Molecular Characterization of a Novel Family VIII Esterase with β-Lactamase Activity (PsEstA) from Paenibacillus sp.

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Abstract: Molecular information about family VIII esterases, which have similarities with class C β-lactamases and penicillin-binding proteins, remains largely unknown. In this study, a novel family VIII esterase with β-lactamase activity (PsEstA) from Paenibacillus sp. was characterized using several biochemical and biophysical methods. PsEstA was effective on a broad range of substrates including tertiary butyl acetate, glyceryl tributyrate, glucose pentaacetate, olive oil, and p-nitrophenyl esters. Additionally, PsEstA hydrolyzed nitrocefin, cefotaxime, and 7-aminocephalosporanic acid. Interestingly, two forms of immobilized PsEstA (CLEAs-PsEstA and mCLEAs-PsEstA) showed high recycling property and enhanced stability, but hybrid nanoflowers (hNFs) of PsEstA require improvement. This study provides a molecular understanding of substrate specificities, catalytic regulation, and immobilization of PsEstA, which can be efficiently used in biotechnological applications.

Keywords: PsEstA; family VIII esterase; β-lactamase; antibiotics; immobilization

1. Introduction

Bacterial β-lactamases hydrolyze chemical compounds containing a β-lactam ring and are the primary cause of bacterial resistance against different classes of β-lactam antibiotics. Therefore, the production of one or more types of β-lactamase is the most effective strategy through which clinically important gram-negative bacteria effectively hydrolyze the β-lactam rings of antibiotics, such as penicillins, cephalosporins, or carbapenems [1–3]. Based on amino acid motifs, protein sequence, molecular size, and 3D structure, β-lactamases are classified into four molecular classes termed A, B, C, and D [4]. Interestingly, classes A, C, and D hydrolyze their substrates through the formation of an acyl-enzyme with a catalytic serine, whereas class B utilizes a zinc ion to facilitate a hydrolytic reaction. Among β-lactamases, class C enzymes are of high importance, because they pose significant threats for antibiotic treatments due to their occurrence in many gram-negative pathogens, such as Enterobacteriaceae and Pseudomonas spp. [5,6]. Class C β-lactamases play a role in the resistance of these pathogens to cephalosporins, cephemycins, and carbapenems, and are not inhibited by clavulanic acid [7,8]. The molecular structure of class C β-lactamases consists of two domains of different sizes. The bigger domain contains a central antiparallel beta sheet flanked by alpha helices at each side,
while the smaller domain is entirely composed of alpha helices and contains the catalytic serine residue [7,9–11].

There is an increasing need to comprehend class C β-lactamases at the molecular level to develop potent and novel inhibitors against them given that they are one of the major causes of antibiotic resistance in clinically significant bacterial pathogens. Interestingly, several esterases, collectively known as family VIII esterases, have recently been reported to have similarities with class C β-lactamases and hydrolyze nitrocefin or other β-lactam antibiotics [12–15]. These proteins have a Ser-Lys-Tyr catalytic triad, and their nucleophilic serine residue is present in the N-terminal region, instead of the G-X-S-X-G motif located in the N-terminal region, of most bacterial lipases/esterases [16,17]. A novel family VIII esterase with β-lactamase activity (PsEstA) has recently been identified in Paenibacillus sp. PBS-2 [18]. Here, we biochemically characterized PsEstA to lay the foundation for the understanding of this enzyme at the molecular level.

2. Materials and Methods

2.1. Sequence Analysis and Model Building

The primary sequences of PsEstA and other related bacterial esterases/lipases, as well as class C β-lactamases, were retrieved from the NCBI database or Swiss-Prot. The phylogenetic tree was constructed using the neighbor-joining (NJ) method of MEGA 7.0 package [19]. Multiple sequence alignments were performed using Clustal Omega [20], and the results were rendered using ESPript [21]. The PsEstA structure was modeled using the crystal structure of EstU1 from a metagenome (PDB code: 4IVK, 31% sequence identity) as a template on the SWISS-MODEL server. Molecular docking was performed using the flexible side chain method [22,23]. The coordinate of the tetrahedral intermediate was built using the PRODRG server [24]. Processing of PsEstA and the tetrahedral intermediate was performed with AutoDock Vina [25]. Graphical representations were prepared using PyMOL software [26].

2.2. Protein Purification

Cloning and expression of PsEstA were performed as previously described [18]. Transformed E. coli BL21(DE3) cells were grown in LB medium until OD600 reached 0.4–0.6. After incubation with 1 mM isopropyl-β-d-1-thiogalactoside for 4 h, the cells were harvested by centrifugation at 6000 rpm and at 4 °C for 20 min, and then resuspended in lysis buffer (20 mM Tris-HCl, 20 mM imidazole, and 1 mM EDTA; pH 7.4). Next, the cell suspension was lysed by sonication and cell debris were precipitated out by centrifugation at 20,000 rpm for 30 min. The supernatants were then loaded onto HisTrap HP columns in an AKTA Prime Plus Liquid Chromatography System (GE Healthcare, Chicago, IL, USA). After extensive washing, the recombinant PsEstA protein was eluted using an imidazole gradient, and the pooled fractions were buffer-exchanged into a storage buffer (20 mM Tris-HCl and 1 mM EDTA; pH 8.0) without cleaving the N-terminal His-tag. Protein purity and molecular weight were confirmed by SDS-PAGE. Protein concentrations were determined using a Bio-Rad Protein assay kit (Bio-Rad Laboratories, Hercules, CA, USA).

2.3. Characterization of PsEstA

Activity staining was performed using native-PAGE followed by staining of the gel with Coomassie Brilliant Blue R-250 and 4-methylumbelliferone (4-MU) acetate [27,28]. For the overlay activity assay, native-PAGE analysis was performed. Subsequently, the gel was washed three times with a storage buffer, and then incubated with 4-methylumbelliferyl (4-MU) acetate to detect the fluorescent signal under UV illumination [16,17]. Hydrolysis of 4-methylumbelliferone (4-MU) acetate or phosphate by PsEstA was also investigated in an Eppendorf tube in an ultraviolet (UV) illumination box. For gel filtration analysis, purified PsEstA was applied onto Superdex 200GL columns in AKTA UPC-900 system (GE Healthcare, Chicago, IL, USA). Matrix-assisted laser desorption time-of-flight mass spectrometry
was performed in the positive ion mode on a Voyager™ BioSpectrometry™. Urea-induced unfolding was performed following 1 h incubation of PsEstA with urea at 25 °C. The emission spectra from 300 nm to 400 nm were recorded after excitation at 295 nm. Thermal unfolding was monitored from 15 °C to 90 °C at 222 nm using circular dichroism (CD) signals with a thermostatic cell holder.

The esterase activity of PsEstA and its S58A mutant was analyzed using p-nitrophenyl (p-NP) esters and naphthyl esters. Specifically, substrates included p-nitrophenyl acetate (p-NA), p-nitrophenyl butyrate (p-NB), p-nitrophenyl hexanoate (p-NH), p-nitrophenyl octanoate (p-NO), p-nitrophenyl decanoate (p-ND), and p-nitrophenyl phosphate (p-NPP). For naphthyl esters, 1-naphthyl acetate (1-NA), 2-naphthyl acetate (2-NA), and 1-naphthyl butyrate (1-NB) were used. The effects of the chemicals (urea, NaCl, glycerol, Triton X-100, and Tween 20) on the activity of PsEstA were investigated after 1 h incubation of each chemical with PsEstA. The standard assay solution included 5 μM of p-NB in 20 mM Tris-HCl (pH 8.0) with 0.5 μg PsEstA, and the assay was run for 5 min at 25 °C. All the assays described above were performed using an Epoch 2 Microplate spectrophotometer (BioTek, Winooski, VT, USA) and the enzyme activity of PsEstA in buffer alone was defined as 100%.

2.4. pH Indicator-Based Hydrolysis Assay

For pH indicator-based colorimetric assays, PsEstA was added into a phenol red-containing substrate solution [27–29]. For these assays, 10 μg of PsEstA was added to substrate solutions at 37 °C containing 0.13 g L⁻¹ phenol red. The substrates included glyceryl lipids (glyceryl tributyrate and glyceryl trioleate), natural oils (olive oil and fish oil), tertiary alcohol esters (tertiary butyl acetate, α-terpinyl acetate, and linalyl acetate), and carbohydrate esters (glucose pentaacetate, cellulose acetate, and glucosamine acetate). A rhodamine B–olive oil (or fish oil) mixture was prepared, and the fluorescence spectra were recorded from 500 nm to 600 nm after excitation at 350 nm [30]. For enantioselectivity analysis, PsEstA was incubated with enantiomeric solutions containing 300 mM methyl-(R)-(−)-3-hydroxy-2-methylpropionate or methyl-(S)-(+)3-hydroxy-2-methylpropionate.

2.5. β-Lactamase Assay

The β-lactamase activity of PsEstA was determined using the chromogenic β-lactam substrate nitrocefin [17]. PsEstA (0.1 mg/mL) was reacted with 484 μM nitrocefin solution in 20 mM Tris-HCl pH 8.5 at 25 °C, and the color change in the reaction mixture was observed. In addition, pH indicator-based colorimetric assay was performed to investigate the activity of PsEstA on 1 mM cefotaxime (CTX) and 7-aminocephalosporanic acid (7-ACA) at 25 °C for 1 h.

2.6. Immobilization of PsEstA

For preparing crosslinked enzyme aggregates (CLEAs), 0.5 mg mL⁻¹ of PsEstA was coprecipitated by 80% (w/v) ammonium sulfate at 4 °C. Crosslinking was then performed by the dropwise addition of glutaraldehyde to the final concentration of 25 mM at room temperature. After an overnight incubation followed by centrifugation at 15,000 g for 10 min, the pellet (CLEA-PsEstA) was resuspended and repeatedly washed until no activity was observed in the supernatant. The preparation and addition of the magnetic Fe₃O₄ nanoparticles were carried out as previously described [27,31]. Nanoparticles were mixed with PsEstA solution containing ammonium sulfate and glutaraldehyde. After gentle agitation for 12 h, magnetic CLEAs of PsEstA (mCLEA-PsEstA) were precipitated by centrifugation at 15,000×g for 10 min and stored in 50 mM sodium phosphate buffer (pH 7.5) at 4 °C. Scanning electron microscope images were obtained at various magnifications (50,000–100,000×) using a Carl Zeiss SUPRA 55VP microscope. The chemical stabilities of the three forms of PsEstA (free PsEstA, CLEA-PsEstA, and mCLEA-PsEstA) were determined by measuring the residual activity toward p-nitrophenyl butyrate (p-NB) after 1 h incubation in the presence of the chemical compounds [ethanol, iso-propanol, sodium dodecyl sulfate (SDS), urea] at 25 °C. For the recycling process, the two immobilized forms of PsEstA (CLEA-PsEstA and mCLEA-PsEstA) were extensively washed and then reused in the next cycle. For preparing enzyme-inorganic hybrid nanoflower (hNF), PsEstA (0.05, 0.1,
and 0.5 mg·mL⁻¹) was added into 800 μM of the Cu²⁺ metal solutions for a total volume of 3 mL [32]. The resulting mixtures were incubated at 25 °C and centrifuged at 15,000× g for 10 min. The pellets were resuspended and transferred to 1.5 mL microtubes. In the reusability tests, hNF-PsEstA was recovered by centrifugation and extensively washed before being reused in the next cycle. Then, fresh substrate of 1 mM p-nitrophenyl butyrate (p-NB) was reacted for another cycle and activity was measured. In these experiments, the enzyme activity of free PsEstA was defined as 100%.

3. Results and Discussion

3.1. Bioinformatic Analysis

For phylogenetic tree analysis of PsEstA, the 26 representative bacterial lipases/esterases sequences were analyzed. As shown in Figure 1, PsEstA belongs to family VIII esterases, which include EstC [12], Est-Y29 [33], EstB [34], and EstU1 [35].

![Figure 1. A phylogenetic tree analysis of PsEstA. The phylogenetic tree was constructed with MEGA v 7.0 using neighbor-joining method with 3000 iterations.](image)

As shown in Figure 2A, three motifs were found to be highly conserved among class C β-lactamases based on multiple sequence alignment. These three motifs (motif I, II, and III) have also been identified in family VIII esterases. In motif I, a nucleophilic Ser⁵⁸ is located in the characteristic S⁵⁸-x-x-K⁶¹ tetrapeptide, and the alkoxide ion formation at Ser⁵⁸ is believed to be enhanced by Lys⁶¹. The Tyr¹⁷² residue in motif II is involved in the substrate recognition and specificity regulation. Although motif III of class C β-lactamases is composed of the highly conserved K-T-G sequence, a W-x-G motif is observed instead in PsEstA and other family VIII esterases. The third residue of motif III has to be glycine because otherwise, other amino acids would be sterically unfavorable for the binding of the substrate. In PsEstA, the W-S-G sequence was observed in motif III. In sequence analysis, PsEstA was
found to contain a higher percentage of acidic (Asp and Glu, 12.0%) than basic (Lys and Arg, 9.3%) amino acids.

| Class C β-lactamase | Sequence |
|---------------------|----------|
| O24824 | ELGSVSTFT |
| Q1HPG6  | ELGSVSTFN |
| Q93CA2 | ELGSVSTFT |
| Q48435 | ELGSVSTLT |
| Q51578 | EIGSVSTLT |

| Family VIII esterase | Sequence |
|----------------------|----------|
| PsEstA | NLYSMTEMTT |
| Est-Y29 | RITYSMKPVV |
| EstB | RLYSMKPVV |
| EstC | RLYSMKPVV |
| EstU1 | RLYSMKPTT |

**Figure 2.** Multiple sequence alignment and structural representation of PsEstA. (A) Multiple sequence alignment was performed among class C β-lactamases and family VIII esterases. Three characteristic motifs (motif I, II, and III) with important catalytic residues are shown. Highly conserved catalytic residues are shown in red triangles. (B) A molecular model of PsEstA is shown in ribbon diagram, and α-helices and β-strands of PsEstA are represented in light green and light red, respectively. Catalytic residues and putative substrate binding pocket residues are shown as a ball-and-stick model. (C) (left) Modeling of the complex formation of p-nitrophenyl butyrate (pNB, cyan) in the pocket of PsEstA. The amino acid residues interacting with pNB are shown as sticks (green). (right) LigPlot analysis between p-nitrophenyl butyrate and PsEstA.
The structural model of \( \text{PsEstA} \) consisted of a large \( \alpha/\beta \) domain and a small \( \alpha \)-helical domain, and these features are frequently observed in other \( \beta \)-lactamases and family VIII esterases [14,17,34,35]. The nucleophilic serine was found to be located near the surface of the tunnel formed by two domains (Figure 2B). The putative catalytic triad made of Ser\(^{58} \), Lys\(^{61} \), and Tyr\(^{169} \) is positioned within a catalytic pocket close to the surface (Figure 2B, rectangular region). The substrate-binding pocket is mainly delineated by the aromatic amino acids Tyr\(^{127} \), Tyr\(^{169} \), and Trp\(^{333} \), presumably regulating the entrance of substrates via hydrophobic interactions. These residues are also highly conserved in most family VIII esterases. In molecular docking analysis, Tyr\(^{169} \) and Trp\(^{333} \) were shown to stabilize the \( p \)-nitrophenol ring, while side chains of Asp\(^{302} \) and Ser\(^{334} \) formed hydrogen bonds with the nitro moiety (-NO\(_2\)) of \( p \)-nitrophenyl butyrate (Figure 2C). In addition, backbone nitrogen of Ala\(^{336} \) is involved in the formation of an oxyanion hole. This model analysis suggested a number of active-site residues seem to be involved in hydrogen bonding networks among other conserved residues or substrate [34]. In addition to conservation of primary sequences, spatial orientation of catalytic residues between class C \( \beta \)-lactamases and family VIII esterases are very similar. As shown in Figure 3, catalytic residues of \( \text{PsEstA} \) have comparable conformation to those of AmpC [36] and blaMOX-1 [37], as well as EstU1 and Est-Y29.

![Figure 3. Structural comparison of the catalytic amino acids between class C \( \beta \)-lactamases and family VIII esterases. The key residues in three motifs are shown in sticks. Two class C \( \beta \)-lactamases [AmpC (PDB code: 1S6R) and blaMOX-1 (PDB code: 3W8K)] and two family VIII esterases [EstU1 (PDB code: 4IVI) and Est-Y29 (PDB code: 4P6B)] are colored in cyan, magenta, yellow, and pink, respectively. The structure superimposition was performed using PyMOL software.](image-url)

### 3.2. Characterization of \( \text{PsEstA} \)

The recombinant \( \text{PsEstA} \) was highly purified using an immobilized metal-affinity column, and a single band was observed (Figure 4A). The molecular mass of \( \text{PsEstA} \) was estimated to be approximately 45 kDa by SDS-PAGE, and this value is similar to those of Est-Y29 [33] and EstB [34], but smaller than EstU1 [35]. An overlay activity assay showed high fluorescence at the position where purified \( \text{PsEstA} \) was located in native PAGE (Figure 4B). Furthermore, strong fluorescence intensity was observed for 4-MU acetate and \( \text{PsEstA} \), but not for 4-MU phosphate and \( \text{PsEstA} \) (Figure 4C,D). In native PAGE, the molecular mass of active \( \text{PsEstA} \) was found to be substantially greater than 200 kDa, implying that \( \text{PsEstA} \) is in an oligomeric conformation. A similar behavior indicating an oligomeric state of \( \text{PsEstA} \) was also observed in gel filtration chromatography (Figure 4E), which has been reported for other family VIII esterases [38–40]. The mass spectrometric analysis showed a major peak \((m/z)\) at 47.7 kDa with all additional amino acids including the His-tag (Figure 4F).
3.2. Characterization of PsEstA

The recombinant PsEstA was highly purified using an immobilized metal-affinity column, and its enzymatic activity was measured. (A) Purified PsEstA was analyzed by SDS-PAGE. (B) 4-MU overlay assay of PsEstA. CBB staining (left) and fluorescence (right). (C) Hydrolysis of 4-methylumbelliferyl (4-MU) acetate and -phosphate. PsEstA only (I), 4-MU acetate with PsEstA (II), and 4-MU phosphate with PsEstA (III). (D) Fluorescence intensity of PsEstA only, 4-MU acetate with PsEstA, and 4-MU phosphate with PsEstA were measured. (E) Gel filtration analysis of PsEstA. The column was calibrated with β-amylase (200 kDa), BSA (66 kDa), peroxidase (44 kDa), carbonic anhydrase (30 kDa), and cytochrome C (12 kDa). (F) Mass analysis of PsEstA. The [M + H]+ ion peak was observed at m/z value of 47,738.

3.3. Biochemical Assay

The hydrolytic activity of PsEstA was analyzed using p-nitrophenyl (p-NP) esters with acyl chain lengths of different length [27–29,41]. As shown in Figure 5A, PsEstA had a strong substrate preference for p-NA or p-NB. However, almost no enzyme activity was observed for the long-chain substrates p-NO and p-ND. Similarly, other members of family VIII esterases such as Est22 [13], EstU1 [42], or EstM-N1 [43] showed substrate preference for p-NB. When naphthyl esters were used as the substrates, the highest enzymatic activities were observed with 1-naphthyl acetate (1-NA), followed by 1-naphthyl butyrate (1-NB) and 2-naphthyl acetate (2-NA). However, mutation of Ser 58 residue abolished most of the hydrolytic activity (Figure 5B). As shown in Figure 5B, PsEstA exhibited approximately 75% activity on 2-NA. In contrast, it showed almost no activity on 1-naphthyl phosphate (1-NP). PsEstA was also shown to be highly stable in the presence of NaCl and glycerol. Specifically, it was shown to retain approximately 100% and 70% of its enzymatic activity in the presence of 3.0 M and 4.0 M NaCl, respectively (Figure 5C). In addition, PsEstA exhibited strong tolerance to glycerol with almost no loss of enzymatic activity after incubating with up to 30% glycerol.

The chemical stability of PsEstA was investigated by monitoring the intrinsic fluorescence spectra. In its native form, PsEstA exhibited a λmax at 334 nm, implying that all the tryptophan residues of PsEstA were located in the hydrophobic interior. However, a red shift of λmax to 344 nm was observed with a remarkable increase in fluorescence intensity in the presence of 5 M urea, suggesting that the tryptophan residues were mostly exposed to the solvent (Figure 5D). The chemical stability of PsEstA in urea was also investigated by measuring the enzymatic activity in the presence of increasing concentrations of urea. At 4.0 M, ~50% of the initial activity was retained (Figure 5E). Next, the thermal...
stability of PsEstA was investigated by monitoring its thermal denaturation from 15 to 90 °C using far-UV CD at 222 nm. PsEstA showed only minor changes up to 50 °C, and its melting temperature was determined as 63 °C (Figure 5F).

Figure 5. Biochemical properties of PsEstA. (A) Substrate specificity of PsEstA toward using p-nitrophenyl esters (acetate: p-NA; butyrate: p-NB; octanoate: p-NO; decanoate: p-NDe; phosphate: p-Np). The hydrolase activities are shown relative to the activity toward p-NB. (B) Regioselectivity of PsEstA toward naphthyl esters. The hydrolase activities are shown relative to the activity toward 1-NA. Activities of wild-type PsEstA (black) and S58A (gray) are shown for comparison. (C) Activity of PsEstA toward p-NB in the presence of NaCl (left) and glycerol (right). (D) Intrinsic fluorescence spectra of PsEstA with or without 5 M urea. (E) Effects of urea on the activity of PsEstA. (F) Thermal unfolding of PsEstA was monitored by circular dichroism at 222 nm from 10 to 90 °C. The standard assay solution included 1 mM p-nitrophenyl butyrate (p-NB) as a substrate in 20 mM Tris-HCl (pH 8.0).

3.4. Substrate Analysis

The hydrolytic properties of PsEstA for acetylated carbohydrates, tertiary alcohol esters, and lipids were evaluated using a colorimetric assay [28,29,41]. The tertiary alcohol esters tert-butyl acetate, α-terpinyl acetate, and linalyl acetate were used. As shown in Figure 6A, PsEstA could effectively hydrolyze tert-butyl acetate, but not linalyl acetate or α-terpinyl acetate. Additionally, a significant hydrolytic activity of PsEstA was detected only for glyceryl tributyrate based on the yellow color of the solution (Figure 6B). Furthermore, PsEstA displayed a high activity for glucose pentaacetate, although no significant activity was detected for cellulose acetate or glucosamine acetate (Figure 6C). In addition, a high level of enzymatic activity of PsEstA was observed toward olive oil, judging from the fluorescence spectra (Figure 6D,E). For enantioselectivity analysis, (R)- and (S)-methyl-β-hydroxyisobutyrate were used [27,32]. Following incubation with PsEstA, only the reaction mixture containing the (S)-enantiomer turned yellow, showing the (S)-selectivity of PsEstA (Figure 6F). This indiscriminate nature of PsEstA could be useful for industrial applications as a biocatalyst.
3.4. Substrate Analysis

The hydrolytic properties of PsEstA for (A) acetylated carbohydrates (GPA: glucose pentaacetate; CA: cellulose acetate; NAG: N-acetyl-glucosamine), (B) triacyl glycerols (GTB:glyceryl tributyrate; GTO:glyceryl trioleate), (C) tertiary alcohol esters (tert-buty acetate, α-terpinyl acetate, and linalyl acetate). Hydrolysis of olive oil (D) and fish oil (E) was investigated using fluorescence spectra. The excitation wavelength was set at 350 nm, and emission was recorded from 500 to 600 nm. (F) Enantioselectivity of PsEstA with methyl-(R)-(−)-3-hydroxy-2-methylpropionate or methyl-(S)-(−)-3-hydroxy-2-methylpropionate.

3.5. β-Lactamase Activity of PsEstA

To investigate the β-lactamase activity of PsEstA, a chromogenic β-lactamase substrate (nitrocefin) was used as a substrate [12,17]. As shown in Figure 7, PsEstA apparently showed β-lactamase activity toward nitrocefin. The β-lactamase activity of PsEstA was further investigated using 7-ACA and CTX. PsEstA showed significant catalytic activities toward cefotaxime (CTX) and 7-aminocephalosporanic acid (7-ACA) in a pH shift assay (Figure 7E,F). In accordance with PsEstA, Est22 and EstU1 were shown to possess a noteworthy β-lactam hydrolytic activity [13,35]. In contrast, EstC showed a significant hydrolyzing activity for nitrocefin, but none of these esterases displayed any activity for other β-lactam substrates [12].

Figure 6. Substrate analysis of PsEstA. The pH shift assay was carried out toward (A) acetylated carbohydrates (GPA: glucose pentaacetate; CA: cellulose acetate; NAG: N-acetyl-glucosamine), (B) triacyl glycerols (GTB:glyceryl tributyrate; GTO:glyceryl trioleate), (C) tertiary alcohol esters (tert-buty acetate, α-terpinyl acetate, and linalyl acetate). Hydrolysis of olive oil (D) and fish oil (E) was investigated using fluorescence spectra. The excitation wavelength was set at 350 nm, and emission was recorded from 500 to 600 nm. (F) Enantioselectivity of PsEstA with methyl-(R)-(−)-3-hydroxy-2-methylpropionate or methyl-(S)-(−)-3-hydroxy-2-methylpropionate.
3.6. Immobilization of PsEstA

Enzyme immobilization has been explored in a large variety of industrial applications [44,45]. Crosslinked enzyme aggregates (CLEAs) constitute one of the highly studied methods to generate highly efficient immobilized biocatalysts [46]. Toward this end, CLEAs-PsEstA was prepared by precipitating PsEstA with ammonium sulfate and glutaraldehyde. The SEM images of CLEAs-PsEstA showed formation of globular structures with a diameter of 1–2 nm (Figure 8A). The stabilities of both free PsEstA and CLEAs-PsEstA in the presence of several chemical compounds were compared (Figure 8B). For 5 M urea and 0.1% (v/v) SDS, there were no significant differences between these two forms. However, for alcohols such as ethanol and iso-propanol, CLEAs-PsEstA was highly active compared with the free form of the enzyme. Specifically, with 30% ethanol, free PsEstA retained only ~3% of its original activity, whereas CLEAs-PsEstA showed ~35% of the original activity. The operational stability of CLEAs-PsEstA was studied up to 20 cycles. As shown in Figure 8C, CLEAs-PsEstA was highly stable for 20 cycles, retaining at the end ~75% of the original activity.

Furthermore, PsEstA was immobilized as CLEAs on magnetite nanoparticles for an efficient separation during industrial applications [27,32]. To obtain magnetic CLEA form of PsEstA (mCLEA-PsEstA), PsEstA was coaggregated with nanoparticles, and then chemically crosslinked using glutaraldehyde. Transmission electron microscopy showed that the magnetic nanoparticles had a diameter of 5–8 nm (Figure 8D). The effects of the chemical compounds on the activities of mCLEA-PsEstA and free PsEstA were investigated by measuring the residual activities in the presence of each chemical. As shown in Figure 8E, the enzymatic activities of mCLEA-PsEstA were largely remarkably higher than those of the free form of PsEstA in the presence of the alcohols tested. Specifically, free PsEstA showed a relative activity of 21% in the presence of 10% EtOH, whereas mCLEA-PsEstA retained 47% of the initial activity. Moreover, although the free PsEstA retained less
than 5% of its original activity in the presence of 30% EtOH, mCLEA-PsEstA showed 62% activity at the end. However, even 0.1% (v/v) SDS was sufficient to inactivate both free PsEstA and mCLEA-PsEstA almost completely. Similar behavior was also observed in the presence of 5 M urea. In reusability analysis, mCLEA-PsEstA retained ~75% of its original activity after the 4th cycle, but lost a substantial level of activity at the 5th cycle (Figure 8F). Taken together, immobilization of PsEstA (CLEAs-PsEstA and mCLEAs-PsEstA) was effectively performed, and these immobilized forms could function more effectively than the free form of PsEstA.

Figure 8. Characterization of immobilized PsEstA. (A) Field emission scanning electron microscopic (FE-SEM) images of crosslinked enzyme aggregates (CLEAs). (B) Chemical stabilities of free PsEstA and CLEAs-PsEstA were compared to ethanol, iso-propanol, SDS, and 5 M urea. (C) Reusability of CLEAs-PsEstA was investigated for 20 cycles. (D) Transmission electron microscopic images of magnetic nanoparticles. (E) Chemical stabilities of free PsEstA and mCLEA-PsEstA. (F) Reusability of mCLEA-PsEstA was investigated for 20 cycles. In these experiments, the enzyme activity of PsEstA in buffer alone was defined as 100%. The standard assay solution included 1 mM p-nitrophenyl butyrate (p-NB) as a substrate in 20 mM Tris-HCl (pH 8.0).

3.7. Formation of Organic–Inorganic Hybrid PsEstA Nanoflower

Recently, syntheses of organic–inorganic hybrid nanoflowers (hNFs) with greatly enhanced catalytic activities and stabilities have been reported [47,48]. Herein, the formation and catalytic activity of hybrid nanoflowers containing PsEstA were pursued. Scanning electron microscopy showed that the hybrid nanoflowers of PsEstA were hierarchical peony-like structures assembled from interlaced nanoplates (Figure 9A). Reusability tests were also carried out for 10 cycles showing that the residual activity was ~25% by the 3rd cycle (Figure 9B).
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