Nature versus nurture in two highly enantioselective esterases from Bacillus cereus and Thermoanaerobacter tengcongensis

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Summary

There is an increasing need for the use of biocatalysis to obtain enantiopure compounds as chiral building blocks for drug synthesis such as antibiotics. The principal findings of this study are: (i) the complete sequenced genomes of Bacillus cereus ATCC 14579 and Thermoanaerobacter tengcongensis MB4 contain a hitherto undescribed enantioselective and alkali-liphilic esterase (BcEST and TtEST respectively) that is specific for the production of (R)-2-benzyloxy-propionic acid ethyl ester, a key intermediate in the synthesis of levofloxacin, a potent antibiotic; and (ii) directed evolution targeted for increased thermostability of BcEST produced two improved variants, but in either case the 3–5°C increase in the apparent melting temperature (Tm) of the mutants over the native BcEST that has a Tm of 50°C was outperformed by TtEST, a naturally occurring homologue with a Tm of 65°C. Protein modelling of BcEST mapped the S148C and K272R mutations at protein surface and the I88T and Q110L mutations at more buried locations. This work expands the repertoire of characterized members of the α/β-fold hydrolase superfamily. Further, it shows that genome mining is an economical option for new biocatalyst discovery and we provide a rare example of a naturally occurring thermostable biocatalyst that outperforms experimentally evolved homologues that carry out the same hydrolysis.

Introduction

Levofloxacine (LFC) is the biologically active levo- or S-isomer of ofloxacin, a tricyclic fluoroquinolone, a potent antibiotic against Gram-negative and Gram-positive bacteria (Hayakawa et al., 1986). The mechanism of action of LFC is through targeting the essential bacterial type II topoisomerases, DNA gyrase and topoisomerase IV that are involved in DNA replication and metabolism. As a third-generation fluoroquinolone antibiotic, LFC has a broader spectrum of antibacterial activity than the second-generation ciprofloxacin and related antibiotics (Fu et al., 1992; Ernst et al., 1997). The clinical use of LFC includes long-term treatment of multidrug-resistant tuberculosis, community-acquired pneumonia, acute chronic bronchitis and acute bacterial sinusitis (Mehlhorn and Brown, 2007; Moadebi et al., 2007). Due to increasing resistance of staphylococci for example, new LFC-containing hybrid chemicals have been synthesized and shown to be potent against these microorganisms (Foroumadil et al., 2006).

There are at least five synthetic routes to LFC that involve numerous chemical steps and solvents (Kang et al., 1996; 1997; Miyadera and Imura, 1999; Kleemann and Engel, 2001). One of the key chemical intermediates is (R)-2-benzyloxy-propionic acid ethyl ester or O-benzyl lactic acid ethyl ester (BnLAE, Fig. 1). Chemical synthesis of BnLAE would produce a racemic mixture and therefore additional steps for the separation of the isomers are required. The availability of a biocatalyst, an enantioselective esterase capable of resolving the racemic esters, would provide a green and chemoenzymatic route towards the production of LFC.

In this article, we describe the identification, cloning and expression in Escherichia coli of an S-specific esterase (BcEST) from Bacillus cereus ATCC 14579 that is capable of racemic resolution of BnLAE. Furthermore, we have used directed evolution to attempt improvement the thermostability of this biocatalyst. In addition, a predicted open reading frame (ORF) from a thermophilic microorganism, Thermoanaerobacter tengcongensis, with a presumed hydrolase or acyltransferase function (TTE0556; GenBank NC_003869) was cloned and characterized. Biochemical properties of these biocatalysts are compared and discussed in the context of enhanced
thermostability and of a naturally derived biocatalyst versus genetic variants nurtured in the laboratory.

Results

Preliminary microbial screening

Four bacterial strains (Pseudomonas sp. IFO 3925; B. cereus DSC 0007, Microbacterium laevaniformans IFO 14471 and B. cereus ATCC 14579), out of approximately 500 microorganisms screened for the enantioselective hydrolysis reaction using rac-BnLAE as substrate, were found to have excellent enantiomeric ratio, $E$, in excess of 100 (not shown). $E$ is the ratio of one enantiomer in a mixture to that of the other (Chen et al., 1982). An initial attempt to purify the desired esterase from Bacillus sp. IFO 3925 proved difficult and therefore attention was turned to strain ATCC 14579 which catalysed the same reaction with an $E$ value > 200.

Cloning and identification of the BcEST-encoding gene

A 10.5 kb recombinant plasmid, designated pGEMHindIll2, was propagated in E. coli DH5α as a result of cloning a 7.3 kb HindIII genomic fragment of B. cereus ATCC 14579 in plasmid pGEM3zf (see Experimental procedures). Using synthetic primers, the cloned insert was sequenced completely on both strands. This sequence, which encompasses at least eight ORFs, has been deposited in the NCBI GenBank under Accession No. GQ243224. Identification of the coding sequence of BcEST was aided by determining peptide sequences by Edman degradation of the purified enzyme. These sequences consist of the N-terminal 17 amino acids, and three internal peptide sequences following protease digestion, one of 7 aa and two of 12 aa. In the overall BcEST sequence of 300 aa, the internal peptide sequences were located at positions 139–145, 152–163 and 176–187. Except for one discrepancy (second D in the peptide 139DTDGQPI should be an N), all other peptide sequences are in complete agreement with the DNA-derived data.

The gene context of BcEST is such that it is flanked by a potential repressor of the MarR-type (Grkovic et al., 2002; Schumacher and Brennan, 2002) that is divergently transcribed, and downstream where two hypothetical proteins, one being a putative transport protein, are found. This gene organization is in keeping with the annotation in the B. cereus ATCC 14579 genome database (Ivanova et al., 2003).

Plasmid pBcEST was constructed as a subclone of pGEMHindIll2 containing only the est gene in the same orientation as the lac promoter of expression vector pSD80. Cell cultures of E. coli BL21(pBcEST) and DH5α(pGEMHindIll2) were assayed for their enantioselective esterase activity. Both clones produced a pH change causing bromothymol blue to turn yellow when using the substrate (S)-BnLAE but not the (R)-substrate. The control cells with only the plasmid vector, IPTG-induced or non-induced, did not impart any colour change. The specific activities of the crude enzyme extract prepared from DH5α(pGEMHindIll2), BL21(pBcEST) and the native B. cereus ATCC 14579 were shown to be 0.22, 0.60 and 0.015 U mg$^{-1}$ protein respectively. This indicated a 15-
to 40-fold improvement of activity in the recombinant strains. It is noted that the est gene in pGEMHindIII2 is in the reverse orientation relative to the lac promoter, suggesting that the Bacillus promoter is functional in E. coli.

**Characteristics of BcEST and identification of TtEST**

The calculated Mₐ of BcEST of 34,321 is in agreement with that estimated by SDS-PAGE of an overproduced 32.5 kDa protein prepared from a crude extract of E. coli BL21(pBcEST) (Fig. S1). Gel filtration indicated an apparent Mₐ of 40 kDa implying a monomeric enzyme (data not shown). The predicted isoelectric point (pI) is 4.8. The NCBI Conserved Domain Search predicted an α/β-fold hydrolase enzyme of the esterase_lipase superfamily, with the characteristic catalytic triad consisting of S102-D251-H279.

A phylogenetic analysis of BcEST is presented in Fig. S2. As expected, the sequence clusters with a number of predicted hydrolases of the α/β-fold family, invariably annotated as 3-oxoadipate enol-hydrolase, that originate from a number of Bacillus spp.; the two closest homologues are from B. cereus (Fig. 2). The structural homologues of BcEST include: an aryl esterase from Pseudomonas fluorescens (PDB 1VA4, 27% identity; Cheeseman et al., 2004); a C–C bond hydrolase (MphC) of the phenylpropionate degradation pathway of E. coli (PDB 1U2E, 25% identity; Dunn et al., 2005); and, a bromoperoxidase A1 from Streptomyces aureofaciens (PDB 1A8Q, 26% identity; Hofmann et al., 1998). The sequence alignment of BcEST, together with TtEST and the three structurally similar enzymes, is given in Fig. 2.

Characteristically conserved sequences around the predicted catalytic triad include residues S102 and D251 that are found within the tetrapeptide SMGG and a motif L-X₁-G-X₂-D-X₃-V. The sequence around H279 is less cons-

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served, although BcEST and TIEST share a common (G/A)HS(P/V)FID sequence. In the classification of micro-
bial lipolytic enzymes and carboxylesterases, BcEST and
TIEST appear to be members of the Group V family that
consists of mostly non-lipolytic enzymes (Arpigny and
Jaeger, 1999; Bornscheuer, 2002).

**Evolving the thermostability of BcEST**

A preliminary stability assay with crude BcEST enzyme
preparation indicated that it is unstable above 40°C as
expected of a mesophilic enzyme. We used a thymolblue-
based colorimetric assay to carry out error-prone PCR
(epPCR) with the aim of generating thermostable mutants
by screening at two elevated temperatures, first at 50°C
and then at 55°C. In the first round, more than 4600
transformants were screened, and the best clone, desig-
nated EH5, was subjected to a second round of mutagen-
esis. The result of screening >2300 transformants in the
second round culminated in the isolation of 2IC12.

DNA sequencing identified three single-base changes
in EH5 that resulted in two amino acid substitutions: S148C due to AGT to TGT; and K272R due to AAG to
AGG; the ATT to ATC change at 148 is silent. The 2IC12
mutant was the result of two additional single-amino-acid
substitutions: I88T (ATT to ACT) and Q110L (CAA to CTA),
in addition to S148C and K272R derived from the first
round of mutagenesis.

Stability assays for BcEST and the two variants were
carried out at 55°C. After a 20 min treatment of the
respective crude enzyme, 2IC12 and EH5 retained 80%
and 40% of their original activity compared with 25% in
the case of the wild-type BcEST. A 5 min treatment at
60°C resulted in almost total loss of wild-type activity,
whereas 2IC12 retained about 50% and EH5 25% of their
activity (Fig. S3).

**Properties of purified BcEST, EH5, 2IC12 and TIEST**

The four specified esterases were purified to electro-
phoretic homogeneity (Fig. 3) using a three-step chro-
matographic procedure as described in Experimental
procedures. By SDS-PAGE (12%) gel, the proteins
appear as single bands with estimated Mr between 35
and 38 kDa, in good agreement with the calculated Mr(s).
The 2IC12 and TIEST proteins showed a slightly higher
Mr than expected (34 336 and 34 388 respectively). The puri-
fied esterases eluted at 36–40 kDa by gel filtration on
Superose-12 indicating monomeric enzymes. Analysis by
UV-Vis showed no other absorbance peak besides that at
280 nm.

The specific activity of the purified BcEST, EH5, 2IC12
and TIEST towards rac-BnLAE, assayed at 25°C, was
found to be very similar for the former three enzymes

![Fig. 3. SDS-PAGE of purified esterases. Lanes 1–4: BcEST, EH5,
2IC12 and TIEST respectively. The following standard proteins
were used: lysozyme (14 400), trypsin inhibitor (21 500), carbonic
anhydrase (31 000), ovalbumin (45 000) and bovine serum albumin
(66 200).](image)

| Esterase | $k_{cat}$ (s$^{-1}$) | $K_m$ (mM) | $k_{cat} / K_m$ (M$^{-1}$s$^{-1}$) | Mr (kDa) | M (SDS) (kDa) | $T_m$ (°C) |
|----------|------------------|-----------|-------------------------------|---------|---------------|-------------|
| BcEST    | 18.0             | 0.22      | $8.2 \times 10^4$             | 34.319  | 34.7          | 50          |
| EH5      | 14.8             | 0.21      | $7.0 \times 10^4$             | 34.363  | 35.4          | 53          |
| 2IC12    | 18.7             | 0.13      | $1.4 \times 10^4$             | 34.336  | 37.9          | 55          |
| TIEST    | 4.9              | 0.69      | $7.1 \times 10^4$             | 34.388  | 38.2          | 65          |
Although racemic BnLAE was used in the assay, selective cleavage of the \((S)\)-enantiomer producing pure \((R)\)-enantiomer was confirmed using HPLC with a chiral column for the four esterases. The enantiomeric excess of \((R)\)-BnLAE was found to be 99% with high yield (close to 50%) and an \(E\) value of > 200.

The specificity of TtEST was evaluated for various 4-methylumbelliferyl esters, \(\alpha\)-naphthyl esters, indoxyl acetate and \(p\)-nitrophenyl acetate and butyrate (Fig. S6). All the activities were approximately 3–11 times lower than for BnLAE. Methyl cinnamic acid was a poor substrate. No activity was found towards \(\beta\)-naphthyl caprylate, indicating some regio-selectivity of TtEST. Tributyrin and esters of fatty acids are not substrates. Similarly, BcEST activities towards \(\alpha\)-naphthylacetate, \(\alpha\)-naphthylbutyrate and \(\alpha\)-indoxylacetate were found to be 8–18 times lower than for the BnLAE substrate (Table S1).

**Stability of cloned esterases**

Organic solvents such as DMF (up to 30%) and DMSO (up to 50%) did not effect the stability (20 h incubation) nor the activity of the esterases (not shown). After several months of storage as lyophilized powder at room temperature, the enzymes retained full activity. The thermostability, thermodenaturation constant \(k_d\) and half-life time \(t_{1/2}\) of these biocatalysts are shown in Fig. 4. TtEST was fully active at 50°C for an extended period of time but lost about 40% of its activity when incubated at 70°C for 20 min. At 50°C, the \(t_{1/2}\) of BcEST was about 40 min, but lost its activity very quickly at higher temperatures. The EH5 and 2IC12 variants have greater \(t_{1/2}\) at 50°C (130 and > 200 min respectively), and also displayed increased stability at 55°C (17 and 59 min).

According to the Lumry and Eyring model (Lumry and Eyring, 1954; Sanchez-Ruiz, 1992), the irreversible denaturation of a protein is a two-step process, in which the native enzyme reversibly unfolds and is then eventually and irreversibly altered to its final non-native state. Based on the experimentally determined rate constants of denaturation \(k_d\) for the esterases, the transformed Arrhenius equation (similarly to above using \(k_{cat}\)) can be used to estimate the activation energy \(E_d\) of denaturation (unfolding). The \(E_d\) was estimated to be 18.6 ± 1.6, 21.0 ± 0.4, 30.0 ± 0.8 and 39.4 ± 3.6 kcal mol\(^{-1}\) for BcEST, EH5, 2IC12 and TtEST respectively.

**Circular dichroism spectroscopy**

Thermal unfolding experiments monitored by CD spectroscopy revealed apparent melting temperatures \(T_m\) for BcEST, EH5, 2IC12 and TtEST of 50°C, 53°C, 55°C and 65°C respectively (Fig. S5). The CD spectra of the esterases in the far UV (200–250 nm) revealed typical helical proteins. All four enzymes showed little differences in their far UV spectra and were found to consist of at least 50% \(\alpha\)-helices (not shown). Despite the low sequence identity (38%) between TtEST and BcEST, these proteins share a very similar pattern of secondary structure as predicted by PSIPRED (Jones, 1999) (Fig. 2).

**A qualitative interpretation for the enhanced thermostability in BcEST mutants EH1 and 2IC12**

In order to find a plausible qualitative explanation for the improved thermostability of the two improved variants of BcEST, we generated a homology model of BcEST using the aforementioned three structural templates. The
BcEST homology model displays the \( \alpha/\beta \)-fold around the catalytic triad, a characteristic of this superfamily of hydrolyases (Fig. 5). Three of the four mutated residues are predicted to be found in \( \alpha \)-helices (I88, Q110 and S148) while the fourth one (K272) in an extended \( \beta \)-strand region (Fig. 5). The S148 and K272 residues mutated in the first round appear surface-exposed, while the additional two residues, I88 and Q110, from the second round are presumably partially buried. The two mutations in the double mutant, S148C and K272R, are highly conserved. Based on a visual examination of the BcEST homology model one can speculate that the S148C surface mutation would slightly improve hydrophobic packing against neighbouring hydrophobic residues (e.g. I145, V253) owing to a larger, more polarizable S atom compared with the O atom at this position. The K272R surface mutation may lead to some gain in H-bonding and electrostatic interactions due to a closer approach to neighbouring acidic residues (e.g. E270, D295). However, the net stabilizing effect may not be large since such gains may also incur desolvation and entropic costs. Given the more buried location of the additional I88T and Q110L mutations from the quadruple mutant, the effects of those mutations taken individually may be relatively larger than those of the S148C or K272R surface-exposed mutations. However, these additional two mutations introduce opposite changes in the physicochemical properties at the site of mutation: I88T increases the polarity while Q110L increases the hydrophobicity. In a plausible scenario one would expect I88T and Q110L to have relatively large opposing effects on protein stability (I88T – destabilizing, Q110L – stabilizing). Hence, the net stabilizing effect of the combined buried mutations I88T and Q110L (large opposing effects) may end up as comparable to that afforded by the combined surface mutations S148C and K272R (small synergistic effects), as observed experimentally.

**Discussion**

BcEST and TtEST are two new members of the growing superfamily of \( \alpha/\beta \)-fold hydrolases (Ollis et al., 1992; Nardini and Dijkstra, 1999; Holmquist, 2000). Otherwise annotated as either a 3-oxoacid enol-hydrolase in *B. cereus* ATCC 14579 (NC_004722; GenBank AE016877) or simply a hydrolase or acyltransferase in the case of *T. tengcongensis* (NC_003869; AAM 23832.1; Bao et al., 2002), the two enzymes are characterized as alcaliphilic esterases and are remarkable for the racemic resolution of BnLAE to produce R-BnLAE with an \( E \) value > 200. Whereas BcEST has a higher catalytic activity towards BnLAE, TtEST has greatly improved overall thermostability. This pair of enzymes represents the second example of an enzymatic step in the chemoenzymatic synthesis of LFC as shown in Fig. 1, the other example being the resolution of the benzoxazine intermediate by either Bacillus sp. DSC 1012 (Miyadera and Imura, 1999) or Baker’s Yeast (Kang et al., 1996).

In the context of comparative genomics, it is noteworthy that BcEST is not an ORF specific to ATCC 14579, unlike two members with unknown function that have the \( \alpha/\beta \)-hydrolase fold (Mois et al., 2007). On the other hand, the BcEST-encoding gene and its surrounding sequences that include a putative major facilitator transport protein and a 6-aminohexanoate-dimer hydrolase homologue, are very well conserved among several of the completely sequenced genomes of the *B. cereus* group of bacteria, including the recently described *B. cereus* Q1 extremophile, isolated from a deep subsurface oil reservoir (GenBank Accession No. CP000227; Xiong et al., 2009). Unfortunately, the physiological substrate and the significance of this potential metabolic pathway are not known at this time. However, one can speculate that the BcEST-encoding gene is likely controlled by a divergently transcribed repressor of the MarR family, as suggested by the presence of two inverted repeat sequences in the 188 bp intergenic space between the marR homologue and est (Grkovic et al., 2002; Schumacher and Brennan, 2002).

In the *T. tengcongensis* genome (NC_003869), TtEST (TTE0556) is flanked by a LipA homologue (putative acetyltransferase and hydrolase with an \( \alpha/\beta \)-hydrolase fold) and a putative FabH\(_2\) [3-oxo-acyl-(acyl-carrier

**Fig. 5.** Homology model of wild-type and mutant BcEST. The overall fold of the enzyme is rendered as a yellow ribbon. The side-chains of the active-site catalytic triad residues (S102, D251 and H279) are rendered as red spheres. The side-chains of mutated residues are rendered as blue spheres.
protein) synthase III]. We speculate that this pathway is involved in fatty acid metabolism. Previously, Zhang and colleagues (2003) had cloned a thermostable esterase from *T. tengcongensis* strain MB4. Although it can hydrolyse tributyrin, it does not hydrolyse olive oil but only hydrolyses short-chain *p*-nitrophenol esters and is therefore not considered to be a true lipase. The enzyme is optimally active at 70°C and at pH 9 (Zhang et al., 2003) similar to TtEST. Indeed, the *T. tengcongensis* MB4 genome has been used as a source of thermostable proteins, such as *tengconglysin*, a new serine protease (Koma et al., 2007) and a highly thermostable single-strand DNA-binding protein (Olszewski et al., 2008).

Characterization of TtEST represents a contribution to the global proteomics of the *T. tengcongensis* genome by Wang and colleagues (2007).

We modelled BcEST according to three structural homologues in an attempt to explain the added thermostability in the double and quadruple mutants EH5 and 2IC12 respectively. The stabilizing effects observed experimentally are undoubtedly the result of a complex interplay of intramolecular interaction, desolvation and entropy contributions. One interesting hypothesis is that the I88T and Q110L mutations in the quadruple mutant may have actually large opposing effects, which when combined lead to a relatively small stabilization as observed experimentally by a mere 2°C increase of *Tm*. This leads to the prediction that a triple mutant consisting of Q110L, S148C and K272R would have an improved thermostability compared with the quadruple mutant, a hypothesis that remains to be tested.

A modelled structure of TtEST was also constructed and found to have a very similar structural environment around the catalytic triad as in BcEST and the three template structures (not shown). There appears to be no major structural differences between the modelled TtEST and BcEST structures. As amino acid composition is believed to play an intrinsic role in the stability of proteins (Farías et al., 2004; Zhou et al., 2008), the sequence of TtEST was analysed and compared with the mesophilic BcEST. For TtEST, the E+K/Q+H ratio is 4.8, which falls in the range of thermophiles (3.2–4.6) and hyperthermophiles (>4.5). This ratio is 1.5 for BcEST that fits the category of mesophiles of <2.5. However, explaining thermostabilization is more complex since it is known that a multitude of factors are involved. These include additional formation of disulfide bridges, hydrophobic or aromatic interactions, contact order, hydrogen bonding, ion pairing, dimer–dimer interaction, stability of α-helices, reduction of the entropy of unfolding etc. (Vieille and Zeikus, 2001; Robinson-Rechavi and Godzik, 2005; Zhou et al., 2008).

The experimental values of activation energies for denaturation of the BcEST and variants, and TtEST esterases are in good agreement with the respective *Tm* values. It has been noted that thermostable enzymes, although adapted to work at high temperatures, are not optimized in terms of activity, whereas their mesophilic counterparts often show optimized catalytic activities but low stability (Vieille and Zeikus, 2001). The BcEST and TtEST described in this study are another example.

Owing to the wide-ranging use of hydrolases in industry as well as academic interest, protein engineering efforts applied to this family of enzymes have been intense and have utilized a number of approaches (Jaeger et al., 1999; Qian et al., 2007). Using directed evolution and selection at higher temperature we have obtained two thermostable variants. However, the modest increase of 5°C in *Tm* in one quadruple mutant was less than the 15°C increase in a genome-mined homologue derived from a naturally occurring thermophilic organism. This suggests that nature has outrun directed evolution in this thermostabilization case, contrary to many examples of directed evolution efforts where *bona fide* thermophilic counterparts (unlike BcEST–TtEST pair) are not readily available (Woodyer et al., 2004). In any case, judicious genome mining can be a productive and economical way of new biocatalyst discovery. More importantly, functional characterization of the sequence space is key, even though the ‘road’ can be difficult (White, 2006).

Presently, a popular approach to discovering new and improved proteins is through metagenomics, with the marine environment as a rich source for the discovery of novel enzymes (Ferrer et al., 2005; Kennedy et al., 2008). With reference to racemic resolution of building blocks or precursors for pharmaceutical compounds such as antibiotics, Park and colleagues (2007) had reported on a *Vibrio*–derived esterase (Vip509) that is capable of hydrolysing the 5-enantiomer of racemic ofloxacin propyl ester. In a more recent study, two esterases (EstAT1 and EstAT11) from the Arctic sediment metagenome were found to hydrolyse 5–ofloxacin butyl ester (Jeon et al., 2009). In either case, the enantiomeric excess (e.e.) value is modest at 42.9% for Vip509 and up to 70.3% for EstAT11.

In summary, this study illustrates that despite an enormous database on esterase sequences (ESTHER; http://bioweb.ensam.inra.fr/esther) there is ample room for new findings and it is particularly rewarding for potential industrial applications. There is an increasing demand for the environmentally benign synthesis of enantipure compounds as chiral building blocks for drug synthesis such as antibiotics (Agranat et al., 2002). Given a multistep synthesis for molecule such as LFC, a chemoenzymatic route appears attractive since it combines proven economics of some unavoidable chemical processes with environmental care through green chemistry.
Experimental procedures

The description of the experimental procedures is contained in Appendix S1 in Supporting information.

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Supporting information

Additional Supporting Information may be found in the online version of this article:

**Fig. S1.** Coomassie blue-stained protein gel of *E. coli* crude extracts separated on SDS-10% PAGE.

**Fig. S2.** Phylogenetic analysis of BcEST. The GenBank accession numbers for the respective proteins, annotated as either α/β-fold family hydrolase, 3-oxoadipate enol-lactonase, or in some cases, unnamed protein product or conserved hypothetical protein, are as follows: (1) ACK89548.1, (2) AAP27115.1, (3) AAS42215.1, (4) ACM13522.1, (5) AU17266.1, (6) AAT62319.1, (7) ACO26908.1, (8) ABY44194.1, (9) ACK96374.1, (10) BcEST (AAP10252.1), (11) ACK59771.1, (12) ABS22367.1, (13) ABX31253.1, (14) ACB86291.1, (15) ABV36133.1, (16) ABZ77297.1, (17) ACL19790.1, (18) ABX81126.1, (19) TtEST (AAM23832.1), (20) EAS01633.1, (21) EAR92522.1, (22) CAAK6695.1, (23) BAH42062.1, (24) BAD39794.1, (25) ACL97306.1, (26) AAK25687.1, (27) ABI78668.1, (28) ABC62566.1, (29) ABS62973.1, (30) ABN96204.1, (31) ABG51905.1, (32) BAH44442.1, (33) ABK86474.1, (34) ABM15056.1, (35) ABP44844.1, (36) ACO24661.1 (1U2E), (37) ACC84816.1, (38) AAB60168.1 (1VA4), (39) AAC43253.1 (1A8Q). The tree was generated using MEGA4 (Kumar et al., 2008).

**Fig. S3.** Heat inactivation of BcEST and variant enzymes.

**Fig. S4.** Estimation of free energy of activation in the enzymatic reaction of (A) BcEST and variants and (B) TtEST.

**Fig. S5.** *T. maritima* of the BcEST and variant enzymes, and TISTEST.

**Fig. S6.** Colorimetric assay of esterase activity using Phenol-Red.

A. Time-course of enzyme-induced colour shift of Phenol-Red.

B. Standard curve for the quantification of enzymatic reaction.

C. Linear increase of A<sub>435</sub> over time due to esterase activity.

**Table S1.** Comparison of the substrate specificity of BcEST and TIEST

**Appendix S1.** Experimental procedures

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