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Nylon-Oligomer Hydrolase Promoting Cleavage Reactions in Unnatural Amide Compounds

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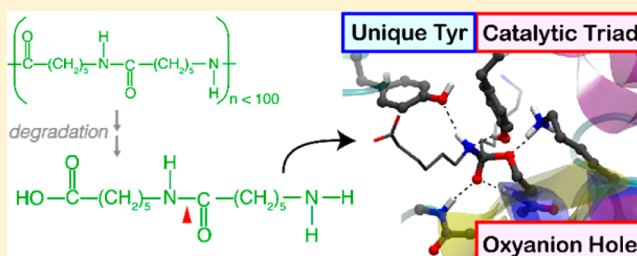
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Supporting Information

ABSTRACT: The active site of 6-aminohexanoate-dimer hydrolase, a nylon-6 byproduct-degrading enzyme with a β -lactamase fold, possesses a Ser112/Lys115/Tyr215 catalytic triad similar to the one of penicillin-recognizing family of serine-reactive hydrolases but includes a unique Tyr170 residue. By using a reactive quantum mechanics/molecular mechanics (QM/MM) approach, we work out its catalytic mechanism and related functional/structural specificities. At variance with other peptidases, we show that the involvement of Tyr170 in the enzyme–substrate interactions is responsible for a structural variation in the substrate-binding state. The acylation via a tetrahedral intermediate is the rate-limiting step, with a free-energy barrier of ~ 21 kcal/mol, driven by the catalytic triad Ser112, Lys115, and Tyr215, acting as a nucleophile, general base, and general acid, respectively. The functional interaction of Tyr170 with this triad leads to an efficient disruption of the tetrahedral intermediate, promoting a conformational change of the substrate favorable for proton donation from the general acid.

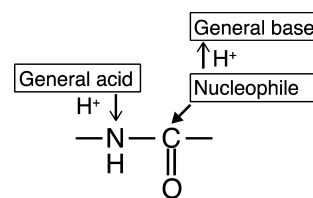
SECTION: Biophysical Chemistry and Biomolecules



Amide bond cleavage is a fundamental reaction of living organisms,^{1–9} and an important case is represented by peptidases catalyzing the hydrolysis of peptide bonds.^{1–8} Such an enzyme is ubiquitous and exhibits either nonspecific protein degradation or highly specific cleavage of protein substrates in physiological processes, such as activation of zymogens and hormone release.^{1,2,5–8} From the evolutionary standpoint, proteases with purely digestive functions have developed a high degree of specificity and efficiency to degrade amide compounds, in virtue of a characteristic set of functional amino acid residues in the active site organized in a specific configuration.^{1,7} The most common amide bond cleavage mechanism of peptidases is the polarization of a peptide bond upon nucleophilic attack on the carbon–oxygen bond, assisted by a proton abstraction from the nucleophile and following a proton donation to the peptide nitrogen in a general base/acid (Scheme 1).^{1–8}

Besides natural compounds, a peptidase exists having the ability to cleave the amide bond in artificial compounds. Such a hydrolase promotes the degradation of a nylon-6 manufacture byproduct, namely, 6-aminohexanoate (Ahx) oligomers

Scheme 1. Common Mechanism of Amide Bond Cleavage by Peptidases



(termed nylon-oligomers).^{10–19} Nylon-oligomer hydrolases were discovered in a bacterium (*Arthrobacter* sp. KI72) from soil near nylon factories in 1972.¹⁰ The bacterium evolved under environmental conditions developing a specific ability to use nylon-oligomers as a source of C and N, exploiting three types of nylon-oligomer hydrolases: Ahx-cyclic-dimer hydrolase

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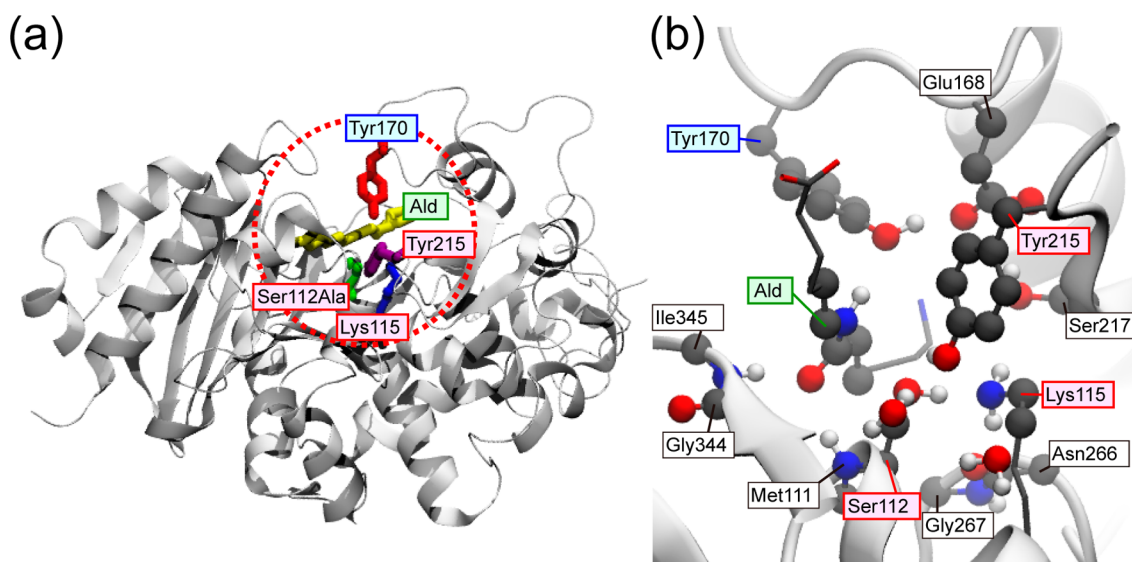


Figure 1. (a) X-ray structure of the NylB-type nylon-oligomer hydrolase. The Ser112/Lys115/Tyr215 catalytic triad and the unique Tyr170 residue are shown in green, blue, purple, and red, respectively, while the Ald substrate is yellow. (b) Atomic-level detail of the active site for the QM/MM model, corresponding to the portion inside the red dotted circle of panel a. The QM atoms are shown as balls and the MM subsystem is shown as ribbon model. Solvent water molecules are not shown for clarity. Here and in the following figures the color code for the atoms is gray for C, red for O, blue for N, and white for H.

(NylA), Ahx-linear-dimer (Ald) hydrolase (NylB), and endotype Ahx-oligomer hydrolase (NylC).¹¹ Among the identified nylon-oligomer hydrolases, NylB has recently been the target of extensive investigations.^{13–19} The NylB enzyme has been shown to possess not only a degrading activity of amide compounds, for example, $\text{NH}_2-(\text{CH}_2)_5-\text{CO}-\text{NH}-(\text{CH}_2)_5-\text{COOH} + \text{H}_2\text{O} \rightarrow 2\text{NH}_2-(\text{CH}_2)_5-\text{COOH}$ but also an esterase activity on various carboxyl esters with different acyl chains, similar to other peptidases (e.g., serine proteases).^{13,14} X-ray structures of NylB revealed similarities with several enzymes of the penicillin-recognizing family of serine-reactive hydrolases, specifically, D-alanyl-D-alanine (DD) carboxypeptidase (DD-peptidase) from *Streptomyces* sp. R61, carboxylesterase (EstB) from *Burkholderia gladioli*, and Class C β -lactamase.^{15–19} The crystal structure also evidenced that the binding site of an Ald substrate includes a catalytic triad Ser112/Lys115/Tyr215 and an oxyanion hole region (Figure 1a), located at positions close to those of the penicillin-recognizing enzymes.

However, in the NylB active site, a tyrosine residue Tyr170 exists, having no analog in any member of the penicillin-recognizing family for the serine-reactive hydrolases.¹⁶ The NylB substrate-binding X-ray structure showed that Tyr170 forms a hydrogen bond (H bond) with the Ald substrate near the Ser112/Lys115/Tyr215 catalytic triad (Figure 1a).

Concerning its functional role, a site-directed mutagenesis of Tyr170 to a phenylalanine exhibited a decrease in Ald degradation activity to 1.4% with respect to the wild type, indicating that Tyr170 is directly involved in the degradation process.¹⁷ Moreover, a comparison between substrate-binding/-unbound NylB crystal structures evidenced an induced-fit motion of Tyr170.¹⁷ This indicates that Tyr170 enhances Ald binding, a conclusion supported also by our former simulations.²⁰ However, no clear understanding of the Tyr170 catalytic role and of the structure–function relationships of the catalytic triad has been obtained to date. To shed light on these issues, we studied the catalytic mechanisms of Ald degradation by NylB-type nylon-oligomer hydrolase using hybrid QM/MM dynamics in combination with free-energy sampling^{21–34} for the activated processes.

The equilibrium structure of the active site of the wild-type (WT) NylB in a substrate-binding state was determined via QM/MM unbiased dynamics. Results are reported in Table 1 as averages over ~ 8 ps at 300 K. For comparison, the corresponding parameters of the NylB X-ray structures¹⁸ and structurally similar enzymes of the penicillin-recognizing family of the serine-reactive hydrolases^{35–39} are included. The calculated distances for the WT NylB with the Ald substrate show a rather good agreement with the available NylB crystal structures, providing support to our approach. One exception is the distance of Lys115- N_ϵ and Tyr215- O_η . Such a discrepancy might be due to S112A mutation effects on the Ald-binding X-ray structure to inactivate the enzyme (details in the Supporting Information).

A comparison with other penicillin-recognizing enzymes shows that the H-bond pattern in the catalytic triad and oxyanion hole of the substrate-binding NylB is basically similar to those enzymes (Table 1). In the NylB catalytic triad, the nucleophile Ser112- O_γ forms H bonds with both Lys115- N_ϵ and Tyr215- O_η . The distance between the side chains of Lys115 and Tyr215 (~ 3.7 Å) is slightly above the typical H bond,⁴⁰ and the estimation of H nuclear magnetic resonance (NMR) chemical shift indicates that no H bonds exist between these residues. (See the Supporting Information.) The backbone N atoms of Ser112 and Ile345 also form H bonds with the amide oxygen of the Ald substrate. At variance with the common features discussed so far, NylB exhibits a structural diversity in the substrate binding. In our simulated NylB, the Ald amide binds to Tyr170- O_η via a H bond (Table 1).

This hydrogen bonding leads to two structural differences around the substrate amide group between NylB and the other penicillin-recognizing enzymes. The first one stems from the absence of any H bond of the amide of the Ald substrate with side chains of the NylB catalytic triad (Table 1). Conversely, the substrate-binding X-ray structures of the DD-peptidase and the class C β -lactamase showed a clear binding of the substrate to the side chains of their catalytic triads.^{35,37} The second one stems from the distances of the carbonyl O of the substrate from the oxyanion hole region: These distances in the WT

Table 1. Main Geometrical Parameters for the Active Site of NylB, R61 DD-Peptidase, EstB Carboxylesterase, and Class C β -Lactamase^a

Δ /deg	NylB QM/MM substrate-binding (WT) ^c	NylB X-ray substrate-binding (S112A) ^b	NylB X-ray substrate-unbound (WT) ^c	R61 DD-peptidase substrate-binding (WT cross-linking) ^d	R61 DD-peptidase X-ray substrate-unbound (WT) ^e	EstB carboxyl-esterase substrate-binding (WT tetrahedral intermediate analog) ^f	EstB carboxyl-esterase X-ray substrate-unbound (WT) ^g	class C β -lactamase X-ray substrate-binding (S64G) ^h	class C β -lactamase X-ray substrate-unbound (WT) ⁱ
S112-O _γ -K115-N _ε [Å]	2.72 ± 0.11	none	2.60	4.08	4.13/5.30	3.18	2.98	none	2.82
S112-O _γ -Y215-O _η [Å]	2.88 ± 0.17	none	2.90	3.17	3.76/3.68	2.96	2.53	none	2.93
K115-N _ε -Y215-O _η [Å]	3.74 ± 0.23	2.60	3.71	3.27	2.76/3.21	2.73	2.69	2.91	3.33
Y170-O _γ -Ald-N [Å]	3.16 ± 0.19	3.08/3.04	none	none	none	none	none	none	none
S112-N-Ald-O [Å]	3.23 ± 0.21	2.96/3.03	none	2.72	none	2.92	none	2.80	none
I345-N-Ald-O [Å]	2.97 ± 0.14	2.81/2.86	none	2.79	none	2.66	none	2.86	none
S112-O _γ -Ald-C [Å]	3.01 ± 0.16	none	none	2.91	none	1.64	none	none	none
Ald-N-Ald-C [Å]	1.35 ± 0.02	1.33	none	1.40	none	none	none	none	none
S112-O _γ -Ald-C-Ald-O [deg]	87.23 ± 5.42	none	none	93.10	none	109.34	none	none	none

^aThe corresponding amino acid residues constituting the catalytic triad and oxyanion hole region in the latter three enzymes are given in Table S1 in the Supporting Information. ^bPDB entry is 2ZMA (1.51 Å resolution), the S112A mutant with Ald (two conformations). ^cPDB entry is 1IKG (1.90 Å resolution) including cross-linking of Lys65 and Tyr159, complex with glycyl-L- α -amino- ϵ -pimelyl-D-Ala-D-Ala fragment. ^dPDB entry is 3PTE (1.60 Å resolution) including two conformations for Lys65 and Tyr159. ^ePDB entry is 1CJ9 (1.80 Å resolution), the complex with diisopropyl-fluorophosphate. ^fPDB entry is 1KVL (1.53 Å resolution). ^gPDB entry is 1KE4 (1.72 Å resolution).

NylB are longer by 4–19% than those in the other penicillin-recognizing enzymes (Table 1), and the net effect is that the position of the Ald amide group toward the nucleophile is slightly altered in the NylB active site. As shown in Table 1, the distance Ser112-O_γ-Ald-C and angle Ser112-O_γ-Ald-C-Ald-O are 3 Å and 87°, respectively, whereas in the DD-peptidase case the corresponding geometrical parameters are 2.9 Å and 93°. Although all of these values are close to the energetically preferred distance and angle reported in ref 41 (~3 Å and ~100–110°), the difference between NylB and the DD-peptidase indicates that the involvement of Tyr170 in the enzyme–substrate interactions in NylB leads to an alternative stable state for the substrate binding in the amide bond cleavage.

To get a deeper insight into the Ald amide bond cleavage by NylB and the functional interactions between the catalytic triad and Tyr170, we investigated the cleavage reaction of the Ald amide bond in acylation via QM/MM dynamical simulations coupled to metadynamics.^{28–30} As reaction coordinates (RCs), two distances were selected: (1) the distance between Ser112-O_γ and the Ald carboxyl carbon and (2) the distance between the Ald amide nitrogen and carboxyl carbon. The results are summarized in Figures 2 and 3 in terms of the free-energy

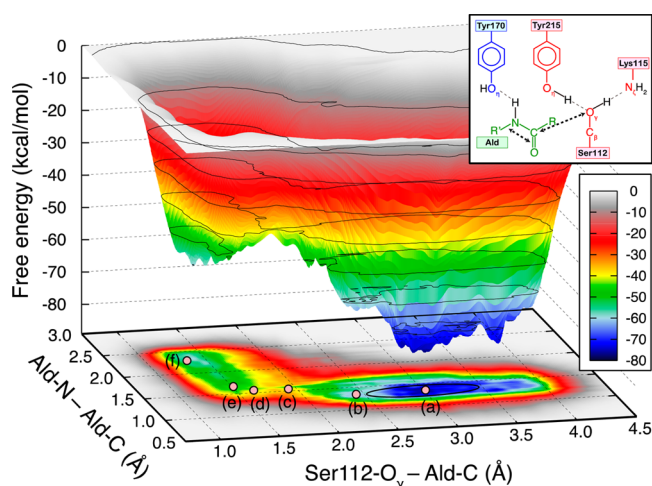


Figure 2. Free-energy surface as a function of the Ser112-O_γ-Ald-C and the Ald-N-Ald-C distances. The circle in the contour map shows the area to which the system is thermally accessible by standard unbiased dynamics.

surface (FES) as a function of the two RCs and key configurations characterizing the cleavage reaction.

The cleavage mechanism of the Ald amide bond in acylation of NylB is as a two-step reaction from the viewpoint of FES. The first step occurs along the reaction path joining the initial local minimum (a) to the adjacent local minimum (b) of the FES (Figure 2). The initial minimum is located in a deep basin, where the enzyme–substrate complex is in thermal equilibrium and fluctuates inside the area enclosed by a circle in Figure 2. The corresponding representative structure around the active site is sketched in Figure 3a. The FES local minimum (a) corresponds to a remarkable stabilization of the Ald substrate binding, induced by the unique H-bond interactions of the Ald amide group with Tyr170 and the oxyanion hole region in NylB. The subsequent local minimum (b) is positioned in a shallow basin, energetically located at ~14 kcal/mol above (a).

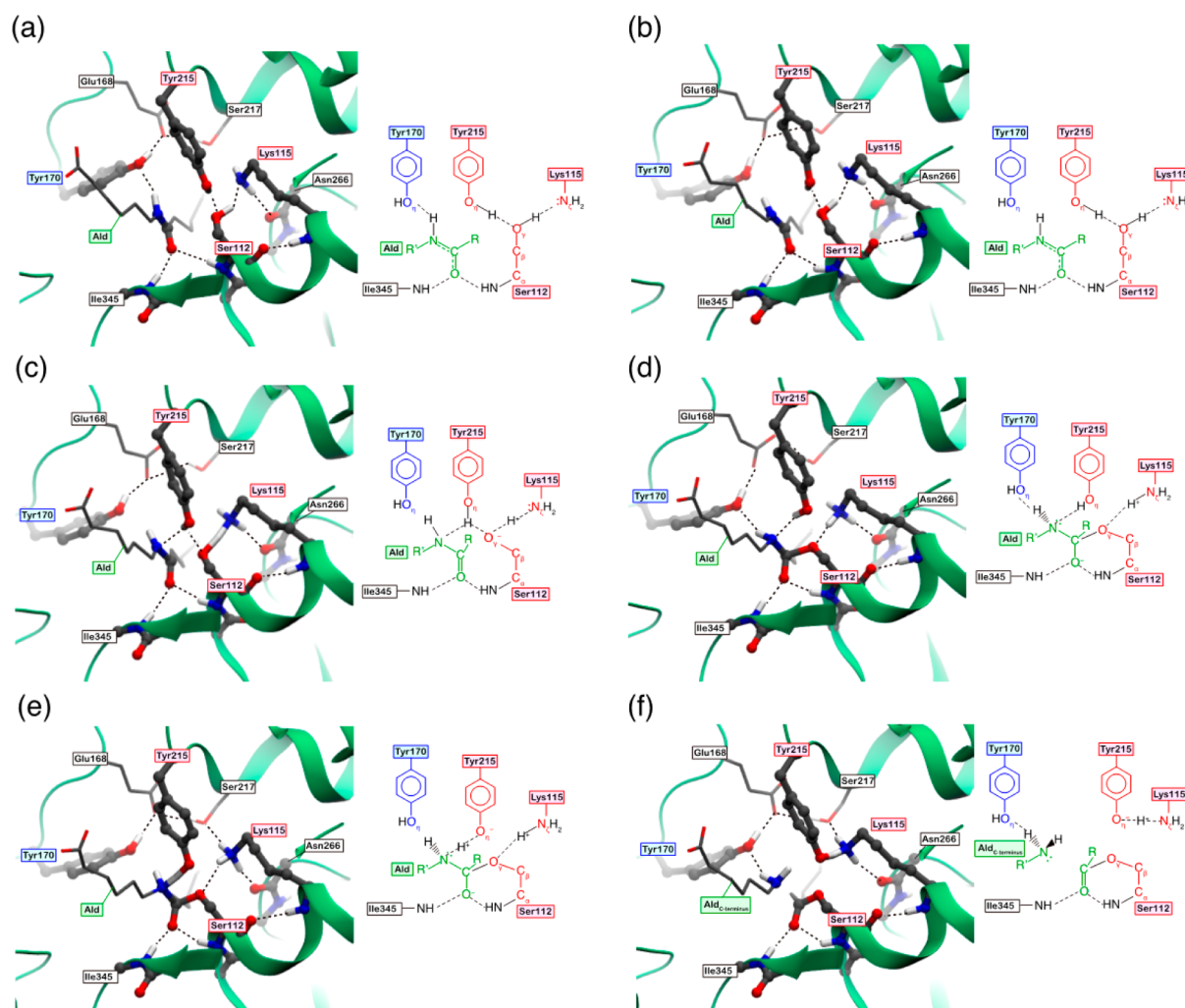


Figure 3. Snapshots of the structures of the various intermediate states realized by the system during the acylation. The labels a–f refer to the points marked by dots in Figure 2. The most relevant atoms are drawn as balls and sticks, and H bonds are indicated by dashed lines.

As shown in Figure 3b, the corresponding structure shows that Tyr170 is detached from the Ald substrate and the amide group slides into a pocket formed by the oxyanion hole.

The second reaction step is the cleavage of the amide bond and the formation of the acyl-enzyme, along the reaction path from (b) to (f) in Figure 2. This is the rate-limiting step and is characterized by a free-energy barrier, from (b) to (c), of ~ 21 kcal/mol, in quite good agreement with the reported experimental value of ~ 17 kcal/mol estimated from the transition-state theory assuming the experimental k_{cat} value of 3.2 s^{-1} for WT NylB.¹⁸ This value is also in the range of activation barriers for the acylation step in serine-reactive hydrolases (18–26 kcal/mol).^{42–47} Hence, we can infer that the reaction-ready state of NylB is likely to correspond to the substrate-binding state of other serine-reactive hydrolases.

The resulting reaction mechanism is mainly the one typical of a catalytic triad, but it involves the crucial participation of Tyr170 (Figure 3, forms c–f). A proton abstraction from the nucleophile of Ser112- O_γH is triggered by Lys115 (Figure 3c). Simultaneously, the nucleophilic attack of Ser112- O_γ^- on the amide carbon occurs, leading to the formation of a tetrahedral intermediate (Figure 3d). This intermediate is then stabilized by the oxyanion hole consisting of Ser112 and Ile345 backbone

NHs as well as Lys115- $\text{N}_\zeta\text{H}_3^+$, as in the conventional mechanism of the serine-reactive hydrolases.^{1–9}

However, the functional interaction of Tyr170 with the catalytic triad leads to a remarkable stereoelectronic effect on the tetrahedral intermediate. As shown in Figure 3d, the formation of a Ser112- O_γ –Ald C single bond after the nucleophilic attack leads to a conversion of the N–C partial double bond in the amide into a single bond. In contrast, Tyr170 works as a H-bond acceptor for the Ald amide N, forming another H bond with the adjacent Glu168 residue. Simultaneously, Tyr215 of the catalytic triad acts as a H-bond donor for the Ald amide. As a result of these three interactions, the NH moiety in the amide group takes an antiperiplanar configuration with respect to the Ser112- O_γ –Ald C bond, thus assuming a stereochemically favorable conformation to accept the proton from Tyr215- O_ηH without requiring large positional changes. In addition to this stereochemical effect of Tyr170 on the tetrahedral intermediate, electrostatic interactions of Tyr215- O_ηH with a positively charged Lys115- $\text{N}_\zeta\text{H}_3^+$, located between O_η and N_ζ at distances of ~ 3.5 Å, contribute in a concerted way to the proton transfer from Tyr215- O_η to the amide N. This, in turn, leads to the cleavage of the Ald amide bond and the formation of the acyl-enzyme (Figure 3e,f). Although the reaction path was sampled until the departing

amine (6-aminohexanoate, i.e., carboxyl residue in Ald) reached the border of the QM cell, it is noteworthy that this leaving amine is caught by Tyr170 via a H bond, suggesting that the release of the amine is accompanied by a Tyr170 structural change. Furthermore, Tyr215-O_η⁻ forms a strong H bond with Lys115-N_εH₃⁺ in the acyl-enzyme, analogous to the X-ray structure of covalent acyl enzyme complex of the DD-peptidase.⁴⁸

Concerning the functional role of Tyr170 in the catalytic-triad-driven NylB reaction, our calculations indicate that Tyr170 enhances the proton transfer from Tyr215 (general acid) to the Ald amide N in the tetrahedral intermediate. This is consistent with the results of site-directed mutagenesis, where a conversion of Tyr170 to a phenylalanine decreases the Ald degradation activity with respect to the WT.¹⁷ The importance of such functional role stems from the stereoelectronic requirements for an efficient breakdown of the tetrahedral intermediate, an important issue in other serine proteases.^{4,43,49–51} In both cases, NylB and serine protease, a nucleophile and a general acid in a catalytic triad, are located on the same side of the scissile amide bond. When the nucleophile attacks the amide bond, the NH of the amide group rotates around the scissile bond axis to accept the H⁺ from the acid, provided that the acid does not undergo large displacements. In the case of NylB, this rotation proceeds according to the nucleophilic attack by functional interactions among the Tyr170 residue and the catalytic triad along three steps: (1) the distance between Tyr215-O_η and the Ald amide N decreases during the nucleophilic attack (Figure 3b,c), (2) Tyr215 starts pointing its O_η-H bond toward the amide N to form an H bond (Figure 3c), and (3) Tyr170 moves back toward the amide N, forming an H bond by rotating the NH moiety around the scissile bond toward Tyr170 (Figure 3d). This mechanism, driven by the concerted action of the Tyr170 residue and the NylB catalytic triad, could efficiently induce the disruption of the tetrahedral intermediate because it does not require large displacements of the involved amino acid residues. This represents a significant novelty in the structure–function relationships of NylB active site, where the formation of a preorganized environment suited to realize conventional catalytic-triad-driven mechanisms was already evidenced for other proteins.^{52,53}

The importance of the Tyr170 functional role can also be rationalized by comparison with a class C β-lactamase sharing a structural similarity with the Ald hydrolase in the penicillin-recognizing family of the serine-reactive hydrolases. In the class C β-lactamase, the tyrosine residue corresponding to Tyr215 forms two H bonds with two nearby lysine residues, considered to be a general base/acid.⁵⁴ Concerning its proton donation to the substrate amide, such positively charged lysine residues could increase the acidity of the tyrosine. Conversely, a cooperation of the positively charged lysine would be negligible in NylB because only one Lys115 residue is present near Tyr215 and no direct H bond to Tyr215 exists. In this situation, the acidity of Tyr215 does not increase significantly, thus hindering the proton transfer.

In summary, we studied, via reactive simulations, the acylation of the Ald degradation by the NylB-type nylon-oligomer hydrolase. The depicted reaction mechanism evidence two crucial functional roles of a unique residue, Tyr170: (1) a stabilization of the substrate-binding state and (2) an enhancement in the proton-donation from a general acid to the substrate amide N in the tetrahedral intermediate. The free-energy barrier for the rate-limiting step of the acylation is ~21 kcal/mol, providing also a quantitative information to the inspected reaction. Moreover,

this discloses environmentally friendly and industrially appealing ways of disposing nylon wastes.

■ COMPUTATIONAL METHODS

The structural model of the WT NylB was constructed from the substrate-binding X-ray structure of the S112A mutant (PDB ID: 2ZMA) after replacing S112A with serine. The protonation states of Lys115 and Tyr215 in the catalytic triad were set to be neutral, corresponding to a general base for the nucleophile of Ser112. The system was thermalized at 300 K by classical molecular dynamics with AMBER ff99 force field.²¹ The equilibrated system was then used for hybrid QM/MM simulations, where the QM driver was the density functional theory framework^{22,23} in the generalized gradient approximation on the exchange and correlation functional after Hamprecht, Cohen, Tozer, and Handy (HCTH).²⁴ A Car–Parrinello molecular dynamics^{25,26} approach was used with a Hamiltonian QM/MM coupling scheme according to ref 27. The QM region includes the Ser112/Lys115/Tyr215 catalytic triad, Tyr170, and their surrounding residues (Figure 1b). The reaction pathways were sampled by the metadynamics approach.^{28–34} Additional computational details are provided in the Supporting Information.

■ ASSOCIATED CONTENT

§ Supporting Information

Further details about the computational setup, the reaction coordinates, the scatterplot of sampled points in reaction coordination space, and free-energy analyses. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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