for 5 min at 4 °C. 200 µl of supernatant 340 were mixed with 200 μ l of mobile phase (0.005 N H₂SO₄). Sample was filtered through 13 341 342 mm Millipore PES syringe membrane filter (0.02 µm pore diameter) and analysed by HPLC Shimadzu, Prominence-I (Milton Keynes, UK) with an ion exchange column Hi PlexH (300 343 x 7.7 mm) (Agilent, Cheadle, UK) and 0.6 mL min⁻¹ flow rate at 55 °C (oven temperature) 344 with UV detector set at 190-210 nm. 345

346 Protein crystallization and structure determination. Native metagenomic esterases were 347 purified using metal-chelate affinity chromatography, and crystallization was performed at room temperature using the sitting-drop vapor diffusion method. For IS11, protein 348 concentration was 25 mg mL⁻¹, reservoir solution 0.1 M citric acid, pH 3.5 and 19 % PEG 349 350 3350). The crystal was cryoprotected by transferring into paratone oil and flash frozen in liquid nitrogen. Diffraction data for the IS11 crystal was collected at 100 K at a Rigaku home 351 352 source Micromax-007 with R-AXIS IV++ detector and processed using HKL3000 (49). The 353 structure was solved by molecular replacement using Phenix.phaser (50) and a model built by 354 AlphaFold2 (51). Model building and refinement were performed using Phenix.refine and 355 Coot (52). TLS parameterization was utilized for refinement, and B-factors were refined as 356 isotropic. Structure geometry and validation were performed using the Phenix Molprobity 357 tools. Data collection and refinement statistics for this structure are summarized in Table S3.

Accession numbers. SSU rRNA gene sequences were deposited to GenBank as BioProject ID: PRJNA881593. Sequences of IS10-IS12 proteins were deposited to GenBank under accession numbers OL304252, OL304253, and OL304254. The atomic coordinates of IS11 have been deposited in the Protein Data Bank (PDB), with accession code 7SPN.

362

363 **RESULTS and DISCUSSION**

364 Natural microbial communities of terrestrial hydrothermal vents of Ischia and effect of 365 polyester enrichments. To provide insights into the composition of natural microbial communities and thermophilic enzymes of hydrothermal vents of the island of Ischia, four 366 367 sediment samples were collected from the Cavascura hot spring (samples IS1 and IS2) and from Maronti beach near Sant'Angelo (samples IS3 and IS4) (see Materials and Methods). 368 Both sites represent thermophilic habitats with slightly different environmental conditions: 369 IS1 (pH 7.0, 45 °C), IS2 (pH 8.5, 55 °C), IS3 (pH 4.5, 75 °C), and IS4 (pH 5.0, 85 °C) (Table 370 S1). From each sample, total DNA was extracted and subjected to barcoded amplicon 371

372 sequencing of the V4 region of 16S rRNA gene. Sequence analysis revealed that the IS1 373 community comprised mainly *Pseudomonas* (17.2 %), class Anaerolineae (Chloroflexi) (12.3%), class Armatimonadota (10.0%), *Elizabethkingia* (phylum Bacteroidota) (9.5%), 374 other Myxococcota (9.1 %), Sphingobacterium (order Sphingobacteriales, class Bacteroidia, 375 376 phylum Bacteroidota) (6.7 %), and class Nitrospirota (6.4%), whereas the IS2 community was dominated by *Caldimonas* (order Burkholderiales, class Gammaproteobacteria) (63.9 %), 377 378 Cutibacterium (order Propionibacteriales, class Actinobacteria) (17.2%), and Thermus 379 (phylum Deinococcota) (16 %) (Fig. 1). In contrast, the IS3 community was mainly represented by Bacillales (Firmicutes), namely Brevibacillus (48.3%) and Geobacillus (42 380 381 %), and other Bacilli (4.4 %), whereas IS4 comprised Sphingobacterium (Sphingobacteriales, 382 Bacteroidetes) (31.9 %), Thermobaculum (Thermobaculales, Chloroflexi) (17.4 %) and Geobacillus (10.7 %), followed by Pseudomonas (7%) and Bacillus (6.1%) (Fig. 1). The 383 observed differences in the taxonomic composition of the Cavascura (IS1 and IS2) and 384 Maronti (IS3 and IS4) samples can be attributed to different environmental conditions 385 386 (temperature and pH) at the sampling sites.

387 Using the four sediment samples from two Ischia sites, twelve enrichment cultures were 388 established with different polyester plastics as carbon substrate including PHB, PLA and 389 commercial polyester blend (Table S1). After two weeks of incubation with polyesters, the 390 IS1 enrichment culture showed a drastic increase in the relative abundance of members of the 391 order Burkholderiales within the families Comamonadaceae and Rhodocyclaceae (relative abundance 15.2-35.1 % across the three plastic enrichments), Fontimonas (Solimonadaceae, 392 16.9-27.5%), and Schleiferia (order Flavobacteriales, 15.6-34.4%) (Fig. 1). Likewise, IS2 393 enrichment showed an increase in Fontimonas (11-26%), Schleiferia (21% in PHB 394 395 enrichment), whereas the relative content of the Caldimonas decreased from 63 % to 5.7 %, in favour of members of other families of the order Burkholderiales, namely Rhodocyclaceae, 396 Hydrogenophilaceae and Comamonadaceae (18-43%) Kapabacteriales (phylum Bacteroidota, 397 398 2.9-8%) and Rehaibacterium (order Xanthomonadales, 0.3-9.3%) (Fig. 1). The enrichment 399 culture with the compostable P3 blend stimulated the growth of Rhodocyclales, as both IS1 400 and IS2 showed a strong increase in *Thauera* compared to experiments with PHB and PLA 401 (Fig. 1). In the enrichment cultures IS3 and IS4, higher incubation temperature $(75^{\circ}C)$ 402 selected for thermophilic bacteria, and the nature of polyester used for enrichments 403 influenced the microbial composition (Fig. 1). The PHB enrichment stimulated growth of 404 Thermus (Deinococcota), which accounted for 66.7 % (92-fold increase) and 90.9% (1,280-405 fold increase) of the total reads in IS3 and IS4, respectively, followed by Geobacillus and 406 other members of Firmicutes. In contrast, the PLA culture favoured growth of Geobacillus, 407 which reached a relative abundance of 95.8% in IS3 (2.3-fold increase) and 91.8% in IS4 408 (8.6-fold increase), followed by Thermus and Brevibacillus. Finally, the commercial polyester blend promoted growth of both Geobacillus (accounted for 68 % or 1.6-fold 409 410 increase) and *Thermus* (accounted for 31.5% or 43.8-fold increase) in the IS3 enrichment, 411 whereas the IS4 culture was dominated by Firmicutes, Geobacillus (81 %), Paenibacillus (11.9%), Brevibacillus (5.9%), and Thermus (1.18%). As expected, the Shannon index of 412 413 microbial diversity (a measure of diversity of species in a community) (Fig. S1) revealed an 414 overall tendency to decrease after incubation with polyester plastics, with the exception of 415 IS2, which also showed low diversity in the native sample with the flattened rarefaction curve 416 (Fig. S1).

417

418 Activity-based screening of the hydrothermal metagenome library from Ischia for 419 **carboxylesterase activity.** After two weeks of incubation with polyesters, total DNA was 420 extracted from the enrichment cultures and combined for the construction of the 421 metagenomic fosmid libraries IS Lib1 and IS Lib2. In order to identify carboxylesterases 422 with high-temperature profiles, this library was screened for esterase activity with tributyrin 423 as substrate (for carboxylesterases and lipases) at three temperatures: 37, 50 and 70 °C. 424 Emulsified tributyrin gives a turbid appearance to the plates, and the presence of active 425 metagenomic esterases or lipases is seen as a clear zone around the colony. After screening 3,456 clones from the IS Libr2 library on tributyrin agar plates, 64 positive hits were 426 427 identified with 19 positive clones observed at 37 °C, 27 clones at 50 °C, and 18 clones at 70 428 °C. Furthermore, eight esterase positive clones detected at 50 °C were found to be unique for 429 this temperature, whereas one unique clone was found at 70 °C suggesting that these 430 esterases are mostly active only at elevated temperatures. Following endonuclease digestion 431 profiling and Sanger sequencing analysis, 14 non-redundant fosmids were selected for insert 432 sequencing using the Illumina platform, and fosmid inserts were assembled with an average 433 size of 39 kbp. Sequence analysis revealed 12 putative ORFs encoding predicted hydrolases 434 (including peptidases, carboxylesterases, β -lactamases, serine proteases) homologous to 435 proteins from Chloroflexi and metagenome assembled genome (MAG) affiliated to 436 thermophilic Chloroflexi. From candidate proteins cloned in E. coli, three putative carboxylesterases (IS10, IS11, and IS12) were soluble, when expressed in E. coli cells, and 437 438 the presence of carboxylesterase activity in purified proteins was confirmed using tributyrin agarose plates assay (Table 1) and were further selected for detailed biochemical 439 440 characterisation. Amino acid sequences of IS10 (314 amino acids), IS11 (455 aa), and IS12 (318 aa) showed no presence of recognizable signal peptides. Both IS10 and IS12 belonged 441 to the α/β hydrolase superfamily and had 56.8% sequence identity one to another, whereas 442 443 IS11 showed no significant sequence similarity to IS10 and IS12 as it was a member of the 444 large family of β -lactamases and penicillin-binding proteins (Table 1). A blastP search of the 445 nrNCBI database revealed that amino acid sequences of IS10 and IS12 were identical to two 446 putative α/β hydrolases from uncultured Chloroflexi bacteria (GenBank accession numbers 447 HEG24678.1 and HHR50377.1, respectively), whereas the IS11 sequence exhibited the highest identity (99.1%) to the putative "class A β -lactamase-related serine hydrolase" 448 449 HDX58025.1 from uncultured Dehalococcoidia. Interestingly, the top homologous proteins 450 of Ischia esterases were the proteins identified in metagenome from a deep-sea hydrothermal 451 vent (black smoker) in the Mid-Atlantic Ridge (South Atlantic Ocean) (53). The comparison 452 with previously characterised proteins showed the thermostable arylesterase, Are, from 453 Saccharolobus solfataricus (UniProt ID B5BLW5, 306 aa) being the top homologue for IS10 (42 % sequence identity), whereas the metagenome-derived esterase Est8 (KP699699, PDB 454 455 4YPV, 348 aa) was the top characterised homologue for IS12 (56 % sequence identity) 456 (54,55) (Fig. S2). The IS11 sequence was homologous to penicillin-binding proteins and β -457 lactamases with low sequence similarity to the CmcPBP from Actinobacteria Amycolatopsis 458 lactamdurans (Q06317, 36 % identity) and esterase EstB from Burkholderia gladioli 459 (Q9KX40, 32 % identity) (56,57). Domain and multiple sequence alignment confirmed the presence of conserved regions and motifs linked to esterase activity in lipolytic families 460 previously described (Fig. S2 and S3). IS10 and IS12 contained an α/β hydrolase fold 461 (PF07859), displaying the characteristic catalytic triad composed of Ser¹⁵⁵, Asp²⁵⁰ and His²⁸¹ 462

and the conserved consensus motif G-x-S-x-G around the active site serine (22), clustering
together with representatives of family IV (Fig. S2 and S3).

The protein IS11 contained a β -lactamase domain (PF00144) and the consensus tetrapeptide S-x-x-K, perfectly conserved among all penicillin-binding enzymes and β lactamases, surrounding the active serine Ser68. In addition, Lys71 and Tyr160 were also conserved as part of the catalytic triad of family VIII esterases, which groups enzymes with homology to class C β -lactamases and penicillin-binding proteins (Fig. S2 and S3).

470 Biochemical characterisation of purified metagenomic carboxylesterases using model 471 esterase substrates. The esterase activity of purified proteins (IS10, IS11, IS12) was 472 initially evaluated using model esterase substrates with different chain lengths (C2-C16) at 30 473 °C (to diminish spontaneous substrate degradation at high temperatures). The proteins were 474 found to be active against several short acyl chain substrates with IS10 and IS11 showing a 475 preference to pNP-butyrate, α N-butyrate, and pNP-hexanoate, whereas IS12 was most active 476 with pNP-acetate and α N-propionate (Fig. 2). All enzymes were active within a broad pH range (pH 6.0-9.0) with maximal activities at pH 9 (Fig. S4a). The purified metagenomic 477 478 carboxylesterases exhibited saturation kinetics with model esterase substrates at optimal pH 479 9.0 and 30 °C (Table 2). IS10 appeared to be the most efficient esterase compared to IS11 and 480 IS12, with the highest substrate affinity (lowest $K_{\rm M}$) and catalytic efficiency ($k_{\rm cat}/K_{\rm M}$) towards 481 the tested model substrates. IS12 showed higher substrate affinity to pNP-butyrate and higher 482 activity with pNP-acetate than IS11, whereas the latter was more active against pNP-butyrate (Table 2). 483

484 Since the selected carboxylesterases originated from thermophilic environments, we investigated the effect of temperature on the activity (temperature profiles) and 485 thermostability of purified carboxylesterases using p-NP-butyrate as substrate (Fig. 3). All 486 enzymes showed considerable activity at 20 °C, but reaction rates increased 5-10 times at 487 488 higher temperatures with IS10 showing the highest activity at 60-70 °C, whereas IS12 was 489 most active at 70°C-80°C and IS11 at 80-90 °C (Fig. 3). The thermostability of purified 490 enzymes was analysed using 20 min preincubation at different temperatures (from 30 to 95°C) followed by esterase assays with pNP-butyrate at 30 °C. IS10 retained 60% activity 491 492 after preincubation at 50 °C and showed a complete loss of activity at 80 °C (Fig. 3). In 493 contrast, both IS11 and IS12 revealed a significant decrease of activity only after 20 min preincubation at 90 °C and 70 °C, respectively. After two hours of incubation at 70 °C, IS12 494 retained 50% of initial activity, but was completely inactivated at 80 °C (Fig. 4). However, 495 496 IS11 showed no loss or a small reduction of activity at 70 °C and 80 °C, respectively, and 497 required over three hours of incubation at 90°C for inactivation (Fig. 4). Thus, the 498 metagenomic carboxylesterases from the Ischia hydrothermal vents are the thermophilic enzymes highly active at 70-80 °C with IS11 and IS12 also showing significant 499 500 thermostability at temperatures from 60 to 80 °C. Furthermore, the thermostability of IS11 501 and IS12 was comparable with, or exceeded the, thermostability of other esterases identified 502 in high-temperature environments (41,58-61).

503 Esterase activity of purified metagenomic esterases was inhibited by high concentrations 504 of NaCl (50-67% of remaining activity in the presence of 0.5 M NaCl) with IS11 showing a 505 slightly higher resistance (Fig. S4). Similarly, IS11 retained higher activity in the presence of 506 non-ionic detergents (43% and 53% in the presence of 2% Triton X-100 and Tween 20) (Fig. 507 S4). With organic solvents, IS10 was inhibited by acetone, acetonitrile, ethanol, and 508 isopropanol (10 %, v/v) (Fig. S5). In contrast, IS11 was more tolerant to these solvents (10-509 50 %) and was stimulated by 10% ethanol (60% increase) and 30% methanol (84% increase). 510 Furthermore, low concentrations of these solvents (10%, v/v) stimulated esterase activity of 511 IS12 (26-34 % increase), whereas higher concentrations of acetone (50 % v/v) and 512 isopropanol (30% v/v) were inhibiting. Finally, DMSO (10-30 %) stimulated esterase activity 513 of enzymes (20-46 % increase) but was inhibitory to IS10 at 30% (Fig. S5).

514 Substrate range of purified carboxylesterases. To analyse the substrate range and preference of metagenomic carboxylesterases from the Ischia hydrothermal vents, the 515 516 purified proteins were examined for the presence of hydrolytic activity against chemically 517 and structurally diverse esters, including alkyl and aryl esters (Materials and Methods). Both 518 IS10 and IS12 revealed a broad substrate range with significant activity against all 44 esters 519 tested ester substrates and the highest activity with phenyl acetate, phenyl propionate, glyceryl tripropionate, tributyrin, and α N-acetate (Table S2). IS12 was also highly active 520 toward vinyl propionate. The broad substrate range of IS10 and IS12 correlates with 521 relatively large effective volumes of their active sites, 650.23 Å³ and 780.5 Å³, respectively 522 (calculated as cavity volume/solvent accessible surface area) (23). These volumes are the 523 524 largest calculated for prokaryotic esterases experimentally characterised so far, with only 525 CalA lipase (Novozym 735) from the yeast C. antarctica having a larger value (23). IS11 had a more restricted substrate range, showing detectable activity against 22 ester substrates 526 of 44 tested with a preference for benzyl (R)-(+)-2-hydroxy-3-phenylpropionate (Table S2). 527 In this study we found this ester being hydrolysed by the three esterases (IS10, IS11, IS12), 528 529 but preferentially by IS11, suggesting that either the lipocalin domain of IS11 or the hydrophobic and polar residues located at the active site of this esterase (see structural 530 531 features below) may have a role for the preference of this ester, not only as compared to the 532 other two esterases but also as compared to other esters. The three metagenomic esterases 533 revealed no apparent enantio-preference and hydrolysed both enantiomers of several tested 534 commercially available chiral substrates.

535

The purified metagenomic esterases were also tested for hydrolytic activity against the T-536 537 2 mycotoxin, which contains three ester groups on its side chains. The T-2 and deacetylated HT-2 toxins are members of the large group of trichothecene mycotoxins (over 190 538 539 derivatives) containing a tetracyclic ring system (62). Mycotoxins are highly toxic fungal 540 metabolites frequently contaminating food and feed and causing negative effects on human 541 health, animals, and economy (63,64). While physical and chemical methods have been used to detoxify mycotoxins, biological detoxification using enzymes or microbes is more 542 attractive due to specificity, safety, and costs. With T-2 as substrate, both IS10 and IS12 543 showed high esterase activity based on a pH-shift assay with phenol red (2.3 U/mg and 4.4 544 545 U/mg, respectively, at 37°C and pH 8.0), whereas IS11 was found to be inactive. Hydrolytic activity of IS10 and IS12 against T-2 was confirmed using HPLC, which also revealed the 546 547 formation of different reaction products (Fig. 5). IS10 produced HT-2 as the main product, 548 whereas HT-2 was present as the minor product in the reaction mixture with IS12, which 549 produced mostly the T-2 triol as the main product (Fig. 5). Since the T-2 triol is known to be 550 less toxic than T-2 and HT-2 (36,65), IS12 might represent a promising candidate for the 551 biodetoxification of T-2 and HT-2.

552

553 Since our metagenomic libraries were prepared using enrichment cultures with synthetic 554 polyesters, the purified esterases were also tested for the presence of polyesterase activity. 555 Although recent studies on biocatalytic depolymerization of synthetic polyesters including PLA and polyethylene terephthalate (PET) have shown the potential of microbial 556 carboxylesterases, there is an urgent need to identify novel robust polyesterases for 557 applications in plastics recycling (35,48,66). The purified esterases were screened for the 558 559 presence of polyesterase activity using an agarose plate assay with the emulsified PET model substrate, 3PET. These screens revealed the presence of polyesterase activity against 3PET in 560 both IS10 and IS12, as indicated by the formation of a clear zone around the wells with 561 562 loaded enzymes after incubation at 37 °C (Fig. 6A). Purified IS11 did not show a visible clearance zone on the 3PET plate, however the *in vitro* assay of hydrolysis of 3PET by IS11 563 564 and HPLC analysis of reaction products, showed an increase in MHET, which was the main 565 hydrolysis product while IS10 and IS12 produced BHET as the principal hydrolysis product (Fig. 6B). Furthermore, enzymes exhibited activity toward PLA, with a clear substrate 566 preference toward P(DL)A, over P(L)LA, over P(D)LA (Fig. 6C). To sum up, both IS10 and 567 IS12 exhibited broad substrate profiles and were able to degrade both mycotoxins and 568 569 polyesters.

570 Structural studies of metagenomic carboxylesterases. To provide structural insights into the active site and activity of metagenomic carboxylesterases, purified proteins (IS10, IS11, 571 572 and IS12) were subjected to crystallization trials. IS11 produced diffracting crystals, and its 573 crystal structure was determined by molecular replacement (Table S3, Materials and 574 Methods). The overall structure of IS11 revealed a protein dimer with protomers composed of 575 two structural domains, an N-terminal β -lactamase-like serine hydrolase domain (1-345 aa) 576 connected via a flexible linker (346-358 aa) to a C-terminal lipocalin domain (Fig. 7 and Fig. S6). Protein oligomerization has been suggested to contribute to thermostability of several 577 578 thermophilic carboxylesterases (e.g. AFEst, PestE, EstE1) (33,61,67). Accordingly, the results of size-exclusion chromatography of purified IS11, as well as IS10 and IS12, suggest 579 that these proteins exist as dimers in solution (Fig. S7). 580

581 The serine β -lactamases (classes A, C, and D) are structurally and evolutionary related to penicillin-binding proteins (the targets of β -lactam antibiotics), which also include hydrolytic 582 583 DD-peptidases (68,69). The overall structure of the IS11 β -lactamase domain is composed of 584 a mostly α -helical (all- α) sub-domain inserted into an $\alpha/\beta/\alpha$ sandwich (or an α/β sub-domain) 585 (Fig. 7 and 8 and Fig. S6). The $\alpha/\beta/\alpha$ sandwich sub-domain includes a nine-stranded antiparallel β -sheet flanked by two helices on each side, whereas the mostly helical sub-586 587 domain comprises nine α -helices (Fig. 8). The search for structural homologues of IS11 using 588 the Dali server (70) identified numerous β -lactamase-like proteins with low sequence identity including the Pyrococcus abyssi peptidase PAB87 (PDB code 2QMI) and Pseudomonas 589 *fluorescens* β -lactamase AmpC (PDB code 2QZ6) as the top structural homologues (Z-score 590 36.0-40.2, r.m.s.d. 2.1-2.8 Å, sequence identity 22-29%). The two sub-domains form a 591 groove accommodating the catalytic residues including Ser68 (a nucleophile) and Lys71 (a 592 general base accepting the proton from Ser68 O^{γ}) (1st motif S-x-x-K), Tyr160 and Asn162 593 (2nd motif Y-x-N/S), and His299 (3rd motif H/R/K-T/S/G-G). Accordingly, the IS11 structure 594 revealed the presence of an additional electron density positioned near the side chains of 595 596 Ser68, Tyr160, and His299, represents an unknown ligand covalently attached to Ser68 597 (could not be modeled with various components of the protein purification or crystallization solutions) (Fig. 9a). The positioning of these catalytic residues was also conserved in the 598 599 active sites of the biochemically characterised carboxylesterases with a β -lactamase fold

(family VIII): EstB from *Burkholderia gladioli* and Pab87 from *Pyrococcus abyssi* (71,72),
suggesting a common catalytic mechanism with acylation-deacylation. The catalytic cleft of
IS11 also contains several hydrophobic and polar residues potentially involved in substrate
binding (Asp126, Phe128, Trp158, Asn304, Ile307, Leu309) (Fig. 9).

604 The C-terminal domain of IS11 represents a typical lipocalin fold with one α -helix and an 605 eight-stranded antiparallel β -barrel containing a hydrophobic core (Fig. 10). Lipocalins are a diverse family of small individual proteins or domains (160-180 aa), which bind various 606 607 hydrophobic molecules (e.g. fatty acids) in a binding pocket located inside the barrel (73). Although lipocalins are very divergent in their sequences and functions, their structures 608 609 exhibit remarkable similarity. The lipocalin α -helix of IS11 closes off the top of the β -barrel, 610 whose interior represents a ligand-binding site coated mostly with hydrophobic residues (Fig. 611 10). In the IS11 protomer, the lipocalin domain covers the β -lactamase domain shielding the 612 catalytic cleft with the extended proline-rich strand (Pro391-Ser409) containing eight Pro 613 residues (Pro391, Pro396, Pro401, Pro402, Pro404, Pro406, Pro407, and Pro408) (Fig. 10). In 614 the thermophilic carboxylesterase Est2 from *Alicyclobacillus acidocaldarius*, the increased 615 number of Pro residues has been suggested to be important for thermostability, because they 616 reduce the flexibility of loops and other structural elements making them more resistant to 617 denaturation (74). The side chains of several residues of the lipocalin domain and proline-rich 618 strand are positioned close to the IS11 active site suggesting that they can be involved in 619 substrate binding (Phe395, Arg397, Lys398, Arg403, Arg449). Typically for all lipocalins, 620 the interior of the IS11 β-barrel is coated by mostly hydrophobic and polar residues (Leu373, Ser374, Ile376, Leu387, Gln389, Leu426, Ser429, Phe444, Phe446, Phe451). Proline-rich 621 sequences are also known to be directly involved or facilitating protein-protein interactions or 622 623 oligomerization (75). However, the IS11 dimer structure revealed no obvious interactions 624 between the individual lipocalin domains (Fig. S6) suggesting that the lipocalin domain of IS11 participates in substrate binding, rather than in the oligomerization. 625

626 High-quality structural models of IS10 and IS12 proteins constructed using the Phyre2 server (Fig. S8) revealed the presence of a core domain with a classical α/β hydrolase fold 627 628 and an all-helical domain, as well as a serine hydrolase catalytic triad (Ser155, His280, and 629 Asp250 in both proteins) (Fig. S9). The putative catalytic nucleophile Ser155 is located on 630 the classical nucleophilic elbow, a short sharp turn between a β -strand and α -helix. It is located at the bottom of the active site, which is mostly covered by the all-helical lid domain 631 632 (Fig. S9). Both acyl- and alcohol-binding pockets of IS10 and IS12 include several 633 hydrophobic and polar residues potentially involved in substrate binding (IS10: His81, Trp85, 634 His93, Asn159, Tyr183, Val185, Leu252; IS12: Trp85, Ile87, His93, Asn159, Tyr183, 635 Leu252, Ile279, Val283, Thr284, Leu285) (Fig. S9). Furthermore, the lid domains of both enzymes contain additional hydrophobic residues, which can contribute to substrate binding 636 (IS10: Phe34, Met38, Phe203, Leu204, Met208, Met209, Tyr211; IS12: Phe22, Met34, 637 638 Tyr195, Leu203, Leu204, Met209, Phe212, Trp213).

639

640 CONCLUSION

Present work has demonstrated a high value of high-temperature microbial habitats,
particularly of the volcanic island of Ischia (Italy), Terme di Cavascura and Maronti Beach
hydrotherms populated by taxonomically diverse microorganisms, as a resource for discovery

644 of high-temperature active enzymes. As revealed by an in-depth characterization of three metagenomics-derived carboxylesterases (IS10, IS11 and IS12) they were active at 645 temperatures as high as 70-90 °C and were capable to degradation of bio-based and synthetic 646 polyester plastics. The 3PET was hydrolysed by IS10 and IS12 to predominantly BHET, 647 648 while IS11 produced MHET as a main product. Interestingly, IS12 further degraded 649 mycotoxin T-2, a common agent causing poisoning the animal feed, to the less toxic T-2 triol. 650 The three wild-type enzymes may readily be applicable in pilot trials in industrial processes 651 relevant to the circular bioeconomy for plastics and/or in the production of toxin free foods 652 and feeds. This study can also serve as a starting point for deepening our knowledge on 653 structural determinants for substrate specificity in carboxylesterases and for rational 654 engineering to further improve their catalytic efficiencies to make them accepting PET 655 oligomers larger than 3PET.

656

657 ACKNOWLEDGMENTS

658 This study was conducted under the auspices of the FuturEnzyme Project funded by the 659 European Union's Horizon 2020 Research and Innovation Programme under Grant Agreement No. 101000327. M.F. and F.J.P. also acknowledge grants PID2020-112758RB-660 661 IO0 (M.F.), PDC2021-121534-IO0 (M.F.), TED2021-130544B-IO0 (M.F.) and PID2019-662 105838RB-C31 (F.J.P.) from the MCIN/AEI/10.13039/501100011033 and the European Union ("NextGenerationEU/PRTR"). M.A.D., T.T.C., R.B., A.N.K., O.V.G., A.F.Y. and 663 664 P.N.G. thank the support from the European Regional Development Fund (ERDF) through 665 the Welsh Government to the Centre for Environmental Biotechnology (CEB), Project Nr 666 81280. P.N.G. and A.F.Y. acknowledge the Natural Environment Research Council UK 667 (NERC)-funded Plastic Vectors project NE/S004548/1 and the Sêr Cymru programme partly 668 funded by the ERDF through the Welsh Government for the support of the project 669 BioPOL4Life. We are indebted to Connie Tulloch and Gwion Williams for their technical 670 support.

671

FIGURE LEGENDS

Fig. 1. The composition of microbial communities of native samples from the Ischia hydrothermal vents (IS1 (green), IS2 (orange), both from Cavascura; IS3 (purple) and IS4 (magenta), both from Maronti Beach) and their enrichment cultures set up with PHB, PLA and polyester blend and incubated for 4 days at 50 °C (IS1-IS2) or 75 °C (IS3-IS4), with consequent transfer into the fresh medium and incubation at same temperatures for 11 days. The relative abundance of barcoded V4-region 16S rRNA gene amplicon reads derived from particular taxa, is reflected in the sizes of circles. For reference, s. the panel in the top left corner.

Fig. 2. Hydrolytic activity of purified IS10 (A), IS11 (B) and IS12 (C) against model esterase substrates. The reaction mixtures contained the indicated *p*-nitrophenyl esters (*p*NP, white bars) and α -naphthyl esters (α N, grey bars) with different acyl chain lengths (reaction temperature 30 °C, see Materials and Methods for details).

Fig. 3. Activity temperature profiles and thermostability of purified metagenomic carboxylesterases from Ischia. (A) Esterase activity of purified enzymes with *p*NP-butyrate at different temperatures. (B) Thermostability of purified enzymes measured as residual activity after 20 min preincubation at different temperatures. Esterase activity was determined with *p*NP-butyrate as substrate at 30 °C. "Ctrl" corresponds to the activity measured at 30 °C without 20 min of pre-incubation.

Fig. 4. Thermoinactivation of purified IS11 (A) and IS12 (B) at different temperatures. Activity data are presented as relative activity from triplicate measurements \pm SD. Residual activity was determined with *p*NP-butyrate at 30 °C.

Fig. 5. Hydrolytic activity of purified IS10 and IS12 against the mycotoxin T-2: HPLC analysis of reaction products. Purified IS10 and IS12 were incubated with T-2 (at 37 °C and pH 8.0), and reaction products were analysed using HPLC (see Materials and Methods for experimental details).

- **Fig. 6.** Polyesterase activity of metagenomic esterases against PLA and 3PET. (A) plate assay
- 673 with emulsified 3PET as substrate. The formation of a clear zone around the wells with
- loaded enzyme indicates the presence of polyesterase activity. Agarose plates (1.5%)
- 675 containing 0.2 % emulsified 3PET and loaded proteins (50 μg/well) were incubated at 37 °C
- and monitored for three days. Porcine liver esterase (PLE), bovine serum albumin (BSA) and
- elution buffer (EB) were used as a negative, esterase MGS0105 characterised earlier (45) as a
- positive control. (B) HPLC assay of 3PET hydrolysis products after 16 h of incubation at 30
- ^oC, elution buffer was used as a negative control (not shown). (C) HPLC analysis of
- 680 hydrolysis of PLA incubated with metagenomic esterases for 48 hrs at 30 °C.
- 681 Fig. 7. Crystal structure of IS11. (A), Schematic representation of the IS11 domains: the N-
- terminal β -lactamase related Ser hydrolase domain is coloured cyan with all-helical sub-
- 683 domain shown in light blue, whereas the C-terminal lipocalin domain in orange. (B), overall
- fold of the IS11 protomer shown in three views related by 90° rotations. The protein domains
- are shown as ribbon diagrams with the core domain (β -lactamase) coloured pale cyan,
- 686 whereas the C-terminal lipocalin domain is coloured light orange. The position of the active
- site is indicated by the side chains of catalytic Ser68, Lys71, and Tyr160, whereas the protein
- 688 N- and C-terminal ends are labelled (N and C).

Fig. 8. Crystal structure of the IS11 β -lactamase and lipocalin domains. (A), The N-terminal

- 690 β -lactamase-like domain with two sub-domains coloured pale cyan (α/β) and light pink (all-
- helical). (B), The lipocalin domain. The domains are shown in three views related by 90°
- 692 rotations with the N- and C-termini labelled (N and C).
- **Fig. 9.** Close-up view of the IS11 active site. (A), The core domain showing the active site 10^{-1} 10^{-1} 10^{-1} 10^{-1}

cleft with catalytic residues: motif-1 (Ser68 and Lys71), motif-2 (Tyr160 and Asn162), and
 motif-3 (His299). The magenta-coloured mesh represents an additional electron density (a

- 2Fo-Fc omit map contoured at 2.5 σ) covalently attached to the Ser68 side chain. (B), The
- proline-rich loop of the lipocalin domain covering the active site and residues potentially
- contributing to substrate binding. Protein ribbon diagrams are coloured grey (the β -lactamase
- domain) and light orange (the lipocalin domain), whereas the side chains of residues are
- shown as sticks with green and orange carbons, respectively
- **Fig. 10.** Crystal structure of the IS11 lipocalin domain: ligand binding site and proline-rich
- 102 loop. The protein ribbon diagram is coloured in grey with the residues of ligand binding
- 703 pocket shown as sticks with green carbons and labelled.

704

705 TABLES

Table 1. Novel carboxylesterases from the Ischia polyester enrichment metagenomesselected for biochemical and structural characterisation in this study.

Protein name	Fosmid ID	Protein length	Predicted M.w.	Protein superfamily	Host organism (phylum)
IS10	L2B6_15	314 aa	34.3 kDa	α/β hydrolase	Chloroflexi
IS11	L2F9_18	455 aa	49.4 kDa	β-lactamase	Chloroflexi
IS12	L3G23 11	318 aa	33.9 kDa	α/β hydrolase	Chloroflexi

Table 2. Kinetic parameters of purified metagenomic carboxylesterases from the Ischia
 hydrothermal vents with model esterase substrates^a.

Protein	Substrate	$K_{\rm M}({\rm mM})$	$k_{\text{cat}} (s^{-1})$	$k_{\rm cat}/K_{\rm M}~({\rm s}^{-1}~{\rm M}^{-1})$		
	pNP-acetate	0.05 ± 0.01	41.97 ± 1.79	7.9×10^5		
	pNP-butyrate	0.06 ± 0.01	66.21 ± 3.24	1.2×10^6		
IS10	pNP-hexanoate	0.04 ± 0.01	86.79 ± 3.06	2.0×10^6		
	αN-propionate	0.06 ± 0.02	31.20 ± 1.93	$5.0 \ge 10^5$		
	αN-butyrate	0.12 + 0.04	58.60 ± 5.89	$4.9 \ge 10^5$		
	pNP-acetate	0.53 ± 0.31	1.60 ± 0.30	3.0×10^3		
1011	<i>p</i> NP-butyrate	0.20 ± 0.02	68.81 ± 1.37	3.5×10^5		
1511	pNP-hexanoate	0.08 ± 0.02	40.28 ± 1.39	5.3×10^5		
	αN-butyrate	0.09 ± 0.02	5.93 ± 0.49	6.9×10^4		
	pNP-acetate	0.22 ± 0.05	57.10 ± 3.78	2.6×10^5		
1012	<i>p</i> NP-butyrate	0.08 ± 0.01	8.77 ± 0.29	$1.1 \ge 10^5$		
1512	pNP-hexanoate	0.09 ± 0.01	19.05 ± 0.50	2.1×10^5		
	αN-propionate	0.69 ± 0.19	39.45 ± 3.7	5.7×10^4		

^a Reaction conditions were as indicated in Materials and Methods (pH 9.0, 30°C). Results are mean \pm SD of three independent experiments. $\alpha N = \alpha$ -naphthyl, pNP = p-nitrophenyl.

7	2	4

725

726

727 **REFERENCES**

728 1. Kyrpides NC, Hugenholtz P, Eisen JA, Woyke T, Goker M, Parker CT, Amann R, 729 Beck BJ, Chain PS, Chun J, Colwell RR, Danchin A, Dawyndt P, Dedeurwaerdere T, DeLong EF, Detter JC, De Vos P, Donohue TJ, Dong XZ, Ehrlich DS, Fraser C, Gibbs R, 730 731 Gilbert J, Gilna P, Glockner FO, Jansson JK, Keasling JD, Knight R, Labeda D, Lapidus A, 732 Lee JS, Li WJ, Ma J, Markowitz V, Moore ER, Morrison M, Meyer F, Nelson KE, Ohkuma M, Ouzounis CA, Pace N, Parkhill J, Qin N, Rossello-Mora R, Sikorski J, Smith D, Sogin M, 733 734 Steven R, Stingl U, Suzuki K, Taylor D, Tiedje JM, Tindall, B, Wagner M, Weinstock G, 735 Weissenbach J, White O, Wang J, Zhang L, Zhou YG, Field D, Whitman WB, Garrity GM, 736 Klenk HP. 2014. Genomic encyclopedia of bacteria and archaea: sequencing a myriad of type 737 strains. PLoS Biol 12:e1001920 738 2. Yarza P, Yilmaz P, Pruesse E, Glockner FO, Ludwig W, Schleifer KH, Whitman WB, 739 Euzeby J, Amann R, Rossello-Mora R. 2014. Uniting the classification of cultured and uncultured bacteria and archaea using 16S rRNA gene sequences. Nat Rev Microbiol 12:635-740 741 645 742 3. Rappe MS, Giovannoni SJ. 2003. The uncultured microbial majority. Annu Rev 743 Microbiol 57:369-394 744 Torsvik V, Goksoyr J, Daae FL. 1990. High diversity in DNA of soil bacteria. Appl 4. 745 Environ Microbiol 56:782-787 746 5. Handelsman J. 2004. Metagenomics: application of genomics to uncultured 747 microorganisms. Microbiol Mol Biol Rev 68:669-685 Ferrer M, Golyshina O, Beloqui A, Golyshin PN. 2007. Mining enzymes from 748 6. extreme environments. Curr Opin Microbiol 10:207-214 749 Uchiyama T, Miyazaki K. 2009. Functional metagenomics for enzyme discovery: 750 7. challenges to efficient screening. Curr Opin Biotechnol 20:616-622 751 Turnbaugh PJ, Gordon JI. 2008. An invitation to the marriage of metagenomics and 752 8. metabolomics. Cell 134:708-713 753 754 9. Venter JC, Remington K, Heidelberg JF, Halpern AL, Rusch D, Eisen JA, Wu D, 755 Paulsen I, Nelson KE, Nelson W, Fouts DE, Levy S, Knap AH, Lomas MW, Nealson K, White O, Peterson J, Hoffman J, Parsons R, Baden-Tillson H, Pfannkoch C, Rogers YH, 756 757 Smith HO. 2004. Environmental genome shotgun sequencing of the Sargasso Sea. Science 304:66-74 758 759 10. Rusch DB, Halpern AL, Sutton G, Heidelberg KB, Williamson S, Yooseph S, Wu D, 760 Eisen JA, Hoffman JM, Remington K, Beeson K, Tran B, Smith H, Baden-Tillson H, Stewart C, Thorpe J, Freeman J, Andrews-Pfannkoch C, Venter JE, Li K, Kravitz S, Heidelberg JF, 761 762 Utterback T, Rogers YH, Falcón LI, Souza V, Bonilla-Rosso G, Eguiarte LE, Karl DM,

763 Sathyendranath S, Platt T, Bermingham E, Gallardo V, Tamayo-Castillo G, Ferrari MR, Strausberg RL, Nealson K, Friedman R, Frazier M, Venter JC. 2007. The Sorcerer II Global 764 Ocean Sampling expedition: northwest Atlantic through eastern tropical Pacific. PLoS Biol 765 5:e77 766 11. 767 Yooseph S, Sutton G, Rusch DB, Halpern AL, Williamson SJ, Remington K, Eisen 768 JA, Heidelberg KB, Manning G, Li W, Jaroszewski L, Cieplak P, Miller CS, Li H, 769 Mashiyama ST, Joachimiak MP, van Belle C, Chandonia JM, Soergel DA, Zhai Y, Natarajan 770 K, Lee S, Raphael BJ, Bafna V, Friedman R, Brenner SE, Godzik A, Eisenberg D, Dixon JE, 771 Taylor SS, Strausberg RL, Frazier M, Venter JC. 2007. The Sorcerer II Global Ocean 772 Sampling expedition: expanding the universe of protein families. PLoS Biol 5:e16 773 12. Qin J, Li R, Raes J, Arumugam M, Burgdorf KS, Manichanh C, Nielsen T, Pons N, 774 Levenez F, Yamada T, Mende DR, Li J, Xu J, Li S, Li D, Cao J, Wang B, Liang H, Zheng H, 775 Xie Y, Tap J, Lepage P, Bertalan M, Batto JM, Hansen T, Le Paslier D, Linneberg A, Nielsen 776 HB, Pelletier E, Renault P, Sicheritz-Ponten T, Turner K, Zhu H, Yu C, Li S, Jian M, Zhou 777 Y, Li Y, Zhang X, Li S, Qin N, Yang H, Wang J, Brunak S, Doré J, Guarner F, Kristiansen 778 K, Pedersen O, Parkhill J, Weissenbach J; MetaHIT Consortium, Bork P, Ehrlich SD, Wang 779 J. 2010. A human gut microbial gene catalogue established by metagenomic sequencing. 780 Nature 464:59-65 781 13. Hess M, Sczyrba A, Egan R, Kim TW, Chokhawala H, Schroth G, Luo S, Clark DS, Chen F, Zhang T, Mackie RI, Pennacchio LA, Tringe SG, Visel A, Woyke T, Wang Z, Rubin 782 783 EM. 2011. Metagenomic discovery of biomass-degrading genes and genomes from cow 784 rumen. Science 331:463-467 785 14. Ferrer M, Martínez-Martínez M, Bargiela R, Streit WR, Golyshina OV, Golyshin PN. 786 2016. Estimating the success of enzyme bioprospecting through metagenomics: current status 787 and future trends. Microb Biotechnol 9:22-34 788 15. Levitt M. 2009. Nature of the protein universe. Proc Natl Acad Sci U S A 106:11079-789 11084 790 16. Godzik A. 2011. Metagenomics and the protein universe. Curr Opin Struct Biol 791 21:398-403 792 17. Dinsdale EA, Edwards RA, Hall D, Angly F, Breitbart M, Brulc JM, Furlan M, 793 Desnues C, Haynes M, Li L, McDaniel L, Moran MA, Nelson KE, Nilsson C, Olson R, Paul 794 J, Brito BR, Ruan Y, Swan BK, Stevens R, Valentine DL, Thurber RV, Wegley L, White 795 BA, Rohwer F. 2008. Functional metagenomic profiling of nine biomes. Nature 452:629-632 796 18. Rondon MR, August PR, Bettermann AD, Brady SF, Grossman TH, Liles MR, 797 Loiacono KA, Lynch BA, MacNeil IA, Minor C, Tiong CL, Gilman M, Osburne MS, Clardy 798 J, Handelsman J, Goodman RM. 2000. Cloning the soil metagenome: a strategy for accessing 799 the genetic and functional diversity of uncultured microorganisms. Appl Environ Microbiol 800 66:2541-2547 801 19. Simon C, Daniel R. 2011. Metagenomic analyses: past and future trends. Appl 802 Environ Microbiol 77:1153-1161

- 803 20. Robertson DE, Chaplin JA, DeSantis G, Podar M, Madden M, Chi E, Richardson T, 804 Milan A, Miller M, Weiner DP, Wong K, McQuaid J, Farwell B, Preston LA, Tan X, Snead 805 MA, Keller M, Mathur E, Kretz PL, Burk MJ, Short JM. 2004. Exploring nitrilase sequence space for enantioselective catalysis. Appl Environ Microbiol 70:2429-2436 806 807 21. Lorenz P, Eck J. 2005. Metagenomics and industrial applications. Nat Rev Microbiol 808 3:510-516 809 22. Bornscheuer UT. 2002. Microbial carboxyl esterases: classification, properties and 810 application in biocatalysis. FEMS Microbiol Rev 26:73-81 811 23. Martínez-Martínez M, Coscolín C, Santiago G, Chow J, Stogios PJ, Bargiela R, Gertler C, Navarro-Fernández J, Bollinger A, Thies S, Méndez-García C, Popovic A, Brown 812 813 G, Chernikova TN, García-Moyano A, Bjerga GEK, Pérez-García P, Hai T, Del Pozo MV, 814 Stokke R, Steen IH, Cui H, Xu X, Nocek BP, Alcaide M, Distaso M, Mesa V, Peláez AI,
- 815 Sánchez J, Buchholz PCF, Pleiss J, Fernández-Guerra A, Glöckner FO, Golyshina OV,
- 816 Yakimov MM, Savchenko A, Jaeger KE, Yakunin AF, Streit WR, Golyshin PN, Guallar V,
- 817 Ferrer M, The Inmare Consortium. 2018. Determinants and prediction of esterase substrate
- 818 promiscuity patterns. ACS Chem Biol 13:225-234
- Arpigny JL, Jaeger KE. 1999. Bacterial lipolytic enzymes: classification and
 properties. Biochem J 343(Pt 1):177-183
- 25. Lenfant N, Hotelier T, Velluet E, Bourne Y, Marchot P, Chatonnet A. 2013.
- ESTHER, the database of the alpha/beta-hydrolase fold superfamily of proteins: tools to
 explore diversity of functions. Nucleic Acids Res 41:D423-429
- 26. Littlechild JA. 2017. Improving the 'tool box' for robust industrial enzymes. J Ind
 Microbiol Biotechnol 44:711-720
- 826 27. Popovic A, Hai T, Tchigvintsev A, Hajighasemi M, Nocek B, Khusnutdinova AN,
- Brown G, Glinos J, Flick R, Skarina T, Chernikova TN, Yim V, Brüls T, Paslier DL,
- 828 Yakimov MM, Joachimiak A, Ferrer M, Golyshina OV, Savchenko A, Golyshin PN,
- Yakunin AF. 2017. Activity screening of environmental metagenomic libraries reveals novel
 carboxylesterase families. Sci Rep 7:44103
- 28. Pellis A, Cantone S, Ebert C, Gardossi L. 2018. Evolving biocatalysis to meet
 bioeconomy challenges and opportunities. N Biotechnol 40:154-169
- Antranikian G, Streit WR. 2022. Microorganisms harbor keys to a circular
 bioeconomy making them useful tools in fighting plastic pollution and rising CO2 levels.
 Extremophiles 26:10
- 836 30. Kruger A, Schafers C, Schroder C, Antranikian G. 2018. Towards a sustainable
 837 biobased industry Highlighting the impact of extremophiles. N Biotechnol 40:144-153
- Atomi H. 2005. Recent progress towards the application of hyperthermophiles and
 their enzymes. Curr Opin Chem Biol 9:166-173
- 840 32. Littlechild JA. 2015. Archaeal enzymes and applications in industrial biocatalysts.
 841 Archaea 2015:147671

842 33. Vieille C, Zeikus GJ. 2001. Hyperthermophilic enzymes: sources, uses, and molecular 843 mechanisms for thermostability. Microbiol Mol Biol Rev 65:1-43 844 Alcaide M, Stogios PJ, Lafraya Á, Tchigvintsev A, Flick R, Bargiela R, Chernikova 34. 845 TN, Reva ON, Hai T, Leggewie CC, Katzke N, La Cono V, Matesanz R, Jebbar M, Jaeger 846 KE, Yakimov MM, Yakunin AF, Golyshin PN, Golyshina OV, Savchenko A, Ferrer M; 847 MAMBA Consortium. 2015. Pressure adaptation is linked to thermal adaptation in salt-848 saturated marine habitats. Environ Microbiol 17:332-345 849 35. Wei R, von Haugwitz G, Pfaff L, Mican J, Badenhorst CPS, Liu W, Weber G, Austin 850 HP, Bednar D, Damborsky J, Bornscheuer UT. 2022. Mechanism-based design of efficient 851 PET hydrolases. ACS Catalysis 12:3382-3396 852 36. Heinl S, Hartinger D, Thamhesl M, Vekiru E, Krska R, Schatzmavr G, Moll WD, 853 Grabherr R. 2010. Degradation of fumonisin B1, by the consecutive action of two bacterial 854 enzymes. J Biotechnol 145: 120-129 855 37. Rizzo C, Arcadi E, Calogero R, Sciutteri V, Consoli P, Esposito V, Canese S, 856 Andaloro F, Romeo T. 2022. Ecological and biotechnological relevance of Mediterranean 857 hydrothermal vent systems. Minerals 12:251 858 38. Fadrosh DW, Ma B, Gajer P, Sengamalay N, Ott S, Brotman RM, Ravel J. 2014. An 859 improved dual-indexing approach for multiplexed 16S rRNA gene sequencing on the 860 Illumina MiSeq platform. Microbiome 2:6 861 39. Distaso MA, Bargiela R, Brailsford FL, Williams GB, Wright S, Lunev EA, 862 Toshchakov SV, Yakimov MM, Jones DL, Golyshin PN, Golyshina OV. 2020 High representation of archaea across all depths in oxic and low-pH sediment layers underlying an 863 864 acidic stream. Front Microbiol 11:2871. 865 40. R Core Team. 2020. R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. www.R-project.org/. 866 41. Placido A, Hai T, Ferrer M, Chernikova TN, Distaso M, Armstrong D, Yakunin AF, 867 868 Toshchakov SV, Yakimov MM, Kublanov IV, Golyshina OV, Pesole G, Ceci LR, Golyshin 869 PN. 2015. Diversity of hydrolases from hydrothermal vent sediments of the Levante Bay, 870 Vulcano Island Aeolian archipelago identified by activity-based metagenomics and biochemical characterization of new esterases and an arabinopyranosidase. Appl Microbiol 871 872 Biotechnol 99:10031-10046 873 42. Zhu W, Lomsadze A, Borodovsky M. 2010. Ab initio gene identification in 874 metagenomic sequences. Nucleic Acids Res 38:e132 43. 875 Altschul SF, Madden TL, Schaffer AA, Zhang J, Zhang Z, Miller W, Lipman DJ. 876 1997. Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res 25:3389-3402 877 878 44. Edgar RC. 2004. MUSCLE: multiple sequence alignment with high accuracy and 879 high throughput. Nucleic Acids Res 32:1792-1797 880 45. Kumar S, Stecher G, Li M, Knyaz C, Tamura K. 2018. MEGA X: Molecular 881 evolutionary genetics analysis across computing platforms. Mol Biol Evol 35:1547-1549

882 46. Tchigvintsev A, Tran H, Popovic A, Kovacic F, Brown G, Flick R, Hajighasemi M, Egorova O, Somody JC, Tchigvintsev D, Khusnutdinova A, Chernikova TN, Golyshina OV, 883 884 Yakimov MM, Savchenko A, Golyshin PN, Jaeger KE, Yakunin AF. 2015. The environment 885 shapes microbial enzymes: five cold-active and salt-resistant carboxylesterases from marine 886 metagenomes. Appl Microbiol Biotechnol 99:2165-2178 887 47. Guinta CI, Cea-Rama I, Alonso S, Briand ML, Bargiela R, Coscolin C, Corvini P, 888 Ferrer M, Sanz-Aparicio J, Shahgaldian P. 2020. Tuning the properties of natural 889 promiscuous enzymes by engineering their nano-environment. ACS Nano 14:17652-17664 890 48. Hajighasemi M, Tchigvintsev A, Nocek B, Flick R, Popovic A, Hai T, Khusnutdinova 891 AN, Brown G, Xu X, Cui H, Anstett J, Chernikova TN, Brüls T, Le Paslier D, Yakimov MM, 892 Joachimiak A, Golyshina OV, Savchenko A, Golyshin PN, Edwards EA, Yakunin AF. 2018. 893 Screening and Characterization of Novel Polyesterases from Environmental Metagenomes 894 with High Hydrolytic Activity against Synthetic Polyesters. Environ Sci Technol 52:12388-895 12401 896 49. Minor W, Cymborowski M, Otwinowski Z, Chruszcz M. 2006. HKL-3000: the 897 integration of data reduction and structure solution--from diffraction images to an initial 898 model in minutes. Acta Crystallogr D Biol Crystallogr 62:859-866 899 50. Liebschner D, Afonine PV, Baker ML, Bunkóczi G, Chen VB, Croll TI, Hintze B, 900 Hung LW, Jain S, McCoy AJ, Moriarty NW, Oeffner RD, Poon BK, Prisant MG, Read RJ, 901 Richardson JS, Richardson DC, Sammito MD, Sobolev OV, Stockwell DH, Terwilliger TC, 902 Urzhumtsev AG, Videau LL, Williams CJ, Adams PD.2019 Macromolecular structure 903 determination using X-rays, neutrons and electrons: recent developments in Phenix. Acta 904 Crystallogr D Struct Biol 75:861-877 905 51. Jumper J, Evans R, Pritzel A, Green T, Figurnov M, Ronneberger O, Tunyasuvunakool K, Bates R, Žídek A, Potapenko A, Bridgland A, Meyer C, Kohl SAA, 906 907 Ballard AJ, Cowie A, Romera-Paredes B, Nikolov S, Jain R, Adler J, Back T, Petersen S, 908 Reiman D, Clancy E, Zielinski M, Steinegger M, Pacholska M, Berghammer T, Bodenstein 909 S, Silver D, Vinyals O, Senior AW, Kavukcuoglu K, Kohli P, Hassabis D. 2021 Highly 910 accurate protein structure prediction with AlphaFold. Nature 596:583-589 911 52. Emsley P, Cowtan K. 2004. Coot: model-building tools for molecular graphics. Acta 912 Crystallogr D Biol Crystallogr 60:2126-2132 913 53. Zhou Z, Liu Y, Xu W, Pan J, Luo ZH, Li M. 2020. Genome- and Community-Level 914 Interaction Insights into Carbon Utilization and Element Cycling Functions of 915 Hydrothermarchaeota in Hydrothermal Sediment. mSystems 5:e00795-19 916 54. Park YJ, Yoon SJ, Lee HB. 2008. A novel thermostable arylesterase from the 917 archaeon Sulfolobus solfataricus P1: purification, characterization, and expression. J 918 Bacteriol 190:8086-8095 919 55. Pereira MR, Maester TC, Mercaldi GF, de Macedo Lemos EG, Hyvonen M, Balan A. 920 2017. From a metagenomic source to a high-resolution structure of a novel alkaline esterase.

921 Appl Microbiol Biotechnol 101:4935-4949

- 56. Coque JJ, Liras P, Martin JF. 1993. Genes for a beta-lactamase, a penicillin-binding
 protein and a transmembrane protein are clustered with the cephamycin biosynthetic genes in
 Nocardia lactamdurans. EMBO J 12:631-639
- 925 57. Petersen EI, Valinger G, Solkner B, Stubenrauch G, Schwab H. 2001. A novel
 926 esterase from Burkholderia gladioli which shows high deacetylation activity on
 927 cephalosporins is related to beta-lactamases and DD-peptidases. J Biotechnol 89:11-25
- 58. Lewin A, Strand T, Haugen T, Klinkenberg G, Kotlar H, Valla S., Drablos F, Wentze,
 A. 2016. Discovery and characterization of a thermostable esterase from an oil reservoir
 metagenome. Adv Enzyme Res 4:68-86
- 59. Leis B, Angelov A, Mientus M, Li H, Pham VT, Lauinger B, Bongen P, Pietruszka J,
 Goncalves LG, Santos H, Liebl W. 2015. Identification of novel esterase-active enzymes
 from hot environments by use of the host bacterium *Thermus thermophilus*. Front Microbiol
 6:275
- 60. Miguel-Ruano V, Rivera I, Rajkovic J, Knapik K, Torrado A, Otero JM, Beneventi E,
 Becerra M, Sánchez-Costa M, Hidalgo A, Berenguer J, González-Siso MI, Cruces J, Rúa
 ML, Hermoso JA. 2021. Biochemical and structural characterization of a novel thermophilic
 esterase EstD11 provide catalytic insights for the HSL family. Comput Struct Biotechnol J
 19:1214-1232
- Sayer C, Szabo Z, Isupov MN, Ingham C, Littlechild JA. 2015. The structure of a
 novel thermophilic esterase from the Planctomycetes species, *Thermogutta terrifontis* reveals
 an open active site due to a minimal 'cap' domain. Front Microbiol 6:1294
- 62. Loi M, Fanell, F, Liuzzi VC, Logrieco AF, Mule G. 2017. Mycotoxin
 Biotransformation by native and commercial enzymes: Present and future perspectives.
 Toxins Basel 9:111
- 63. Liu L, Xie M, We, D. 2022. Biological Detoxification of Mycotoxins: Current Status
 and Future Advances. Int J Mol Sci 23:1064
- 64. Lyagin I, and Efremenko E. 2019. Enzymes for detoxification of various mycotoxins:
 origins and mechanisms of catalytic action. Molecules 24:2362
- 950 65. McCormick SP, Price NP, Kurtzman CP 2012. Glucosylation and other
 951 biotransformations of T-2 toxin by yeasts of the trichomonascus clade. Appl Environ
 952 Microbiol 78:8694-8702
- 953 66. Tournier V, Topham CM, Gilles A, David B, Folgoas C, Moya-Leclair E, Kamionka
- 954 E, Desrousseaux ML, Texier H, Gavalda S, Cot M, Guémard E, Dalibey M, Nomme J, Cioci
- 955 G, Barbe S, Chateau M, André I, Duquesne S, Marty A. 2020. An engineered PET
- depolymerase to break down and recycle plastic bottles. Nature 580:216-219.
- 957 67. Palm GJ, Fernández-Álvaro E, Bogdanović X, Bartsch S, Sczodrok J, Singh RK,
- Böttcher D, Atomi H, Bornscheuer UT, Hinrichs W. 2011. The crystal structure of an
- 959 esterase from the hyperthermophilic microorganism Pyrobaculum calidifontis VA1 explains
- 960 its enantioselectivity. Appl Microbiol Biotechnol 91:1061-1072

- 68. Sauvage E, Kerff F, Terrak M, Ayala JA, Charlier P. 2008. The penicillin-binding
 proteins: structure and role in peptidoglycan biosynthesis. FEMS Microbiol Rev 32:234-258
- 69. Lee D, Das S, Dawson NL, Dobrijevic D, Ward J, Orengo C. 2016. Novel
 64 computational protocols for functionally classifying and characterising serine beta65 lactamases. PLoS Comput Biol 12:e1004926
- 70. Holm L. 2022. Dali server: structural unification of protein families. Nucleic Acids
 867 Res 50:W210-W215
- 968 71. Wagner UG, Petersen EI, Schwab H, Kratky C. 2002. EstB from *Burkholderia*969 *gladioli*: a novel esterase with a beta-lactamase fold reveals steric factors to discriminate
 970 between esterolytic and beta-lactam cleaving activity. Protein Sci 11:467-478
- 971 72. Delfosse V, Girard E, Birck C, Delmarcelle M, Delarue M, Poch O, Schultz O, Mayer
 972 C. 2009. Structure of the archaeal pab87 peptidase reveals a novel self-compartmentalizing
 973 protease family. PLoS One 4:e4712
- 73. Flower DR, North AC, Sansom CE. 2000. The lipocalin protein family: structural and
 sequence overview. Biochim Biophys Acta 1482:9-24
- 976 74. De Simone G, Galdiero S, Manco G, Lang D, Rossi M, Pedone C. 2000. A snapshot
 977 of a transition state analogue of a novel thermophilic esterase belonging to the subfamily of
 978 mammalian hormone-sensitive lipase. J Mol Biol 303:761-771
- 75. Kay BK, Williamson MP, Sudol M. 2000. The importance of being proline: the
 interaction of proline-rich motifs in signaling proteins with their cognate domains. FASEB J
 14:231-241

982

983

Relative abundance (%)			×	Before enrichment		Polyhydroxy- butyrate (PHB)			Polylactic acid (PLA)				Commercial polyester blend								
>	90 >75 >50 >40 >30	>20 >	•10 >5	>1 >0.5 <	0.5	IS1	IS2	IS3	IS4	IS1	IS2	IS3	IS4	IS1	IS2	IS3	IS4	IS1	IS2	IS3	IS4
Λ	RCHAEA		Th	ermoprote	ota-									_							
Cutibacteriu																					
Actinobacteriota Conexibacte					ter-						×				···*			*			
		Other Actinobacteriota-				•			•		*	×			*	····×			····×		
	Restargidata			Schleife	eria-			×		•		*			···*	····*		····• ••···			
	Bacteroluota	Other Flavobacteriales- Chitinophagales-			les-	•		.	•										x		
					les-	*				*	*			*	×			••			
				Cytophaga	les-	*		·····		•		*		····•••				····•			
			Sphingobacterium-		um-	•		×	••••												
		Othe	er Sphin	gobacteria	les-					×		*		*				••••			
			lgr	navibacteri	um-	*				*	•			*	••••			*			
			Igna	avibacteria	les-					•				••••				••••			
			Ka	pabacteria	les-	*			×	•		*		•	••••	*		•	•		
			Other	Bacteroid	ota-	*												*	*		
	Chloroflexota	Chloroflexota Ther						•••••	••••			*									
	Chiefonokota		Other	r Chlorofle	xia-			×	••••			••••				*	*			×	
				Anaeroline	eae-	••••				*	*				••••			×			
	Distant		Other C	Chloroflexi	ota-	•				*		×		×		*		••••			
	Deinococcota			Therm	nus-	•	••••		×												
	Firmicutes			Geobacil	lus-		×				*									···· · ···	
				Bacil	ius-														×		
				Brevibacii	ius-				1		*					*		*	*	*	
				Othor Boy			<u></u>														
			Oth	or Eirmicu			<u> </u>											î			
			Our	Haliandi		Î	<u></u>														
	Myxococcota		Other	Myxococc	ota-										*			Î			
			Ac	etobactera	les-				×	×	<u>×</u>	*		×		×		×	x		
	Proteobacteria		Ca	ulobactera	les-									×	x				×		
		α	04	Rhizobia	les-	*		×	×	×	x			*							
			Rho	odobactera	les-									*				*			
	Oth		ther α-P	roteobacte	eria-	×		.			. <mark>*</mark>	*		*		*					
			Caldimonas		nas-		🙆									×		····•			
		Other Burkholderiales- Rehaibacterium-		les-	•		×				*			🖕							
				um-	×			×		×	×		••••				•••••	·····•			
	Other Xanthomonadales-		les-	••••		×	····•••	×				····•••	····••••···	*		····· ••···					
		Fontimonas-			nas-	*	×	×	.					••••		····×		••••	····••••···		
		Pseudomonas-			nas-	••••		.	····•												
		0	Other y-Proteobacteria-		eria-	•			x			*		*	•	*		••••	•		
	Other Phyla	Acidobacteriota-			ota-	•	*			*	*	*		*	••••	***		*	× .		
	other r fiyia		Arn	natimonad	ota-	••••			×					*	*				×		
Desulfobacterota			••••									*									
Nitrospirota				••••					*	*			*	×			× .				
Planctomycetota- Spirochaetota- SAR324- Vorrugomiarobiota									×	••••	×		×	••••	*		×	×			
									×				×	••••				••••			
									×	*			×	***			×	••••			
PAOTEDIA Verrucomicrobiota-			ota	*				×	*			*		*		×	X				
			0	ither Bacte	eria –			·····X····		····X·····	···X····	·····X·····		🔘		·····X····		·····X····			





Assay temperature (°C)





Temperature °C













