Residue-Specific Incorporation of the Non-Canonical Amino Acid Norleucine Improves Lipase Activity on Synthetic Polyesters

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Environmentally friendly functionalization and recycling processes for synthetic polymers have recently gained momentum, and enzymes play a central role in these procedures. However, natural enzymes must be engineered to accept synthetic polymers as substrates. To enhance the activity on synthetic polyesters, the canonical amino acid methionine in Thermoanaerobacter thermohydrosulfuricus lipase (TTL) was exchanged by the residue-specific incorporation method for the more hydrophobic non-canonical norleucine (Nle). Strutural modelling of TTL revealed that residues Met-114 and Met-142 are in close vicinity of the active site and their replacement by the norleucine could modulate the catalytic activity of the enzyme. Indeed, hydrolysis of the polyethylene terephthalate model substrate by the Nle variant resulted in significantly higher amounts of release products than the Met variant. A similar trend was observed for an ionic phthalic polyester containing a short alkyl diol (C5). Interestingly, a 50% increased activity was found for TTL [Nle] towards ionic phthalic polyesters containing different ether diols compared to the parent enzyme TTL [Met]. These findings clearly demonstrate the high potential of non-canonical amino acids for enzyme engineering.

Keywords: polyester modification, enzyme hydrolysis, genetic code engineering, lipase, Thermoanaerobacter thermohydrosulfuricus, TTL, norleucine

INTRODUCTION

With the increasing utilization of synthetic polymers in today’s society, polymer surface functionalization and recycling processes have become enormously important. Surface modification is generally performed with harsh chemical methods, photo grafting, or energy-intensive plasma processes (Mozetić, 2019). These conventional methods are often toxic, expensive and adversely affect the mechanical properties of the polymer (Hetemi and Pinson, 2017). Enzymatic surface functionalization of polyesters has received increased attention as an environmentally friendlier and highly specific process (Pellis et al., 2015; Weinberger et al., 2017; Biundo et al., 2018). Limited enzymatic hydrolysis allows targeted surface functionalization while leaving the
polymer bulk properties unaffected. Complete enzymatic hydrolysis of polyesters, on the other hand, allows specific recovery of the valuable building blocks for further synthesis especially from blends and composite materials which is otherwise quite challenging (Gamerith et al., 2017; Wei and Zimmermann, 2017). Enzymatic hydrolysis has so far been reported for various synthetic polymers such as poly-L-lactic acid (PLLA) (Pellis et al., 2015; Hajighasemi et al., 2016; Bonifer et al., 2019), polyethylene terephthalate (PET) (Herrero Acero et al., 2011; Sulaiman et al., 2012; Ribitsch et al., 2013; Kawai et al., 2014; Dimarogona et al., 2015; Barth et al., 2016; Yoshida et al., 2016; Kawabata et al., 2017), polybutylene adipate terephthalate (PBAT) (Biundo et al., 2016; Wallace et al., 2017) and polyurethanes (PU) (Acero et al., 2012; do Canto et al., 2019). To date, most enzymes used for polyester hydrolysis are esterases, lipases or cutinases from typical mesophilic or thermophilic compost microorganisms (Kleeberg et al., 2005; Herrero Acero et al., 2011; Ribitsch et al., 2011; Ribitsch et al., 2012a; Ribitsch et al., 2012b; Pellis et al., 2015). Only few enzymes were isolated from anaerobic (Biundo et al., 2016; Perz et al., 2016a; Perz et al., 2016b) and aquatic bacteria (Haernvall et al., 2017a; Haernvall et al., 2017b; Wallace et al., 2017; Haernvall et al., 2018).

Over the last years, several protein engineering strategies have been reported to tailor the enzymes for the non-natural polymeric substrates, thus improving the enzymatic hydrolysis of synthetic polymers. Different approaches have been applied to adapt the active site to the bulky polymers (Thumarat et al., 2012) and to reduce inhibition effects caused by release products (Roth et al., 2014). Besides, the adsorption/desorption of the enzymes on the polyester has been improved which turned out to be a crucial and rate-limiting step in enzymatic polymer hydrolysis (Herrero Acero et al., 2013; Ribitsch et al., 2013; Ribitsch et al., 2015). Today it is well known that temperatures higher than the glass transition temperature (Tg) increase the hydrolysis rates due to a higher flexibility of the polymer chains and therefore increased accessibility to enzymes (Kawai et al., 2019). For this reason, the thermostability of polymer hydrolysing enzymes has become a highly important issue and target for protein engineering (Shirke et al., 2016; Son et al., 2019).

The conventional protein engineering approaches use the 20 canonical amino acids (cAAs) prescribed by the genetic code (Steiner and Schwab, 2012). Still providing a very potential tool, conventional protein engineering is limited by the site-chain chemistry offered by the 20 cAAs. An alternative approach for generating more robust enzymes or altering physico-chemical properties is incorporation of non-canonical amino acids (ncAAs) (Hoesl and Budisa, 2012; Wiltschi et al., 2020). The diverse and unusual side chain chemistries (Dumas et al., 2015) make them attractive building blocks for protein engineering (Link and Tirrell, 2003; Voloshchuk and Montclare, 2009; Zheng and Kwon, 2012; Wiltschi et al., 2020). Under tightly controlled conditions, many ncAAs can be incorporated into a target protein by exploiting the translational machinery of the host (Cowie and Cohen, 1957). This residue-specific incorporation (SPI) exploits the natural substrate tolerance of aminoacyl-tRNA synthetases (aaRS) for which various methods have been developed (Budisa, 2004; Hoesl et al., 2011; Ngo and Tirrell, 2011). The non-canonical amino acid norleucine (Nle) is an isosteric carba-analog of methionine (Met) that can be incorporated by the SPI method using a Met auxothrophic strain (Hoesl et al., 2011). The carbon atom of the methylene group of Nle is less electronegative than the corresponding sulfur atom of Met (2.48 vs 2.56) (Xie et al., 1995), resulting in a slightly increased hydrophobicity of the methylene group in comparison to the sulfur atom (Thomson et al., 1994). This subtle difference can exert a cumulative impact when several Met residues are globally exchanged for Nle. For instance, Budisa and co-workers successfully used Nle and other Met analogs to probe protein folding (Budisa et al., 1998) and control the aggregation behavior of prion protein (Wolschner et al., 2009). Nle is a comparably inexpensive ncAA. It can be produced in very good quantities by biosynthesis from glucose and the scalability of this bioprocess was previously demonstrated (Anderhuber et al., 2016).

In the past, several groups demonstrated that the incorporation of a palette of different ncAAs into target proteins could improve...
enzyme activity (Parsons et al., 1998; Cirino et al., 2003; Hoels et al., 2011). NcAA incorporation can increase protein stability (Moroder and Budisa, 2010), especially under harsh conditions, such as tolerance to organic solvents (Deepankumar et al., 2014) or extreme pH values in combination with high temperatures (Votchitseva et al., 2006). For an overview of the engineering of enzymes with ncAAs, the interested reader is referred to (Wiltschi et al., 2020). A whole set of different ncAAs was successfully incorporated into the lipase from the anaerobic extreme thermophilic microorganisms Thermoanaerobacter thermohydrosulfuricus (TTL) by Budisa and coworkers (Acevedo-Rocha et al., 2013), demonstrating the positive effect on enzyme activity and stability in harsh conditions. TTL was first described by Royter and coworkers (Royter et al., 2009) as an enzyme with activity at a broad temperature (40°C–90°C) and pH range (pH 6.5–10). However, incorporation of ncAAs resulted not only in enhanced hydrolytic residual activity upon treatment with organic solvents by up to 450% and surfactants by up to 1630% (Hoels et al., 2011), but has also reduced denaturation, alkylating and inhibition processes (Acevedo-Rocha et al., 2013).

In this study, we explored the incorporation of a ncAA into TTL to tune the physicochemical properties of the enzyme for enhanced polyester hydrolysis. As a proof of concept, the methionines (Met) of TTL were exchanged globally by its slightly more hydrophobic analog Nle. By the residue-specific incorporation of Nle into TTL, we aimed at altering the active site and/or the substrate-entrance channel to better fit the polymeric substrate. Synergistically, the enzyme surface was adapted to promote a better interaction/adhesion to the polymer. Two groups of polyester model substrates with various alkyl and ether diols were subjected to hydrolysis to evaluate the effect of Nle incorporation on polyester hydrolysis.

**MATERIALS AND METHODS**

**Chemicals and Substrates**

Alkyl diols (1,5-pentanediol, 1,8-octanediol and 1,12-dodecanediol) and ether diols (diethylene glycol, triethylene glycol and tetraethylene glycol), terephthalic acid (TA) and 5-sulfophthalic acid (NaSIP) were purchased from Sigma-Aldrich, Merck KGaA (Darmstadt, Germany) or Carl Roth (Karlsruhe, Germany), if not stated differently. Norleucine was purchased from Sigma-Aldrich (St. Louis, MO). Buffer components, bovine serum albumin (BSA) and methanol (HPLC grade) were also purchased from Sigma-Aldrich.

All standard chemicals used in this work were purchased from Sigma-Aldrich, Merck KGaA (Darmstadt, Germany) or Carl Roth (Karlsruhe, Germany), if not stated differently. Norleucine was obtained from IRIS Biotech GmbH (Marktredwitz, Germany). Enzymes for cloning and PCR were from Thermo Fisher Scientific (Waltham, MA). PCRs were performed using TaKaRa Ex Taq® High-Fidelity DNA polymerase (Clontech Laboratories, Inc., Mountain View, CA) or Dream Taq® DNA polymerase (Thermo Fisher Scientific). PCR primers were ordered from IDT Inc. (Coralville, IA) in standard desalted quality.

**Strains and Plasmids**

*E. coli* B834 (DE3) (*E. coli* B F– ompT hsdSB(rB– mB–) dcm+ gal met λ(DE3); Merck KGaA) was the host for the SPI experiment and *E. coli* DH5α (*E. coli* K–12 F– endA1 supE44 thi-1 recA1 relA1 gyrA96 proA q80lacZAM15 Δ(lacZYA-argF)U169 hsdR17 (rK– mK+ λ–); Thermo Fisher Scientific) was used for cloning experiments and plasmid propagation. Transformation of *E. coli* was carried out by electroporation as described by Seidman et al. (2001). pTTL was constructed according to Anderhuber et al. (2016). Briefly, the coding sequence of the lipase from *T. thermohydrosulfuricus* was PCR-amplified from synthetic DNA (IDT Inc.) using primers BPp244 and BPp245 (Anderhuber et al., 2016). The PCR fragment was inserted into pQE80L (Qiagen, Hilden, Germany) cut with EcoRI and HindIII by Gibson isothermal assembly (Gibson et al., 2009).

**Enzyme Expression and Purification**

The SPI experiment was performed as described previously (Anderhuber et al., 2016). Briefly, M9 minimal medium (M9 MM) containing M9 salt buffer (48 mM Na2HPO4, 22 mM KH2PO4, 9 mM NaCl and 19 mM NH4Cl) was used supplemented with 1 mM MgSO4, 7 µM CaCl2, 1 mg/L thiamine and trace elements (9 µM FeSO4, 3.5 µM MnSO4, 2.5 µM AlCl3, 2 µM CoCl2, 0.4 µM ZnSO4, 0.5 µM Na2MoO4, 0.4 µM CuCl2 and 0.5 µM H2BO3). 100 mg/L ampicillin was added for plasmid maintenance. 20 mM glucose was provided as the carbon source. For Met depletion at OD600~ 3, 3.5 g/L yeast extract (Carl Roth) were supplemented in the medium. Cells were grown in baffled shake flasks at 37°C with vigorous shaking. After depletion of Met, as indicated by growth arrest, the cultures were supplemented with 1 mM of either Met (cAA, parent TTL [Met]) or Nle (Nle, synthetic TTL [Nle]). Gene expression was induced by adding IPTG (VWR International, Vienna, Austria) to a final concentration of 0.5 mM and performed with vigorous shaking for 4 h at 30°C. Cells were harvested at 4,000 g for 20 min at 4°C. Cell pellets were reduspended in 30 ml Ni-NTA Lysis Buffer (50 mM Na2HPO4, 300 mM NaCl, 10 mM imidazole, pH 7.4) and incubated for 30 min on ice. Cells were disrupted by sonication on ice (Branson Sonifier 250, Emerson Electric, St. Louis, MO). The sonication was performed for 6 min with the following settings: duty cycle: 70%; output: 7-8, sonication tip Φ~1 cm. After centrifugation for 45 min at 40,000 x g and 4°C the lysates were filtered through 0.2 µm syringe filters and purified on Ni-NTA sepharose according to the manufacturer’s protocol (IBA GmbH, Goettingen, Germany). Finally, the buffer was exchanged for 100 mM Tris-HCl pH 7.0 with PD-10 columns (GE Healthcare, Chicago, IL).

**Protein Quantification**

The Bradford based Bio-Rad Protein Assay (Bio-Rad Laboratories GmbH, Munich, Germany) with bovine serum albumin as standard was used to determine protein concentrations of purified enzymes. The protein assay was performed according to the manufacturer’s instruction.

**Intact Protein Mass Determination**

The protein solutions were desalted using Amicon Ultra 0.5 ml centrifugal filter units (Millipore, Billerica, MA). A final protein concentration of 30 pmol/µL was obtained with water containing 5% acetonitrile and 0.1% trifluoroacetic acid. The separation of
possible protein variants was carried out on a capillary HPLC system (1200 Agilent, Agilent Technologies) using a PepSwift RP monolithic column (50 × 0.5 mm, Thermo Fisher Scientific) at a flow rate of 20 µL/min and a column temperature of 60°C. The gradient of solution A (water +0.05% TFA) and B (ACN +0.05% TFA) was performed as follows: 10% B for 5 min, 10%–100% B for 50 min, 100%–10% B for 1 min, 10% B for 15 min. The injection volume was 5 µL. The Thermo LTQ-FT mass spectrometer (Thermo Fisher Scientific) was operated with an ESI source in positive mode with the following settings: mass range: 300–2000 m/z, resolution 400000, 500 ms injection time, 1 microscan, source voltage 5 kV, capillary voltage 35 V, sheath gas flow 15. The protein mass spectra were deconvoluted by the Thermo Fisher Scientific software Protein Deconvolution 2.0, using the Xtract algorithm. The following main parameters were applied: charge carrier, H+; m/z range, minimal 800 to maximal 2000; minimal detected charge state, 4; s/n threshold, 5; relative abundance threshold, 20%. Trace amounts of unlabeled species might be present but fall below the detection limit of the mass spectrometry method (2–5%).

Structure Modeling
The structure of the TTL was modeled using the CATALOphore platform of Innophore GmbH (www.innophore.com) employing the program Yasara version 20.4.24.L.64 (www.yasara.org). The model is based on the structure of feruloyl esterase (Est1E) from Butyryrivibrio proteoclasticus (PDB 2WTM) (Goldstone et al., 2010) as template, which shares 32% sequence identity and 50% sequence similarity with TTL. The active site cavity in the vicinity of the catalytic triad Ser-113, His-233 and Asp-203 was calculated using the program CavMan (Innophore GmbH, www.innophore.com) employing the LIGSITE algorithm (Hendlich et al., 1997). For the analysis of the hydrophobicity of the cavity, the hydrophobicity module of the program VASCo (Steinkellner et al., 2009) as implemented in CavMan was used.

Polyester Synthesis and Characterization
The oligomeric model substrate bis-(benzoyloxyethyl) terephthalate (3PET) was synthesized as previously reported (Heumann et al., 2009). Other polyesters were synthesized in a two-step process and characterized as described by Haernvall et al. (2017b).

Hydrolysis of Oligomeric and Polymeric Materials
Model Substrates Polyester powders (10 mg/ml) were incubated in 1 ml of 100 mM Tris-HCl pH 7.0 and in the presence of 1 µM enzyme. The reaction mixture was shaken for 3 h at 70°C and 100 rpm (Infors HT Multitron, Infors AG, Bottmingen, Switzerland). In parallel, enzymes and polymers were incubated in pure buffer as blank reactions. All experiments were run in triplicates. Enzymes were precipitated by addition of ice-cold methanol (1:1 v/v), acidified with 0.1 M HCl to pH 4 and sedimented in a tabletop centrifuge (15 min, 0°C, 14,000 rpm; Hermle Labortecnik GmbH, Wehingen, Germany). Supernatants were used for HPLC analysis.

Determination of Hydrolysis Products
After hydrolysis, samples were analyzed by HPLC-UV on a system consisting of a Dionex UltiMate 181 3000 Pump (Dionex Cooperation, Sunnyvale, United States), a Dionex ASI-100 automated sample injector, a Dionex UltiMate 3000 column compartment and a Dionex UVD 340 U photodiode array detector. The hydrolysis products were separated by a reversed-phase column, [XTerra® RP18, (3.5 µm, 3.0 mm × 150 mm)] (Waters Corporation, Milford, United States) using a non-linear gradient where eluent A consisted of water, eluent B of methanol and eluent C of 0.01 N sulfuric acid. The separation was achieved by a non-linear gradient increased from 15% A to 40% A from 13 to 30 min, followed by an increase to 90% A during 5 min which was kept for 10 min to then be re-established to initial conditions within 1 min and equilibrated for 20 min. The injection volume was 5 µL, and the flow rate was 0.4 ml/min. The column compartment had a constant temperature of 40°C. The expected release products TA and NaSIP were detected via UV/VIS spectroscopy, and the release products were qualified and quantified based on calibration curves.

Enzyme Adsorption on PET
Enzyme adsorption on PET film was monitored as previously described (Ribitsch et al., 2013). Briefly, 0.5 × 1 cm² PET films were washed at 50°C in three consecutive steps of 30 min each with 1.5 ml each of 0.5% (v/v) Triton X-100, 100 mM Na₂CO₃ and deionized water. The washed membranes were incubated in 1 ml of 0.6 mg/ml enzyme solution for 2 h at 30°C. The films were washed twice by dipping into 1.5 ml TBST (25 mM Tris, 0.15 M NaCl, pH 7.6, 0.05% (v/v) Tween®-20) at room temperature. Then, the films were incubated with HisProbe (1:2500 dilution in TBST; SuperSignal West HisProbe Kit; Thermo Fisher Scientific, Waltham, MA) for 40 min at room temperature with shaking. The films were washed three times by dipping into 1.5 ml TBST as described above and were developed by incubating in 600 µL of SuperSignal West Pico Substrate Working Solution for 5 min. Chemiluminescence signals were detected using Gbox Chemi HR16 and GeneSnap image acquisition software Version 7.05.01 (Syngene, Cambridge, United Kingdom) and quantified with Colorlite sp850 (Colorlite:Inovative color measurements; Germany).

RESULTS AND DISCUSSION
The increasing interest in enzymatic polymer functionalization and enzymatic polymer recycling processes has resulted in various proposed strategies to improve the enzymatic hydrolysis of synthetic polymers. Here, in order to evaluate a novel approach, the classical protein engineering strategies were expanded to also include ncAAs. To assess the effects on enzymatic polymer hydrolysis, a lipase from an anaerobic extreme thermophilic microorganism was modified by exchanging the cAA methionine with the slightly more hydrophobic ncAA norleucine. The effects were evaluated towards oligomeric and polymeric polyester model substrates with systematically varied compositions to enable a mechanistic
study. Therefore, a set of structurally different polyester substrates was used (Figure 1). The water-insoluble PET model substrate bis(benzoyloxyethyl) terephthalate (3PET) has been widely used in the past for the detection of PET hydrolysing enzymes such as cutinases (Herrero Acero et al., 2011; Ribitsch et al., 2011; Kawai et al., 2019).

Only few enzymes were identified so far that show an ability to hydrolyze ionic phthalic acid based polyesters (Haernvall et al., 2017a). Ionic phthalic polyesters are found in many products in our daily life, such as household products. In this study, we used model polyesters consisting of the aromatic terephthalic acid (TA) and the ionic aromatic 5-sulfoisophthalic acid (NaSIP), which were linked by altering alkyl and ether diols. Two groups of ionic phthalic polyesters were used to evaluate the effect of chain length and hydrophilicity/hydrophobicity of polyesters influencing the hydrolysis. The first group of polyesters contained alkyl diols (1,5-pentanediol (C5), 1,8-octanediol (C8), and 1,12-dodecanediol (C12) (Figure 1B). The second group of polyesters contained ether diols (ethylene glycol (EG1), diethylene glycol (EG2), triethylene glycol (EG3), and tetraethylene glycol (EG4)) (Figure 1C), with systematically varied chain lengths. The ratio of TA and NaSIP was kept constant (70:30 mol%).

Structure Modelling of TTL

The modelled structure of the lipase from *Thermoanaerobacter thermohydrosulfuricus* (TTL) exhibits a typical α/β-hydrolase
fold with a central eight-stranded, mostly parallel β-sheet flanked by six α-helices (Figure 2). A catalytic triad is formed by the amino acid residues Ser-113, His-233 and Asp-203, with the serine being embedded in a GLSMGG sequence motif. The oxyanion hole is built up by the mainchain amide groups of Phe-37 and Met-114. The active site cavity is located at the upper end of the α/β-hydrolase core and delimited by a small cap domain consisting of residues 140 to 180. The TTL-sequence contains a total of 11 methionine residues, of which Met-114 and Met-142 are in close vicinity to the active site residue (Figure 2). It is conceivable that the replacement of those two methionine residues by norleucine modulates the catalytic activity of the enzyme most likely by altering its substrate binding properties. Most of the remaining methionine residues are part of the hydrophobic core of the protein (Figure 2). Amino acid replacements in this region could change the dynamic properties of the enzyme and its stability. Indeed, Hoesl et al. (2011) observed that the global Met → Nle substitution in TTL had a positive impact on the hydrolysis as indicated by a ~30 % higher amounts of hydrolysis products than to the parent enzyme (Figure 1A). Nevertheless, the pattern of the released molecules was similar for the two enzymes TTL (Met) and TTL (Nle). Benzoic acid (BA) was released in the highest concentration, followed by mono-(2-hydroxyethyl) terephthalic acid (MHET), terephthalic acid (TA) and hydroxyethylbenzoate (HEB), respectively. The total amount of released molecules for the hydrolysis of the oligomer model substrates was in the same range as previously reported for

**TTL Expression, Purification and Characterization**

TTL was expressed in *E. coli* and purified from cleared cell lysates over a C-terminal 6xHisTag by affinity chromatography. Expression in the presence of the canonical Met yielded a defined band migrating at the 29.2 kDa molecular weight marker band as calculated for the theoretical molecular weight of TTL (Supplementary Figure S1). In contrast, the lipase expressed in the presence of Nle appeared as a blurred band at slightly lower molecular weight. In agreement with a previous report (Hoesl et al., 2011), the Nle variant exhibited accelerated electrophoretic mobility compared to the Met variant. The altered migration behavior was a first indication for the successful incorporation of Nle into the lipase. Typically, 20 mg/ml TTL [Met] and 10 mg/ml TTL [Nle] were purified from 3 to 4 g of cell pellet obtained from cultures in shake flasks.

To analyze the incorporation efficiency of Nle, Met and Nle variants were subjected to intact protein mass analysis (Supplementary Figure S2). In the mass spectrum of TTL [Nle] the protein species with all 11 Met residues exchanged for Nle was the most prominent peak. A minor peak representing lipase with 10 Met residues exchanged for Nle was detected as well. This is in good agreement with previously published results (Hoesl et al., 2011).

**Hydrolysis of Oligomeric and Polymeric Model Substrates**

The effect of the incorporating Nle into TTL on polymer degradation was assessed in a first step using the oligomeric model substrate 3PET. Interestingly, the global Met→Nle substitution in TTL had a positive impact on the hydrolysis as indicated by a ~30 % higher amounts of hydrolysis products than to the parent enzyme (Figure 1A). Nevertheless, the pattern of the released molecules was similar for the two enzymes TTL (Met) and TTL (Nle). Benzoic acid (BA) was released in the highest concentration, followed by mono-(2-hydroxyethyl) terephthalic acid (MHET), terephthalic acid (TA) and hydroxyethylbenzoate (HEB), respectively. The total amount of released molecules for the hydrolysis of the oligomer model substrates was in the same range as previously reported for
other enzymes, such as esterases from *Clostridium botulinum* (Perz et al., 2016a), a lipase from *Thermomyces lanuginosus* (Eberl et al., 2009), or cutinases from *Fusarium solani pisi* as well as from various *Thermobifida* species (Herrero Acero et al., 2011; Ribitsch et al., 2012b).

To further evaluate the impact of norleucine incorporation on polyester hydrolysis and to obtain a more detailed mechanistic insights into the hydrolysis behavior of TTL [Nle], a set of structurally different ionic phthalic acid polyesters were investigated. In a first step, TTL [Nle] was incubated with polymeric model substrates containing alkyl diols of different chain lengths, C5, C8, and C12 (Figure 1B). TTL [Nle] and the parent enzyme TTL [Met] were both active towards C5 while only a very low activity was detected towards C8 and C12 as indicated by the release of terephthalic acid. TTL [Nle] showed around 5% more activity towards C5 compared to TTL [Met]. However, enzymatic hydrolysis of polymeric substances is influenced by several parameters, such as water solubility of the substrates, crystallinity, molecular weight, glass transition temperature (Tg) and therefore difficult to predict (Chamas et al., 2020). As described recently (Haernvall et al., 2017b), polyesters C8 and C12 have lower water solubility, higher hydrophobicity and a higher crystallinity of 4 and 12%, respectively, than C5 with crystallinity below 1%, which may have influenced the hydrolysis. Low water solubility, high hydrophobicity (Okada et al., 1997) and crystallinity have previously been reported to have a negative impact on the enzymatic hydrolysis of polymers (Donelli et al., 2010). However, the glass transition temperature of the polymeric model substrates is decreasing with increasing chain length which would have been expected to have a positive impact on the enzymatic hydrolysis due to the higher flexibility of the polymer chains (Marten et al., 2003).

In a next step, TTL [Nle] and the parent enzyme TTL [Met] were incubated with the ether diol containing model substrates, EG2, EG3, EG4 (Figure 1C). Surprisingly, an up to 40% increased activity was found for TTL [Nle] towards all three polyesters (EG2, EG3, and EG4) when compared to the parent enzyme TTL [Met]. The highest activity was again detected on the shortest diol chain length, namely EG2, decreasing with increasing chain length. It can also be noted that an overall higher activity was detected for both enzymes towards all three ether diol containing substrates when compared to the alkyl diol analogs. This can be a consequence of the increased water solubility of the ether diol containing polymers and the increased hydrophilicity. It has previously been shown that increased water solubility and increased hydrophilicity is enhancing enzymatic hydrolysis. It has even been suggested as a parameter to tune polymeric biodegradation (Gigli et al., 2013).

In none of the cases, NaSIP was detected after hydrolysis, indicating that TTL [Met] and TTL [Nle] have a limited capacity to cleave ester bonds in close vicinity to the ionic monomer NaSIP. These results are in accordance with previously reported data for cutinase A and an arylesterase from *Pseudomonas pseudoalcaligenes* and a lipase from *Pseudomonas pelagia* (Haernvall et al., 2017a; Haernvall et al., 2017b).

We analyzed the adsorption of TTL [Met] and TTL [Nle] on PET films as described in the methods section. Adsorbed enzymes were detected by binding of horseradish peroxidase labeled HisProbe to the hexahistidine-fusion tag and chemiluminescence detection. TTL [Nle] adsorbed reproducibly better to the PET films than the parent protein (Figure 3). This result supports our notion that Nle improves the interaction of TTL with the polymer surface although a different...
polyster had been used. Both, TTL [Met] and TTL [Nle] survived well the incubation with PET because their esterase activities after the incubation, e.g., with PET, was not lower than before (Supplementary Information, Supplementary Figure S3). Our finding corroborates the extraordinary stability of TTL and its Nle reported previously (Hoesl et al., 2011).

CONCLUSION

Recently, ncAAs have become a valuable asset for protein engineering, e.g., to introduce non-natural chemical functionalities into enzymes. In this study we have shown for the first time that incorporation of the ncAA Nle into a lipase from Thermoanaerobacter thermohydrosulfuricus has a positive impact on the enzymatic hydrolysis of synthetic polyesters. Nle, the carba-analog of Met, is less polar and more hydrophobic than Met while the structures are virtually identical. The global replacement of Met by Nle can tune the hydrophathy of a protein in a subtle way that is difficult to attain by exchanging Met with a hydrophobic canonical amino acid. Nle lacks the sulfur atom which is replaced by a methylene group. Due to the lack of the sulfur atom, Nle cannot form sulfoxides protecting the enzymes from oxidative stress and is also claimed to prevent protein aggregation and chemical degradation. Our structural model of TTL predicts Met-114 and Met-142 to be ocated in close vicinity of the catalytic triad and the active site cavity and two other methionines, Met-147 and Met-158, to be part of the lid. Previously, Budisa et al. (Acevedo-Rocha et al., 2013) hypothesized that replacing methionine residues in the lid could have retained it in an open conformation. This together with the replacement of Met-114 and Met-142 near the catalytic triad might have modified the environment in the substrate entry tunnel and the catalytic site such that the hydrolysis of the hydrophobic synthetic polymers was enhanced. Further studies reveal whether a similar enhancing effect can be elicited in other polyester hydrolyzing enzymes that accommodate Met residues in their substrate entry tunnels and/or active sites. If this is the case, Nle—and eventually also other ncAAs—could complement the existing tools for tuning and improving the activity of enzymes on polymers.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

KH, PF, HS, and SZ performed the experiments. DR, KG, BW, and GG planned the experiments. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fbioe.2022.769830/full#supplementary-material

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