Characterization of an Extracellular Polyhydroxyalkanoate Depolymerase from *Streptomyces* sp. SFB5A

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**Abstract**

A poly(-3-hydroxybutyrate) (PHB) degrading bacterium, *Streptomyces* sp. SFB5A, was isolated from hardwood mulch. It synthesized an extracellular poly(3-hydroxyalkanoate) (PHA) depolymerase during growth on PHB, poly(3-hydroxyvalerate) (PHV), PHB-PHV copolymer (PHBV), or 3-hydroxybutyrate. The purified enzyme had a subunit relative molecular weight of 47,000 and a broad pH optimum of 7.0 to 8.5; was stimulated by Ca2+ and Mg2+; and was inhibited by ethylenediaminetetraacetic acid, dithiothreitol, and non-ionic detergents. The enzyme degraded PHB, PHV, and PHBV to a mixture of monomers, dimers, and trimers, with monomers predominating. The level of trimers peaked and then decreased over time, suggesting that they were produced early and were subsequently degraded to monomers and dimers. The enzyme did not degrade poly(3-hydroxyoctanoate), indicating that it was a short-chain-length PHA depolymerase. The PHA depolymerase gene was cloned and sequenced. The deduced amino acid sequence included three features typical of extracellular PHA depolymerases: a catalytic domain type 1, a fibronectin type III linker domain, and a substrate-binding domain. A mechanism accounting for the transient production of PHA trimers was proposed, based on modeling of the enzyme’s tertiary structure and amino acid sequence homology to related PHA depolymerases.

**Keywords:** PHB; PHV; Polyhydroxyalkanoates; PHA depolymerase; *Streptomyces*; Catalytic type 1 domain; FnIII linker domain; Substrate-binding domain

**Introduction**

Petroleum-based plastics present two major problems: they require a non-renewable fossil fuel for their manufacture and are recalcitrant to degradation upon disposal. Poly(3-hydroxyalkanoates) (PHAs) are water-insoluble polymers of R-3-hydroxyalkanoic acid monomers in chains of 100 to 3000 produced by many bacteria for storage of carbon and energy [1]. They represent a biodegradable alternative to petroleum-based plastics for certain applications and have been produced commercially on a limited scale. Short-chain-length PHAs (PHA<sub>SCL</sub>) consist of monomers with 3 to 5 carbon atoms [2]. The most common of these is poly(3-hydroxybutyrate) (PHB), composed of R-3-hydroxybutanoic acid (3HB) monomers [1]. Another common PHA<sub>SCL</sub> is poly(R-3-hydroxyvalerate) (PHV), composed of R-3-hydroxyvaleric acid (3HV) monomers. Some bacteria produce heteropolymers (PHBV) composed of 3HB and 3HV monomers in various proportions [1]. Medium-chain-length PHAs (PHA<sub>MCL</sub>), such as poly(R-3-hydroxyoctanoate) (PHO), consist of monomers containing 6 or more carbon atoms [2]. Native, intracellular PHAs (nPHAs) from PHA-producing bacteria have an amorphous structure, which is usually altered upon removal from the cell or chemical treatments, forming denatured, i.e., partially crystalline PHAs (dPHAs) [2]. Amorphous analogs of nPHAs, abbreviated aPHAs, can be prepared by treatment of chloroform solutions of dPHAs with detergents [3].

Many microorganisms degrade dPHAs, using extracellular PHA depolymerases. These are secreted into the surrounding medium and hydrolyze PHAs to their monomers, which are transported into the cells and catabolized. Numerous PHA depolymerases have been purified and characterized (reviewed in references 2 and 4). Biosynthesis of PHA depolymerases occurs during growth on PHAs or their monomers, but is repressed by sugars, organic acids, and amino acids [2, 5-8].

*Streptomyces* is a very broad genus of Gram positive, filamentous bacteria with high % G+C (~70%) that produce abundant reproductive spores and various antibiotics [9]. They degrade a variety of polymers including: agar, starch, cellulose, chitin, and xylan, with degradation catalyzed by polymer-specific hydrolyases [10]. *In situ* soil studies show that numerous *Streptomyces* species degrade PHB and PHBV as well [11]. Extracellular dPHA depolymerases are reported in several *Streptomyces* species, some of which specifically degrade PHA<sub>SCL</sub> [7, 12-15], while others degrade PHA<sub>MCL</sub> [16-18].

Despite the taxonomic breadth of *Streptomyces*, comparatively few extracellular PHA depolymerases from this important genus have been described to date. We report here the isolation of a PHB-degrading actinomycete, *Streptomyces* sp. SFB5A, the purification and characterization of an extracellular dPHA<sub>SCL</sub> depolymerase from the organism, and the cloning of its associated gene. We show that the
enzyme produces mostly monomers with some dimers and trimers during PHA degradation and propose a possible model to explain this result, based on biochemical and structural data.

Materials and Methods

Chemicals

PHA and the sodium salts of R,S-3-hydroxybutanoic acid (R,S-3HB) and R-3-hydroxybutanoic acid (R-3HB) were obtained from Sigma-Aldrich, St. Louis, MO, USA. PHBV was prepared by culturing Chromobacterium violaceum with sodium valerate as described [19]. A PHBV copolymer containing 17 mol% 3HV (trade name Biopol® BX P029) was obtained from Zeneca, London, UK. PHO was prepared by culturing Pseudomonas oleovorans with sodium octanoate as described [20]. Homogeneous suspensions of granular PHB and PHBV (3% to 4% w/v) were prepared by sonication, autoclaved, and stored at 4°C. Sonicated PHV suspensions readily aggregated, requiring considerable care in pipetting. For preparation of PHA thin films, PHAs were made to 1% w/v in chloroform and heated at 50°C until dissolved. Amorphous PHB (aPHB) was prepared as described, with 50 mM sodium cholate as the detergent [3]. All other chemicals were of reagent grade or better.

Culture media and growth conditions

A defined mineral salts medium (SNC) [21] was used for experiments on PHA depolymerase synthesis. Carbon sources were added separately as filter-sterilized or autoclaved stocks. SNC containing 0.2% w/v PHB was referred to as SNC-PHB. Nutrient broth (NB), nutrient agar (NA), tryptic soy broth (TSB), trypticase soy agar (TSA) (Becton Dickinson and Company, Franklin Park, NJ, USA) and mannitol soy flour medium (MSF) [9] were used as complex media. Streptomyces sp. SFB5A was maintained on SNC-PHB agar plates. Spore suspensions were prepared from SNC plus 0.5% glucose agar cultures as described [9]. All agar plate cultures of Streptomyces sp. SFB5A were incubated at 30°C. All broth cultures were incubated with rotary shaking (200 rpm) at 30°C for times indicated.

Isolation of strain SFB5A

Well decayed hardwood mulch (1.0 g) was added to 99.0 mL of sterile 0.85% saline solution in a dilution bottle. The mixture was shaken vigorously for 30 sec and allowed to settle for 1 min. Samples (0.1 mL) of the supernatant liquid were serially diluted and plated onto SNC agar plates previously overlaid with 5 mL of SNC-PHB agar. The plates were incubated at 30°C for 48 h. Selected colonies exhibiting large clearing zones in the opaque PHB overlay were streaked from the genomic DNA by the polymerase chain reaction (PCR) with primer pairs StrepB/StrepF [23] and pA/pH [24](Table S1). Reaction mixtures included 100 ng of genomic DNA, 0.5 μM of each primer, 5% v/v dimethylsulfoxide, and an Illustra™ PuReTaq Ready-To-Go™ PCR Bead (GE Healthcare, Pittsburgh, PA, USA) in a final volume of 25 μL. The PCR schedule included an initial denaturation at 94°C for 10 min; 40 cycles of denaturation at 94°C for 30 sec, annealing at 54°C for 30 sec, and extension at 72°C for 1 min; followed by a final extension at 72°C for 10 min. 16S-rDNA amplicons were purified by agarose gel electrophoresis and extracted using a Zymoclean™ Gel DNA Recovery Kit (Zymo Research, Irvine, CA, USA). Sequencing of the amplicons was performed by Eurofins MWG Operon (Louisville, KY). The pA/pH amplicon was sequenced directly using the pA, pH, 16S InL, and 16S InR primers (Table S1). The StrepB/StrepF amplon was first ligated into a pCR®-TOPO® vector and cloned into Escherichia coli One Shot™ Top 10 cells using the TOPO® TA Cloning® Kit for Sequencing (Life Technologies, Grand Island, NY, USA). Plasmid DNA from positive transformants was sequenced using T3 and T7 primers (Table S1). Overlapping sequences were assembled into a contig with the CAP3 Sequence Assembly Program [25], available at: http://doua.prabi.fr/software/cap3. Phylogenetic homologies were determined by pairwise nucleotide sequence alignment of the contig using the EZBioCloud tool available at: https://www.ezbiocloud.net/tools/pairAlign. The assembled 16S rDNA sequence was deposited in GenBank, with the accession number MH392705.2.

Studies of PHA depolymerase synthesis

Degradation of PHB or PHBV by Streptomyces sp. SFB5A was assessed by streaking it on SNC agar plates (20 mL) overlaid with 5 mL of SNC agar containing 0.2% w/v of the polymer and observing the formation of clearing zones. To visualize the degradation of PHV during growth, 20 mg of the polymer was dissolved in 1.0 mL of CHCl₃ and poured onto the surface of an SNC agar plate. The CHCl₃ was allowed to evaporate, forming a thin film of PHV, and 5 mL of molten SNC agar was then carefully poured over the thin film to provide a stable surface for streaking.

For studies of PHA depolymerase synthesis in broth, the organism was grown for 16 h with shaking in NB. Cells were harvested by centrifugation (5,000 × g, 4°C, 10 min) and resuspended in one culture volume of SNC broth containing 20 mM soluble carbon sources (R,S-3HB or R-3HB) or PHA suspensions at 0.2% w/v. Cultures were incubated with shaking. Samples (1.0 to 2.0 mL) were withdrawn at selected times, and clarified by centrifugation. Supernatants were stored at -80°C and analyzed as soon as possible.

Purification of PHA depolymerase

A flask of NB (50 mL) was inoculated with several colonies of Streptomyces sp. SFB5A from a SNC-PHB agar plate culture and incubated with shaking for 16 h. This was used to inoculate two 1 L batches of NB contained in 4 L Erlenmeyer flasks. Cells were harvested by centrifugation (5,000 × g, 4°C, 10 min) and resuspended in one culture volume of SNC broth containing 20 mM soluble carbon sources (R,S-3HB or R-3HB) or PHA suspensions at 0.2% w/v. Cultures were incubated with shaking. Samples (1.0 to 2.0 mL) were withdrawn at selected times, and clarified by centrifugation. Supernatants were stored at -80°C and analyzed as soon as possible.

Taxonomic characterization of strain SFB5A

Genomic DNA of strain SFB 5A was isolated by a salting out method, as described [22]. A portion of the 16S-rRNA gene was amplified from the genomic DNA by the polymerase chain reaction (PCR) with primer pairs StrepB/StrepF [23] and pA/pH [24](Table S1). Reaction mixtures included 100 ng of genomic DNA, 0.5 μM of each primer, 5% v/v dimethylsulfoxide, and an Illustra™ PuReTaq Ready-To-Go™ PCR Bead (GE Healthcare, Pittsburgh, PA, USA) in a final volume of 25 μL. The PCR schedule included an initial denaturation at 94°C for 10 min; 40 cycles of denaturation at 94°C for 30 sec, annealing at 54°C for 30 sec, and extension at 72°C for 1 min; followed by a final extension at 72°C for 10 min. 16S-rDNA amplicons were purified by agarose gel electrophoresis and extracted using a Zymoclean™ Gel DNA Recovery Kit (Zymo Research, Irvine, CA, USA). Sequencing of the amplicons was performed by Eurofins MWG Operon (Louisville, KY). The pA/pH amplicon was sequenced directly using the pA, pH, 16S InL, and 16S InR primers (Table S1). The StrepB/StrepF amplon was first ligated into a pCR®-TOPO® vector and cloned into Escherichia coli One Shot™ Top 10 cells using the TOPO® TA Cloning® Kit for Sequencing (Life Technologies, Grand Island, NY, USA). Plasmid DNA from positive transformants was sequenced using T3 and T7 primers (Table S1). Overlapping sequences were assembled into a contig with the CAP3 Sequence Assembly Program [25], available at: http://doua.prabi.fr/software/cap3. Phylogenetic homologies were determined by pairwise nucleotide sequence alignment of the contig using the EZBioCloud tool available at: https://www.ezbiocloud.net/tools/pairAlign. The assembled 16S rDNA sequence was deposited in GenBank, with the accession number MH392705.2.
Buffer A containing 1 M NaCl was prepared by equilibrating with silica column and subsequent eluxation with 6.6 bed volumes of a linear NaCl gradient from 1.0 to 0 M in Buffer A, and 5.6 bed volumes of Buffer A (flow rate 1.2 mL/min, 3.6 mL fractions). Active fractions, eluted during the Buffer A wash, were pooled, concentrated by dialysis against solid sucrose, and applied to a 1.1 x 59 cm column of Sephacryl® S-100-HR (GE Healthcare) previously equilibrated with Buffer A. The column was developed with Buffer A (flow rate 0.1 mL/min, 0.4 mL fractions). Active fractions showing electrophoretic purity were pooled and stored at -80°C. To test for effects of inhibitors and ionic compounds on activity, the enzyme was first dialyzed against 20 volumes of buffer A for 15 h at 4°C with stirring.

**PHA depolymerase assays**

Routine assays of PHA depolymerase activity were performed turbidimetrically at 650 nm [26,27] at 37°C in a 1.0 mL reaction mixture containing 100 mM Tris-HCl, pH 7.9, 1 mM CaCl\(_2\), 136 µg of PHB suspension, and enzyme sample (40 to 100 ng purified enzyme or up to 200 µL of culture supernatants). Inhibitors and ionic compounds were added as desired. Activity was calculated using an extinction coefficient at 650 nm of 4.3 µL ng\(^{-1}\)/cm\(^{-1}\), with a unit defined as 1 µg of PHB degraded per min. Under these conditions, the assay was linear for up to 30 minutes, and the detection limit was about 0.4 units/mL. To test for aPHB degradation, a volume of aPHB suspension was substituted for PHB in the reaction mixture to give the same initial A650. The effect of pH on activity was determined by substituting buffers of overlapping pH for Tris-HCl pH 7.9 in the assay: 100 mM citrate (pH 3-6), piperazine-N,N'-bis(2-ethanesulfonic acid) (pH 6-7.5), Tris-HCl (pH 7-9.5), or glycine-HCl (pH 8.5-10.5).

Qualitative spot tests for PHA depolymerase activity were done as described [27] using 0.1% w/v PHA granule suspensions in 1.0% w/v agarose made up in turbidimetric assay buffer. Lipase activity was assayed by the rhodamine B-trioleoylglycerol plate assay [28].

Products resulting from degradation of PHAs were identified by electrospray ionization time-of-flight mass spectrometry (ESI-TOF-MS). The reaction mixture contained 180 µg of PHA granules or thin film, 50 mM Tris-HCl (pH 7.9), and 1 mM CaCl\(_2\) in a final volume of 1.0 mL. For thin films, before addition of the other ingredients, 180 µL of PHA from a 1% w/v stock in CHCl\(_3\) was pipetted into a prewarmed 13 × 100 mm glass screw cap tube, vortexed until the CHCl\(_3\) had evaporated, and incubated at 50°C for 30 min to drive off residual CHCl\(_3\). Purified PHA depolymerase (100 ng) was added to initiate the reactions, followed by incubation for 4 h at 37°C. For time course experiments, the reaction volume was increased from 1 mL to 3 mL, and amounts of all reaction components were tripled accordingly. After incubation, the reaction mixture was acidified to pH 2.5 with HCl. Portions (0.5 mL) were removed to microfuge tubes and centrifuged. The supernatant was extracted three times with 0.5 mL of ethyl acetate, and the organic layer dried under a stream of air at 50°C. The extraction residue was redissolved in 50 µL of 0.1% w/v formic acid and injected at 6 µL/min with a Harvard Laboratories Model 22 syringe infusion pump into an Agilent Technologies 6224 ESI-TOF mass spectrometer operated in negative ionization mode. Nitrogen was used as the desolvation gas at a flow rate of 5 L/h and 325°C, with nebulizer pressure at 30 psig. Fragmentor and skimmer voltages were 175 V and 65 V, respectively. Degradation products were identified from the predicted m/z ratios of the primary ions. A plot of 3HB concentration versus spectral abundance was linear up to at least 48 mM (6,000 ppm) and abundance levels of 2.5 million.

**Electrophoretic analyses**

Protein purity and subunit M\(_r\) were determined by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) on 12% polyacrylamide gels according to Laemmli [29], with silver staining as described [30]; Precision Plus Protein™ Unstained Standards (Bio-Rad Laboratories, Hercules, CA, USA) served as molecular mass markers. Native PAGE was performed using the Laemmli system [29] without SDS or 2-mercaptoethanol. To prepare samples for native PAGE, PHB-grown cultures were clarified by centrifugation and the supernatants made to 55% saturation (0°C) ammonium sulfate. Precipitated proteins were pelleted by centrifugation (12,000 × g, 10 min, 4°C), and pellets were redissolved in buffer RB containing 1X Laemmli sample buffer [29] without SDS and 2-mercaptoethanol. Dual zymography versus PHB and PHV was done using the same native gel. After electrophoresis of the samples, the gel was overlaid with a slab of 1% agarose (0.75 × 7 × 9 mm) containing 0.1% w/v PHB in turbidimetric assay buffer (see above). It was then placed on top of a thin film of PHV previously cast by pipetting 2.5 mL of 1% PHV in CHCl\(_3\) evenly onto a (7 × 9 mm) glass plate and allowing the CHCl\(_3\) to evaporate. The gel-zymogram assembly was then incubated at 37°C until bands of clearing appeared, indicating PHA depolymerase activity.

**Other assays and determinations**

Protein concentrations were routinely estimated using the dye-binding assay [31] with bovine serum albumin (BSA) as the standard. For experiments on glucose repression of PHA depolymerase synthesis in broth, glucose concentrations were measured using the dinitrosalicylate assay for reducing sugars [32], with glucose as the standard. Extractable cell protein was determined by pelleting cells from 2.0 mL of culture, washing with deionized water, resuspension in 2.0 mL of 1.0 N NaOH, and heating at 100°C for 10 min. The extracts were clarified by centrifugation, and the protein concentrations in the supernatants were determined using a Bicinchoninic Acid Protein Assay Kit (Sigma-Aldrich) with BSA as the standard. This assay was directly compatible with the NaOH used for extraction.

**Cloning and sequencing of the PHA depolymerase gene**

Restriction digests, agarose gel electrophoresis, culturing of E. coli, and other routine DNA manipulations were done as described [33]. Cloning of the PHA depolymerase gene (phaZ) was done using PCR-based techniques; PCR and sequencing primers used are shown in Table S1. Purified PHA depolymerase (90 pmol) was concentrated by trichloroacetic acid precipitation and subjected to SDS-PAGE, followed by blotting onto Mini ProBlott™ polyvinylidene difluoride membranes (Applied Biosystems, Foster City, CA), and staining with Ponceau S according to the manufacturer’s instructions. The PHA depolymerase protein band was excised, and the N-terminal sequence was similarly obtained by omitting the blotting step from the sequencing procedure, yielding the sequence: AGLTQVTGFGSNPGNLTMHTYVPDGLAAG.
Degenerate PCR primers PHAD-8 (sense) and PHADIN-3 (antisense) were designed using reverse translations of the amino acid sequences from the N-terminus and internal fragment, respectively. Degeneracy in the sense primer was minimized by using G or C in the Wobble positions of the codons, since the G+C content of *Streptomyces* species is known to be about 70% [9]. For the antisense primer, a defined base composition of 53.2% G, 38.1% C, 4.6% T, and 4.1% A was incorporated into the Wobble positions of the amino acids, based on a codon usage table for *Streptomyces coelicolor* available at: http://www.kazusa.or.jp/codon/S.html. These primers were used to amplify a 500 base pair (bp) segment (A20TG) of the PHA depolymerase gene. The PCR reaction contained an Illustra PuReTaq™Ready-To-Go™ PCR bead (GE Healthcare, Piscataway, NJ, USA), 118 ng of genomic DNA template, 25 pmol of each primer, and 5% v/v dimethylsulfoxide in a final volume of 25 μL. The PCR schedule included 30 cycles of denaturation (94°C, 1 min), annealing (62.5°C, 30 sec), and extension (72°C, 1 min), followed by an additional extension at 72°C for 10 min.

A combination of PCR-based gene walking techniques was used to obtain additional sequence within and outside of *phaZ*. These included inverse PCR [34]; single-stranded (ss) DNA ligation PCR [35]; and the method of Siebert et al. [36], using the GenomeWalker™Universal Kit (Clontech, Mountain View, CA, USA). All procedures were performed essentially as described.

PCR products were purified by agarose gel electrophoresis and extracted from gels using a GenElute™ Gel Extraction Kit (Sigma Aldrich) according to the manufacturer’s instructions. In most cases, the products were first ligated into a pCR®2.1 vector or pCR®4-Blunt-TOPO vector (Life Technologies) and introduced into One Shot® TOP10 E. coli cells. Plasmid DNA from positive *E. coli* clones was purified using a GenElute™ Plasmid Miniprep Kit (Sigma-Aldrich), and sequenced with M13 Reverse primer (pCR®2.1 clones) or T3 and T7 primers (pCR®4-Blunt-TOPO clones) by Eurofins MWG Operon, Louisville, KY, USA. PCR products obtained with the GenomeWalker™ Universal Kit were sequenced directly with primers GSP2 and AP2.

As a final check, the entire *phaZ* coding sequence was amplified from genomic DNA by PCR with primers PHAZF9 and PHAZR9, which contained engineered BamHI and HindIII restriction sites, respectively. The PCR product was digested with BamHI and HindIII and ligated with similarly digested plasmid pUC18 [33]. The resulting construct was cloned into *E. coli* DH5α, and plasmid DNA extracted from positive transformants was sequenced with M13 forward and reverse primers plus internal, *phaZ*-specific primers (PHAZL3, PHAZINF, PHAZINR, and PHAZIR4). The nucleotide sequence of *phaZ* was deposited in GenBank, with the accession number MH746934.

**Sequence analysis**

Sequences from *phaZ* were assembled with the CAP3 Sequence Assembly Program [25], and open reading frames identified with the ExpASy Translate utility (https://web.expasy.org/translate/). Initial homology searches of the predicted amino acid sequence were done with the Protein BLAST tool (https://blast.ncbi.nlm.nih.gov/Blast.cgi) against the ReSeq protein database. To create a multiple sequence alignment, the predicted amino acid sequence was first searched with Protein BLAST against the PHA Depolymerase Engineering Database http://www.ded.uni-stuttgart.de/ [4]. Eight sequences with the best homology scores were aligned with the *Streptomyces* sp. SFB5A deduced amino acid sequence using the T-Coffee program (http://tcoffee.vital-it.ch/apps/tcoffee/do:regular) [37]. A potential signal peptide was located using Predisi (http://www.predisi.de/) [38]. A model of the three-dimensional structure of the PHA depolymerase was constructed from its deduced amino acid sequence using PHYRE2 [39], available at: http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index.

**Results**

**Isolation and characterization of *Streptomyces* sp. SFB5A**

Isolate SFB5A formed filamentous, dry, flat, and opaque colonies after 5 d of incubation on SNC-agar plates containing PHB, glucose, or other carbon sources, and emitted the odor of geosmins characteristic of actinomycetes. A substrate mycelium was formed during the first 24 to 36 h of incubation, followed by a powdery white aerial mycelium. After 3 to 5 d, colonies became grey-brown, indicating sporulation, with color more pronounced at the periphery than in the center. Gram stains showed a mixture of Gram positive branched filaments, rounded rods approximately 0.5 × 1.0 μm, and spores in older cultures. A soluble brown pigment was formed after 3 to 5 d of growth on TSA, MSF, or SNC plus glucose agar, but not on SNC-PHB agar. Colonies on complex agar media such as TSA and NA were moister and less filamentous than on SNC-PHB agar, wrinkled, off-white, with undulate margins, and failed to exhibit sporulation even after incubation for over 3 weeks. Growth in SNC broth media was very flocculent, and cells adhered to the sides and bottom of shake flasks. In contrast, growth in complex media such as NB or TS/2 was much faster and more dispersed than in SNC broth. The growth rate and doubling time in NB at 30°C, estimated by turbidometry, were 0.75 h⁻¹ and 56 min, respectively. The temperature range for growth on TSA plates was 15-39°C, with optimal growth between 30 and 35°C.

BLAST searches of a 1324 bp portion of the 16S rRNA gene amplified from isolate SFB5A showed the highest identity to 16S rRNA from *Streptomyces cinereoruber* subsp. *cinereoruber* NBRC 12756 [Accession #AB184121; 99.62% similarity (1319 bp), 99.7% completeness]. However, preliminary results of physiological tests and morphological comparisons between the two strains suggested that they may represent separate species, and further taxonomic identification is in progress. Therefore, the isolate was designated as *Streptomyces* sp. SFB5A.

**Degradation of PHA in cultures**

*Streptomyces* sp. SFB5A grew on SNC agar plate overlays containing PHB or PHBV. Clearing zones appeared after about 30 to 48 h, during the transition from vegetative to aerial mycelium, with full clearing after 3 to 5 d (Figure 1). It also grew slowly on SNC agar plates overlaid with PHV thin films, and full clearing of the PHV required up to 7 d (Figure 1).

When NB-grown cells were resuspended in SNC-PHB broth and incubated in shake culture, PHA depolymerase activity was detected after a lag of 12 to 24 h, peaking at 36 to 48 h, and declining rapidly thereafter, possibly because of proteolytic degradation (Figure 2). Maximal activity levels were obtained with inocula containing 0.4 to 0.7 mg of dry cell weight/mL.
The latter corresponded to resuspension of pelleted NB-grown cells in a volume of SNC-PHB broth equal to that of the original culture, so this inoculum level was used for subsequent experiments on PHA depolymerase production in broth. The lag time with inocula that were washed was about 12 h longer than that for unwashed inocula, although the maximal PHA depolymerase activities observed were similar (not shown). Use of log phase cells versus stationary phase cells in a volume of SNC-PHB broth equal to that of the original culture, so washed was about 12 h longer than that for unwashed inocula, although the maximal PHA depolymerase activities observed were similar (not shown). Use of log phase cells versus stationary phase cells as inocula decreased the maximal activity observed by about 3-fold but did not affect the lag time (not shown).

Activity was also obtained in SNC broth cultures containing R,S-3HB, R-3HB or PHBV but less than with PHB (26.8%, 14.7%, and 4.9% respectively of activity with PHB). Activity from broth containing PHV or lacking any carbon source was at or below the detection limit for the assay. PHA depolymerase synthesis in broth cultures was repressed by glucose and resumed after exhaustion of glucose from the medium (Figure S1). A similar repression phenomenon was observed in agar cultures containing both PHB and 0.5% glucose, clearing zones requiring 4 to 5 d to appear instead of 2 d with PHB alone (not shown). Use of PHB plus 1% glucose increased this time to at least 7 d.

Dual zymography revealed that a PHA depolymerase active against both PHB and PHV was produced during growth in SNC-PHB broth (Figure 3). Identical activity bands of Rf 0.34 appeared on both the PHB and PHV zymograms. A faint band at Rf 0.18 also appeared on the PHV zymogram but not on the PHB zymogram.

**Figure 1:** Growth of Streptomyces sp. SFB5A on PHAs and formation of clearing zones. The organism was streaked for isolation on SNC agar plates overlaid with SNC agar containing PHB granules, PHBV granules, or PHV thin film. PHB and PHBV plates were photographed after 3 d of incubation and the PHV plate after 7 d.

**Figure 2:** Effect of inoculum size on production of PHA depolymerase in liquid cultures of *Streptomyces* sp. SFB5A. Nutrient broth-grown inocula were washed once with SNC and resuspended in 1/6 volumes of 2-fold SNC. Varying volumes of the resuspended cells were combined with 2-fold SNC, sterile water, and PHB (0.2% w/v final concentration) to give the following initial cell densities (g dry cell weight/mL): ●, 0.07; □, 0.40; ▲, 0.70; - - - ○, 1.40; - - - ●, 2.10 with PHB in culture; ●, 2.10 without PHB in culture. PHA depolymerase turbidometric assays were done on duplicate 0.1 mL samples of culture supernatant at each time point.

**Figure 3:** Dual zymography of ammonium sulfate precipitates from culture supernatant of *Streptomyces* sp. SFB5A grown in SNC broth containing 0.2% PHB for 48 h. Lanes were loaded with the amounts of PHA depolymerase activity indicated (turbidometric assay): 1, 4.6 munits; 2, 3.7 munits; 3, 1.8 munits. Abbreviations: O, origin; DF, dye front; RF, relative electrophoretic mobility.

**Purification and characterization of PHA depolymerase**

PHA depolymerase was purified 186-fold to approximately 97% electrophoretic homogeneity from PHB-grown culture supernatants of *Streptomyces* sp. 5A, with a final yield of 11 μg protein and 16% recovery of activity. Its subunit Mr was approximately 47,000 (Figure S2). It was stable during storage for 4 d at 25°C, 7 d at 4°C, at least 14 d at -20°C, and indefinitely at -80°C. It lost only 10% of its original activity after 5 cycles of freezing at -80°C and thawing at 20°C. Optimal activity was obtained from pH 7.0 to 8.5, and at 45°C.

The effects of ionic compounds were tested (Table 1). CaCl2, MgCl2, and KCl stimulated activity, probably because of the cationic moieties of common enzyme inhibitors on PHA depolymerase activity were also tested (Table 1). The enzyme was highly inhibited by the divalent ion Chelator, dithiothreitol (DTT), but only weakly by the sulphydryl alkylating reagent, iodoacetate (IAA). It was not inhibited by KCN. The non-ionic detergents Triton X-100 and Tween-20 completely inhibited activity. Activity was strongly inhibited by the divalent ion chelator, ethylenediaminetetraacetic acid (EDTA), but was restored by addition of 5 mM CaCl2 to the assay mixture, confirming the stimulatory effect of CaCl2. Superficially, the serine hydrolase inhibitor, phenylmethylsulfonyl fluoride (PMSF) inhibited activity at 1 mM, but ethanol used to prepare the PMSF stock solution also inhibited activity
so that the net inhibition was only 2.5%. The enzyme lacked lipase activity (not shown), as determined by a specific plate assay [28].

### Ionic Compounds

| Compound | Final concentration (mM) | % of Control |
|----------|--------------------------|-------------|
| Deionized water (no CaCl₂) | --- | 100 |
| CaCl₂ | 2 | 245 |
| MgCl₂ | 2 | 188 |
| ZnCl₂ | 2 | 33.3 |
| NaCl | 4 | 101 |
| Na₂CO₃ | 2 | 82.3 |
| Na₂HPO₄ | 2 | 83.3 |
| Na₂SO₄ | 2 | 45.8 |
| KCl | 4 | 117 |
| NH₄Cl | 4 | 54.2 |
| NaN₃ | 4 | <0 |

**Enzyme Inhibitors**

| Compound | Final concentration (mM) | % of Control |
|----------|--------------------------|-------------|
| Deionized water (plus CaCl₂) | --- | 100 |
| DTT | 1 | 30.4 |
| EDTA | 1 | 6.1 |
| EDTA plus 5 mM CaCl₂ | 1 | 136 |
| KCN | 1 | 96.6 |
| PMSF | 1 | 66.4 |
| Ethanol | 856 | 68.9 |
| IAA | 1 | 82.6 |
| Triton X-100 | --- | <0 |
| Tween 20 | --- | <0 |

*Table 1: Effects of selected ionic compounds and common enzyme inhibitors on activity of PhaZ<sub>SFB5A</sub>.* The standard turbidometric assay reaction mixture was supplemented with the test compounds at the concentrations indicated. For testing of ionic compounds, CaCl₂ was omitted from the standard assay mixture. For testing of common enzyme inhibitors, the standard assay mixture (i.e., containing 1.0 mM CaCl₂) was used. CaCl₂ was added to a final concentration of 5.0 mM in this assay mixture. Since the PMSF stock was made up in ethanol, a control containing an equivalent volume of ethanol (50 μL per mL of assay mixture) was done. The final concentration of ethanol was thus ~856 mM final concentration. The concentrations of Triton X-100 and Tween 20 were 0.1% v/v. Percentage of activity versus deionized water control without CaCl₂ (for ionic compounds) or with CaCl₂ (for enzyme inhibitors). Values are averages from triplicate assays. Activities were corrected for absorbance changes obtained with minus enzyme controls. Values below zero indicate that final absorbance readings were slightly above that of the minus enzyme control.

Qualitative spot tests indicated that the purified PHA depolymerase degraded suspensions of PHB, PHBV, and PHV (not shown), although formation of clearing zones on PHV was much slower and less complete than with the other two PHAs. The enzyme did not degrade aPHB suspensions as assayed turbidometrically. The products from PHA degradation were identified by ESI-TOF-MS (Table 2; see Figure S3 for representative mass spectra). The enzyme degraded PHB granules and thin films to mostly 3HB monomer, with small and variable amounts of 3HB dimer and trimer; higher order oligomers were not detected. Controls without enzyme showed no significant product formation after 4 h (not shown). The enzyme also degraded granules and thin films of PHBV to primarily the monomers of 3HB and 3HV, with smaller amounts of the dimer and trimer forms. Furthermore, the enzyme degraded PHV granules and thin films, with a similar product pattern of 3HV units. The enzyme did not significantly degrade PHO thin films; total spectral abundance of 3HO obtained was only about 0.9% of total products obtained with PHV thin films.

Time courses of PHB and PHV degradation to monomers and dimers were roughly hyperbolic (Figure 4). The enzyme degraded PHB 9.7 times faster than PHV, based on rates calculated from the formation of total products in the first 15 min, and generated 2.1 times more products from PHB versus PHV after 4 h. With PHB as substrate, production of 3HB trimers peaked at 60 min and then decreased, whereas production of 3HB dimers increased steadily and then stabilized. A similar pattern was obtained with PHV as substrate, although production of 3HV trimers peaked only after 120 min, and not as sharply as with PHB.

#### Cloning of the PHA depolymerase gene

Attempts to clone DNA directly from *Streptomyces* sp. SFB5A into *E. coli*, including preparation of partial and total gene libraries, were unsuccessful. Therefore, *phaZ* was cloned by using PCR-based methods, as illustrated in Figure S4. A 500 bp segment of *phaZ* was obtained by using PCR with degenerate, gene-specific primers designed using amino acid sequence information from purified PHA...
depolymerase protein. Cloning of the rest of the gene was accomplished with three different PCR-based gene walking techniques.

### Table 2: Pattern of products from degradation of various PHAs by PhaZ<sup>S</sup>.

| PHA Substrate<sup>a</sup> | Percent of total<sup>b</sup> | Total Abundance<sup>c</sup> |
|--------------------------|-----------------------------|-----------------------------|
|                          | 3HB Forms                  | 3HV Forms                  | 3HO |
|                          | 1-mer | 2-mer | 3-mer | 1-mer | 2-mer | 3-mer | 1-mer |         |         |
| PHB-G                    | 80.8  | 11.3  | 7.9   | 1-mer | -     | -     | -     | 9,78,272 |
| PHB-TF                   | 82.1  | 10.3  | 7.6   | 1-mer | -     | -     | -     | 10,60,839 |
| PHV-G                    | -     | -     | -     | 72.8  | 13.8  | 13.4  | -     | 9,60,429  |
| PHV-TF                   | -     | -     | -     | 77.6  | 12.9  | 9.5   | -     | 8,37,957  |
| PHBV-G                   | 49.8  | 5.7   | 3.8   | 37.3  | 2.9   | 0.6   | -     | 22,65,534 |
| PHBV-TF                  | 50.5  | 4.9   | 2.1   | 39.1  | 3.1   | 0.4   | -     | 22,23,732 |
| PHO-TF                   | -     | -     | -     | -     | -     | -     | 100   | 7588      |

<sup>a</sup>PHAs (180 µg) in granular (G) or thin film (TF) form were incubated with 100 ng purified PHA depolymerase for 4 h and processed for ESI-TOF-MS as described in Materials and Methods. <sup>b</sup>Data indicate the percentage of total spectral abundance for the forms indicated and are averages from at least two replicates for each PHA tested. Values were corrected for abundances of each form obtained with controls lacking enzyme, which were at or near the background level for the instrument (abundance level 10,000). -, no product detected. <sup>c</sup>Obtained by totaling the spectral abundances of all the products indicated for each PHA.

An open reading frame (ORF) of 1505 bp was ultimately obtained after sequence assembly, encoding a protein containing 501 amino acids and molecular weight of 51,377 Daltons. The ORF contained 71.8% G+C residues, consistent with the high G+C content of streptomycetes [9]. Analysis of the open reading frame with Predisi showed 71.8% G+C residues, consistent with the high G+C content of streptomycetes [9]. Analysis of the open reading frame with Predisi [38] revealed a putative signal peptide containing 48 amino acids. This was followed by sequence encoding a peptide of 453 amino acids, with a calculated isoelectric point of 4.85 and molecular weight of 46,400 Daltons, which corresponded well with the M<sub>r</sub> of the purified PHA depolymerase (47,000) determined by SDS-PAGE (Figure S2). In addition, the first 29 amino acids of this predicted peptide agreed exactly with those obtained from N-terminal amino acid sequencing of purified PHA depolymerase (see Materials and Methods). Initial Protein BLAST searches of the predicted amino acid sequence revealed significant homology to esterases and PHA depolymerases from Streptomyces and other actinomycetes.

**Discussion**

We describe here the isolation of Streptomyces sp. SFB5A, which degrades PHB, PHV, and PHBV in agar overlays and broth culture. PHA depolymerase activity, assayed turbidometrically with PHB as substrate, was detected in culture supernatants from growth of the organism in liquid medium containing PHB, PHBV, or 3HB, but not PHV. Interestingly, PHV was degraded slowly in agar overlays and by purified enzyme (see below), but PHA depolymerase activity was not detected in broth cultures containing PHV, even after extended incubation. Perhaps the enzyme adhered tightly to PHV in broth culture, so that it would not be detectable in culture supernatants. As observed with many other extracellular PHA depolymerases [2,5–7], synthesis of PHA depolymerase was repressed in liquid and agar media containing both glucose and PHB until glucose was exhausted. Zymography suggested that only one PHA depolymerase was produced by this isolate during growth on PHB and that the enzyme could degrade both PHB and PHV. However, we did not entirely rule out the possibility of multiple PHA depolymerases with the same or different electrophoretic mobility on native gels. The faint R<sub>t</sub> 0.18 band on the PHB zymogram (Figure 3) could thus represent a different isoform of PHA depolymerase, but more likely self-aggregates or complexes with other proteins.

A PHA depolymerase (hereofore abbreviated PhaZ<sup>S</sup>) was purified from liquid cultures of Streptomyces sp. SFB5A containing PHB. Combined results of qualitative spot tests, turbidometric assays, and mass spectrometry showed that the purified enzyme could degrade PHB, PHV, and PHBV, but not PHO or aPHB, placing it in the family of extracellular denatured PHA depolymerases [2]. It shared several characteristics with extracellular dPHA depolymerases from other species of Streptomyces (Table 3), and PHA depolymerases in general [2]. These included an alkaline pH optimum (with the exception of S. ascomycinicus), temperature optimum of 40–60°C, high stability, monomer structure of approximately 40–60 kDa, affinity for hydrophobic interaction chromatography resins, and stimulation by divalent cations. Notably, activity was strongly inhibited by NH<sub>4</sub>Cl and SO<sub>4</sub><sup>-2</sup>, which necessitated careful removal of excess (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> from pellets after the ammonium sulfate precipitation step of purification prior to assay. The strong inhibition by the sulphhydril reducing agent, DTT, but weak inhibition by the sulphhydril alkylation reagent, IAA, suggested that disulfide linkages but not free sulphhydril groups were crucial for activity. Lack of inhibition with KCN indicated that iron or other susceptible metals were not involved in catalysis. Non-ionic detergents strongly inhibited its activity, possibly by interfering with binding of the relatively hydrophobic PHB to its binding site within the enzyme. The enzyme was not significantly inhibited by the serine hydrolase inhibitor, PMSE, known to inhibit some PHA depolymerases [2].

Mass spectral analysis showed that the purified enzyme degraded PHB, PHV, and PHBV in either granule or thin film form to a mixture of the corresponding monomers, dimers, and trimers, with monomers as the predominant product (Table 2). Since higher order oligomers...
were not detected, this pattern suggested an exolytic rather than an endolytic cleavage mechanism. The pattern of products arising from degradation of PHB by extracellular dPHA<sub>SCL</sub> depolymerases is variable. Some produce only 3HB monomer [5,40], others produce various proportions of monomer and dimer [6,14,41,42], while still others produce mostly dimer with smaller amounts of monomer and/or trimer [2,43,44]. The enzyme from Comamonas acidovorans forms different proportions of degradation products from PHB depending on enzyme concentration, with mostly dimer and smaller amounts of monomer formed at low enzyme concentrations (0.2 μg/mL) and the opposite proportions at high enzyme concentrations (20 μg/mL) [6].

| Species, reference | PHA Substrates<sup>a</sup> | Subunit M<sub>r</sub> | Optimum temperature (°C) | Optimum pH | Inhibited by: | Stimulated by: | No/weak Inhibition | Products from PHB<sup>b</sup> |
|-------------------|-----------------|----------------|-----------------|-----------|----------------|----------------|----------------|-----------------|
| Streptomyces sp. SFB5A (this study) | PHB, PHV, PHBV | 47,000 | 45 | 7-8.5 | NH<sub>4</sub><sup>+</sup>, SO<sub>4</sub><sup>2-</sup>, DTT, EDTA, Tween 20, Triton X-100 | Ca<sup>2+</sup>, Mg<sup>2+</sup> | IAA, KCN, PMSF | 1, (2, 3) |
| S. exfoliatus K10 [7,41] | PHB only | 49,000 | 40 | 8.5-9 | DTT, DDSC | Ca<sup>2+</sup>, Mg<sup>2+</sup>, Mn<sup>2+</sup> | EDTA, PMSF | 1, (2) |
| S. ascomycinicus [14] | PHB, PHBV | 48,400 | 45 | 6 | EDTA | ND | ND | ND |
| S. bangladeshensis 77T-4 [15] | PHB | 40,000 | 45 | ND | ND | ND | ND | ND |
| S. lydicus MM10 [12] | PHB | 65,000 | 45 | 8 | CuSO<sub>4</sub>, FeSO<sub>4</sub>, EDTA | ND | ND | ND |
| Streptomyces MG [13] | PHB, PHBV | 41,000-43,000 | 60 | 8.5 | ND | ND | ND | ND |

Table 3: Comparison of extracellular, dPHA<sub>SCL</sub> Depolymerases from Streptomyces species<sup>ª</sup>. <sup>a</sup>Abbreviations: ND, not determined or reported; DTT, dithiothreitol; DDSC, dodecansulfonyl-chloride; EDTA, ethylenediaminetetraacetic acid; IAA, iodoacetic acid; PMSF, phenylmethanesulfonylfluoride [14] Other PHAs were not tested. All PHAs listed are dPHAs. 1, 2, and 3 represent 3HB monomer, dimer, and trimer, respectively. Lesser products are in parentheses.

However, most reports of PHA degradation by PHA depolymerases only state one-point composition of the products after several hours of incubation. In contrast, we used ESI-TOF-MS to monitor the change in product composition over time, which might better reveal information about the catalytic mechanism. Time courses of PHB and PHV degradation by PhaZ<sub>Ssp5A</sub> (Figure 4) showed that formation of the corresponding monomers and dimers was essentially hyperbolic, possibly due to substrate depletion or product inhibition. In thin film reaction mixtures, remnants of the reaction mixtures, remnants of the substrate for the enzyme. Indeed, mass spectral assays showed that the initial rate of monomer formation from PHB was considerably greater than from PHV, and the product yields from PHV at 4 h were about 2-fold lower. The slower degradation of PHV versus PHB could be because the 3HV monomer has one more carbon in its side chain than 3HB, resulting in greater steric hindrance or hydrophobic effects at the catalytic site of the enzyme.

The deduced amino acid sequence of PhaZ<sub>Ssp5A</sub> included a putative 48 amino acid signal peptide of 48 amino acids containing the sequence: RRWLT. This is similar to the twin-arginine translocation (Tat) signature found in Streptomyces coelicolor: RRXϕϕ (where ϕ is a hydrophobic amino acid); the Tat pathway has been identified as a major translocation system for secreted proteins in Streptomyces coelicolor [45].

A BLAST search of the PhaZ<sub>Ssp5A</sub> deduced amino acid sequence versus the PHA Depolymerase Engineering Database [4] revealed the highest homology to extracellular dPHA<sub>SCL</sub> depolymerases with catalytic domain type 1, family 11. The best homology was with the PHA depolymerase of Streptomyces ascomycinicus (51% identity with 98% coverage). An abbreviated alignment with eight selected matches from this group is shown (Figure 5; Figure S5 shows the complete alignment). The putative signal peptide of PhaZ<sub>Ssp5A</sub> was followed by a type 1 catalytic domain [2]. This contained a triad of strictly conserved amino acids (numbers indicate the residue number in the secreted PhaZ<sub>Ssp5A</sub>; i.e., excluding the signal peptide): an oxyanion hole histidine (H185), a serine (S166) contained within a lipase box (G-L-S-A-G), as well as an aspartate (D181). Site-directed mutagenesis reveals that these three amino acids are necessary for catalysis (reviewed in reference 2). PhaZ<sub>Ssp5A</sub> also had a second conserved histidine (H305) found in the catalytic domain of PHB depolymerases [2]. A putative type 1 substrate binding domain [2] occurred at the C-terminal end of the predicted amino acid sequence of PhaZ<sub>Ssp5A</sub>. This contained several conserved hydrophobic and other amino acids, notably several tyrosine residues. In addition, PhaZ<sub>Ssp5A</sub> also featured eight cysteine residues, six of which were located in the catalytic domain (C87, C125, C198, C208, C318, and C330) and the other two (C443 and C499) in the substrate binding domain (Figure S5). The positions of these cysteines were highly conserved among the nine enzymes aligned and are likely involved in disulfide bond formation, which would explain...
the strong inhibition of PhaZ_{SFB5A} activity by DTT. A putative linker domain with a fibronectin type III (Fn3) signature [2] was located between the catalytic domain and substrate binding domain of PhaZ_{SFB5A}. However, only PhaZ_{SFB5A} and the enzymes from *Streptomyces ascomyceticus* and *Streptomyces pristinaespiralis* contained the Fn3 signature among those compared; the six others contained threonine-rich regions [2] instead. Further research might reveal whether the type of linker domain in these enzymes influences their reaction mechanisms and composition of their PHA degradation products.

Figure 5: Abbreviated amino acid sequence alignment of PhaZ_{SFB5A} with selected members of extracellular dPHA\_CL depolymerases with catalytic domain type 1, family 11 [4]. Enzyme abbreviations and organisms (accession numbers in parentheses): PhaZ_{SFB5A}, *Streptomyces* sp. SF65A; PhaZ_{Sas}, *Streptomyces ascomyceticus* sp. nov. DSMZ 40822 (formerly known as *Streptomyces hygroscopicus* subsp. *ascomyceticus*) (AAF86381.1); PhaZ_{Spr}, *Streptomyces pristinaespiralis* ATCC 25486 (EDY63333.1); PhaZ_{Th}, *Thermobifida fusca* YY (AAZ54120.1); PhaZ_{Tsp}, *Thermobifida* sp. BCC23166 (ACF17837.1); PhaZ_{Jsp}, *Janibacter* sp. HTCC2649 (EAP98776.1); PhaZ_{Sgr}, *Streptomyces griseus* NBRC 13350 (BAG18022.1); PhaZ_{Ple} and PhaZ_{3Ple}, *Paucimonas lemoignei* PHA depolymerases PhaZ2 and PhaZ3 (AAB17150.1 and AAB48166.1 respectively). Symbols: cons, consensus; H\textsubscript{oxo} oxygen hole histidine; LB, lipase box; D and H, conserved aspartate and histidine. Fully conserved amino acids are in white type with black highlight. Conserved cysteines are boxed (only 3 of 8 are shown). Only the signal peptide region from PhaZ_{SFB5A} is shown. A full alignment of these sequences is available in the supplemental materials available online (Figure S5).

*In silico* modeling of the PhaZ_{SFB5A} tertiary structure revealed a large cleft between the catalytic and substrate-binding domains, bridged by the Fn3-like linker domain (Figure 6). The linker domain is proposed to properly juxtapose the catalytic and substrate binding domains [2]. Site-directed mutagenesis studies of the extracellular dPHA depolymerase of *Ralstonia pickettii* T1 suggest that its substrate-binding domain adsorbs to and disrupts the semi-crystalline PHB surface, solubilizing PHB chains and making them available to the catalytic domain for hydrolysis [46]. We propose that the length of the linker domain may constrain the length of the solubilized chain available to the scission site and thus influence the length of the initial degradation product. The peaking of trimer levels during PHA degradation by PhaZ_{SFB5A} (Figure 4) suggests that they were an initial product of PHA degradation, perhaps due to this constraint, as modeled in Figure 6. Once trimers were formed, they could have reentered the catalytic site by diffusion and been cleaved randomly to monomers or dimers, both of which reached a steady state (Figure 4).

Alternatively, the length of the initial product could have been determined by the monomer-binding capacity of the catalytic site. For example, Ohura et al. demonstrate that the PHB depolymerase of *Pseudomonas stutzeri* preferentially binds three 3HB units, although it recognizes only two 3HB units for hydrolysis [47].

Crystallization of wild type and mutated versions of enzymes is useful for elucidating their substrate-binding and catalytic mechanisms. Only two extracellular PHA depolymerases have been crystallized to date, both of which are considerably different from PhaZ_{SFB5A}: the dPHB depolymerase from *Penicillium* (*Talaromyces*) *funiculosus* [48,49], and the PhaZ7 enzyme from *Pseudomonas lemoignei* [50,51]. The Penicillium enzyme has a type 2 rather than a type 1 catalytic domain, is considerably smaller than PhaZ_{SFB5A} (33,000 Daltons versus 46,400), and lacks the typical linker and substrate-binding domains described above [49]. Its PHA binding site is identified as a crevice on its surface, with several hydrophilic residues and three or possibly four monomer binding subsites. The catalytic triad residues (S39, D121, and H155) are clustered at the mouth of the crevice. Crystallized muteins of PhaZ7 reveal five amino acids (Y105, Y176, Y189, Y190, and W207) located in a cleft shown to be necessary for binding of 3HB tetramer [51]. Unlike PhaZ_{SFB5A}, PhaZ7 is specific for nPHB and aPHB, rather than dPHB. However, nine tyrosine residues occurred in the substrate-binding domain of PhaZ_{SFB5A}, and were well conserved among the PHA depolymerases compared in Figure 5. By analogy, these could be involved in PHA substrate binding.

We are currently attempting to overexpress phaZ in a heterologous host and thereby obtain sufficient quantities of PhaZ_{SFB5A} for crystallization. This will allow us to better study the catalytic, linker,
and substrate domains of the enzyme and to clarify the reaction mechanism of the enzyme.

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Supplemental Information:

https://www.omicsonline.org/supplementary-files/sup-characterization-of-an-extracellular-polyhydroxyalkanoate-depolymerase-from-streptomyces-sp-sfb5a-2155-6199-1000452.pdf

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