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Discovery and Biochemical Characterization of a Novel Polyesterase for the Degradation of Synthetic Plastics †

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Abstract: Plastic waste poses an enormous environmental problem as a result of soil and ocean contamination, causing the release of microplastics that end up in humans through the food web. Enzymatic degradation of plastics has emerged as an alternative to traditional recycling processes. In the present work, we used bioinformatics tools to discover a gene coding for a putative polyester degrading enzyme (polyesterase). The gene was heterologously expressed, purified and biochemically characterized. Furthermore, its ability to degrade polyethylene terephthalate (PET) model substrates and synthetic plastics was assessed.

Keywords: polymer degradation; novel enzyme; plastics

1. Introduction

Since the beginning of the large-scale production of plastics in the 1950s, these materials have found a wide variety of applications and become essential in today’s society. While in the 1960s, the percentage of plastic waste in municipal solid waste was less than 1%, this number increased to around 10% in 2005 in countries with middle/high income [1]. The annual production of plastics worldwide was estimated in 2017 to be 348 Mt, 10% of which was produced from renewable sources. Almost half of this plastic is channelled to the packaging sector, which contributes greatly to municipal solid waste and marine litter [2]. Abandoned plastic waste poses an enormous environmental problem, contaminating the soil and oceans, killing hundreds of thousands of sea animals. The release of microplastic particles, through natural weathering phenomena, can impact the biophysical properties of soil [3].

Even though biodegradable plastics, such as polylactic acid (PLA) and polycaprolactone (PCL), have emerged as an eco-friendly solution to the plastic problem, their uncontrollable use and disposal can pose the exact same problems to the already burdened environment, since for instance, PLA takes over three decades to biodegrade in soil [4]. Unfortunately, recent studies have highlighted the failure of traditional recycling processes for plastics, as only 9% of virgin plastic produced out of 8 billion metric tons of oil was recycled [5]. Lately, however, new approaches for plastic recycling have been
proposed, incorporating microorganisms and their enzymes for the depolymerization of used plastics and synthesis of new ones.

Studies on the enzymatic degradation of polyesters have been performed for over 15 years. However, since 2016, the discussion about polyethylene terephthalate (PET)-degrading enzymes has bloomed. The turning point was the work of Yoshida et al. (2016) [6], who discovered a PET-assimilating bacterium, namely *Ideonella sakaiensis*. The enzyme responsible for the degradation of the polymer was identified and characterized as a PETase. Up until then, enzymes belonging to the family of cutinases had been studied for PET degradation and especially, cutinases from the thermophilic actinomycete bacteria of the genus *Thermobifida*. *Is*PETase shares 45–53% amino acid sequence identity with the actinomycete cutinases. Cutinases have also proven capable of degrading other polyesters (polylactic acid-PLA, polybutylene succinate-PBS, polycaprolactone-PCL) as well [7–10].

Our goal is to discover a novel polyester-degrading enzyme with the ability to hydrolyze both fossil- and bio-based polyesters. The few known PET-degrading enzymes were used as templates for the identification of homologous sequences. The search led to a protein sequence originating from an Antarctic psychrotrophic lipolytic bacterium of the genus *Moraxella*. This was cloned and expressed in *E. coli*. The recombinant enzyme was biochemically characterized and its ability to degrade polyesters was tested.

2. Methods

2.1. Cloning and Expression of MorEst Gene and Purification of Recombinant Enzyme

Gene coding for the putative polyesterase gene from *Moraxella* sp. was codon optimized for expression in *E. coli* and cloned in expression vector pET22b(+). MorEst expression in *E. coli* BL21 was induced by 0.2 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at 16 °C for 20 h, based on the method described previously [11]. After that time, *E. coli* cells were harvested by centrifugation at 4000× g for 15 min at 4 °C and resuspended in 50 mM Tris-HCl pH 8, 300 mM NaCl buffer. Cell suspension was disrupted by sonication during four 1-min cycles (8 s pulses and 5 s pause) at 40% amplitude using a 20 kHz high-intensity (400 W) ultrasonic processor (VC 400, Sonic & Materials, Newtown, CT, USA). Cell debris was removed by centrifugation at 20,000× g, 30 min, 4 °C twice and loaded onto an immobilized metal-ion (Co²⁺) affinity chromatography (IMAC) as described in [12]. The purity of isolated enzymes was checked using SDS-PAGE electrophoresis (12.5% polyacrylamide) and the protein concentration was determined by measuring the absorbance at 280 nm, based on the calculated molar extinction coefficient.

2.2. Biochemical Characterization of Recombinant MorEst

A standard activity assay for MorEst was performed using *p*-nitrophenyl butyrate (*p*NPB) as a substrate at 1 mM concentration in 0.1 M phosphate-citrate buffer at pH 6. Reactions were initiated by adding 20 μL of enzyme preparation in 230 μL of substrate and the release of *p*-nitrophenol was recorded by measuring the absorbance at 410 nm in a SpectraMax-250 microplate reader (Molecular Devices, Sunnyvale, CA, USA) equipped with SoftMaxPro software (version 1.1, Molecular Devices, Sunnyvale, CA, USA) set at 30 °C.

The effect of temperature and pH on the activity of MorEst was tested by performing the standard assay by either altering the temperature or the pH in the range of 20–50 °C and the pH 3–8. Buffer systems used were phosphate-citrate pH 3–6, sodium phosphate pH 6–8 and Tris-HCl pH 8 at a concentration of 0.1 M. The effect of temperature on the stability of MorEst was studied by incubation of the enzyme in 20 mM Tris-HCl pH 8 buffer at temperatures in the range of 30–50 °C for up to 8 h and measuring residual activity by standard assay. The effect of pH on the stability of MorEst was studied by incubating the enzyme at 4 °C in 0.2 M of different buffer systems in the range of pH 5–10 for 24 h and then measuring residual activity by standard assay. Buffer systems used were citrate-phosphate pH 5–6, sodium phosphate pH 6–8, Tris-HCl pH 8–9 and Glycine-NaOH pH 9–10.
The kinetic characteristics of MorEst were calculated on four different p-nitrophenyl fatty-acid esters with varying chain length under standard assay conditions. Esters used were: pNP-acetate (C2), pNPB (C4), pNP-octanoate (C8) and pNP-decanoate (C10).

2.3. Hydrolysis of PET Oligomer, PET and Biodegradable Polymers

MorEst ability to degrade plastics was tested in reactions containing 10 mg mL\(^{-1}\) powdered plastic (particle diameter < 500 \(\mu\)m) in 0.1 M phosphate buffer pH 7 incubated at 30 °C under shaking (1200 rpm) in an Eppendorf Thermomixer Comfort (Eppendorf, Hamburg, Germany) for 3 days. Reactions were initiated by the addition of 25 mg MorEst, while another 12 mg were supplemented 24 and 48 h later. At the end of the reactions, the residual material was removed by centrifugation, washed with ultrapure water twice, freeze-dried and its weight measured. Polymer materials used were: PET (PAPET clear, Lotte Chemical, Lazenby, UK), PLA (4043D, NatureWorks, Minnetonka, MN, USA), polyhydroxy butyrate-PHB (Biomer P226, Biomer, Schwalbach am Taunus, Germany), PCL (CAPA 6500, Ravago Chemicals, Zaventem, Belgium) and PBS (NaturePlast PBE003, NaturePlast, Ifs, France).

Reactions with PET dimer (ethylene glycol-terephthalic acid—ethylene glycol-terephthalic acid) were performed under the same conditions of temperature and pH at a substrate concentration of 1 mg mL\(^{-1}\) for 24 h. These reactions were analyzed on a SHIMADZULC-20AD HPLC equipped with a SIL-20A autosampler. The column used was a C-18 reverse-phase NUCLEOSIL®100-5 (Macherey-Nagel, Düren, Germany) and the mobile phase was 20% acetonitrile, 20% 10 mM sulfuric acid in ultrapure water at a flow rate of 0.8 mL min\(^{-1}\). Detection of terephthalic acid (TPA) and its derivatives took place with a photodiode array detector Varian ProStar at 241 nm. Quantification of TPA, mono-(2-hydroxyethyl) terephthalate (MHET) and bis(2-Hydroxyethyl) terephthalate BHET was performed by constructing calibration curves with standard concentrations in the range of 0.01–1 mM. Prior to analysis, 0.1% v/v of 6 M HCl was added to each reaction and centrifuged at 5000× g, 10 °C. Supernatants were filtered through 0.2 \(\mu\)m syringe filters and analyzed.

3. Results and Discussion

The selected amino acid sequence, namely MorEst, has been designated as a triacylglycerol lipase by UNIPROT and shares the highest similarity with IsPETase (45.2%), followed by the actinomycete cutinases (41.48–46.05%). A synthetic gene was constructed, optimized for expression in E. coli and cloned in pET22b(+) vector. The native signal peptide of the protein was not included in the gene. Instead, the pelB signal sequence of the expression vector was utilized for periplasmic localization of the recombinant protein. Expression took place in E. coli BL21 (DE3) after induction with 0.2 mM IPTG at 16 °C for 20 h. The recombinant protein with an apparent molecular mass of 33 kDa was functionally expressed in this system and purified using immobilized metal affinity chromatography (IMAC), as seen in Figure 1.

![Figure 1](image-url)  
**Figure 1.** SDS-PAGE gel of immobilized metal-ion affinity chromatography (IMAC) purification steps from the E. coli intracellular fraction after sonication. Lanes: (1) crude intracellular protein fraction; (2) IMAC flow through; (3) washing step; elution with (4) 5 mM, (5) 10 mM and (6) 20 mM imidazole; (7) protein marker (Molecular Weights shown on the right side of the figure); and (8) purified MorEst.
Regarding biochemical characterization, MorEst showed maximum activity at 30 °C, while it maintained over 80% of its maximum activity in the range of 20–35 °C (Figure 2a). Concerning pH, MorEst appeared to act optimally at pH 9, but it showed very high activity at pH > 7. However, due to the high autohydrolysis phenomena of pNPB in alkaline pH (especially pH 9), the results cannot be quantified accurately (Figure 2b).

![Figure 2](image)

**Figure 2.** Effect of temperature (a) and pH (b) on the activity of MorEst.

MorEst appeared to be a rather thermolabile enzyme when incubated in 20 mM Tris-HCl pH 8 buffer, losing over 90% of its activity after 1 h at 50 °C. At 40 °C, the half-life of the enzyme was 2 h, while it was more stable at 30 °C, retaining 70% of its initial activity after 8 h (Figure 3a). On the contrary, MorEst appeared to be rather stable at different pH conditions in the range of pH 5–10, retaining or even enhancing its activity after 24 h incubation at 4 °C. What was noticed, however, was that the higher salinity buffer systems (200 mM) provided a better environment compared to low (20 mM) salinity systems (Figure 3b).

![Figure 3](image)

**Figure 3.** Effect of temperature (a) and pH (b) on the stability of MorEst. In (a) relative activity has been calculated at 30 °C (●), 40 °C (○) and 50 °C (▼) based on enzyme activity incubated in ice, while in (b) enzyme incubated in 20 mM Tris-HCl pH 8 buffer.

Kinetic constants of MorEst were calculated on different pNP fatty-acid esters with varying chain-length to determine the specificity of the enzyme. As seen in Table 1, even though MorEst shows the highest catalytic turnover on pNP-C2, it shows the highest affinity and catalytic efficiency for pNP-C4. For carbon chain lengths over 8, the catalytic efficiency of the enzyme drops dramatically (over 20 times).
Table 1. Kinetic constants of MorEst on synthetic substrates.

| Substrate | \( k_{\text{cat}} \) (min\(^{-1}\)) | \( K_M \) (mM) | \( k_{\text{cat}}/K_M \) (min\(^{-1}\) mM\(^{-1}\)) |
|-----------|-----------------|--------------|------------------|
| pNP-C\(_2\) | 69.6 ± 7.7 | 5.4 ± 1.0 | 12.8 ± 2.7 |
| pNP-C\(_4\) | 29.2 ± 1.3 | 1.7 ± 0.2 | 17.6 ± 2.0 |
| pNP-C\(_8\) | 5.8 ± 0.7 | 7.2 ± 1.1 | 0.8 ± 0.2 |
| pNP-C\(_{10}\) | 2.7 ± 0.5 | 7.0 ± 1.8 | 0.4 ± 0.1 |

Recombinant MorEst could cleave soluble BHET, producing mostly MHET and a small amount of TPA (3% of total products concentration). It could also hydrolyze the insoluble PET-dimer, releasing 1.17 mM TPA and 0.13 mM BHET. However, the enzyme did not release any products of detectable concentration from the crystalline PET polymer tested. Regarding biodegradable polyesters, MorEst showed the highest activity on PCL, reducing its weight by 18 ± 2%.

4. Conclusions

Enzymatic degradation of polymers shows a dynamic as a sustainable solution for modern society’s plastic problem. Discovery of novel polymer-degrading enzymes is an important aspect of this solution. In this work, we reported the discovery of a novel putative bacterial polyester-degrading enzyme (polyesterase). The recombinant enzyme was biochemically characterized and its ability to degrade PCL was shown.

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