

Three-dimensional Structure of Nylon Hydrolase and Mechanism of Nylon-6 Hydrolysis^{*S}

Received for publication, November 8, 2011, and in revised form, December 16, 2011. Published, JBC Papers in Press, December 19, 2011, DOI 10.1074/jbc.M111.321992

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Background: Biodegradation of polyamides is important from the industrial and environmental point of view.

Results: We identified the catalytic residue of nylon hydrolase as Thr-267 and enhanced the protein thermostability by 36 °C ($T_m = 88$ °C) by introducing mutations at the subunit interfaces of tetramer structure.

Conclusion: We revealed the mechanism of nylon-6 hydrolysis.

Significance: We established an approach to biodegrade polymeric nylon-6.

We performed x-ray crystallographic analyses of the 6-aminohexanoate oligomer hydrolase (NylC) from *Agromyces* sp. at 2.0 Å-resolution. This enzyme is a member of the N-terminal nucleophile hydrolase superfamily that is responsible for the degradation of the nylon-6 industry byproduct. We observed four identical heterodimers (27 kDa + 9 kDa), which resulted from the autoprocessing of the precursor protein (36 kDa) and which constitute the doughnut-shaped quaternary structure. The catalytic residue of NylC was identified as the N-terminal Thr-267 of the 9-kDa subunit. Furthermore, each heterodimer is folded into a single domain, generating a stacked $\alpha\beta\beta\alpha$ core structure. Amino acid mutations at subunit interfaces of the tetramer were observed to drastically alter the thermostability of the protein. In particular, four mutations (D122G/H130Y/D36A/E263Q) of wild-type NylC from *Arthrobacter* sp. (plasmid pOAD2-encoding enzyme), with a heat denaturation temperature of $T_m = 52$ °C, enhanced the protein thermostability by 36 °C ($T_m = 88$ °C), whereas a single mutation (G111S or L137A) decreased the stability by ~ 10 °C. We examined the enzymatic hydrolysis of nylon-6 by the thermostable NylC mutant. Argon cluster secondary ion mass spectrometry analyses of the reaction products revealed that the major peak of nylon-6 (m/z

10,000–25,000) shifted to a smaller range, producing a new peak corresponding to m/z 1500–3000 after the enzyme treatment at 60 °C. In addition, smaller fragments in the soluble fraction were successively hydrolyzed to dimers and monomers. Based on these data, we propose that NylC should be designated as nylon hydrolase (or nylonase). Three potential uses of NylC for industrial and environmental applications are also discussed.

Nylons are synthetic polymers that contain recurring amide groups (R-CO-NH-R') as integral parts of their main polymer chains. The high strength, elasticity, abrasion resistance, chemical resistance, and shape-holding characteristics of nylons over wide temperature ranges make these polymers suitable for the production of fibers and plastics. Currently, the worldwide production of nylons is estimated to be three to four million tons per year. Nylons tend to be partially crystalline, and the degree of crystallinity affects the properties of nylons, such as the melting points, strength, and rigidity. Two forms of crystals (α and γ) have been reported (1). In the α -form, each polymer chain is stabilized by hydrogen bonds with adjacent chains aligned in an antiparallel orientation, whereas the chains are parallel in the γ -form. The α -form of nylon-6 is generally more stable (1).

Nylon-6 is produced by the ring cleavage polymerization of ϵ -caprolactam and consists of more than 100 units of 6-aminohexanoate (Ahx).⁴ However, during the polymerization reac-

* This work was supported in part by a grant-in-aid for scientific research (Japan Society for Promotion of Science) and grants from the Global Center of Excellence Program and the Japan Aerospace Exploration Agency project.

^S This article contains supplemental Tables S1–S4 and Figs. S1–S5.

The atomic coordinates and structure factors (code 3AXG) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (<http://www.rcsb.org/>).

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⁴ The abbreviations used are: Ahx, 6-aminohexanoate; N-tn, N-terminal nucleophile; NylA, 6-aminohexanoate cyclic dimer hydrolase; NylB, 6-aminohexanoate-linear dimer hydrolase; NylC, 6-aminohexanoate oligomer hydrolase, or nylon hydrolase; r.m.s., root mean square; DmpA, L-amino-peptidase D-Ala-esterase/amidase; BapA, β -peptidyl aminopeptidase; OAT, ornithine acetyltransferase; SIMS, secondary ion mass spectrometry; S¹¹¹, G111S; G¹²², D122G; Y¹³⁰, H130Y; A¹³⁷, L137A; M²²⁵, V225M; G¹²²Y¹³⁰, D122G/H130Y; G¹²²Y¹³⁰A¹³⁷, D122G/H130Y/D36A; G¹²²Y¹³⁰Q²⁶³, D122G/H130Y/E263Q; G¹²²Y¹³⁰M²²⁵, D122G/H130Y/V225M; S¹¹¹G¹²²Y¹³⁰, G111S/D122G/H130Y; G¹²²Y¹³⁰A¹³⁷, D122G/H130Y/L137A; G¹²²Y¹³⁰A¹³⁷Q²⁶³, D122G/H130Y/D36A/E263Q; S¹¹¹G¹²²Y¹³⁰A¹³⁷, G111S/D122G/H130Y/L137A.

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tion, some molecules fail to polymerize and remain oligomers, whereas others undergo head-to-tail condensation to form cyclic oligomers. We have isolated bacterial strains that can degrade Ahx oligomers, which are by-products of nylon-6 production, and use these oligomers as their sole carbon and nitrogen source (2–6). Previous biochemical studies have revealed that three enzymes, NylABC, are responsible for the degradation of the by-products produced during nylon-6 manufacturing. The Ahx cyclic dimer hydrolase (NylA; EC 3.5.2.12), a member of the amidase signature hydrolase family, specifically hydrolyzes one of the two equivalent amide bonds in the Ahx cyclic dimer, generating an Ahx linear dimer (7). The Ahx dimer hydrolase (NylB; EC 3.5.1.46), a member of the penicillin-recognizing family of serine reactive hydrolases, hydrolyzes Ahx oligomers by an exo-type mode (8–12). The Ahx oligomer hydrolase (NylC; EC 3.5.-.-) degrades Ahx cyclic and linear oligomers with a degree of polymerization greater than three by an endo-type mode (5, 13–15).

NylC has been found in *Arthrobacter* (pOAD2 plasmid-encoded NylC; NylC_{p2}), *Agromyces* (NylC_A), and *Kocuria* (NylC_K) (5). These enzymes are each encoded by a single gene that corresponds to a polypeptide chain of 355 amino acids. However, the post-translational cleavage of the nascent polypeptide between Asn-266 and Thr-267 generates a 27-kDa (α)-subunit and a 9-kDa (β)-subunit (5, 14). The autoprocessing of the inactive precursor to an active enzyme is a specific feature of the N-terminal nucleophile (N-tn) hydrolase family (16–33). NylC_A and NylC_K have 5 and 15 amino acid substitutions, respectively, relative to the NylC_{p2} sequence (supplemental Fig. S1), and both have 10–20 °C higher thermostability than NylC_{p2} (5).

Recently, we reported the crystallization conditions of NylC_A that are suitable for x-ray crystallographic analyses (34). In this study, we performed x-ray crystallographic analyses of NylC_A, examined its structural/evolutional relationship to proteins registered in the Protein Data Bank (PDB), and estimated the residues responsible for the catalytic function and autoprocessing. We also analyzed the effect of amino acid mutations on the thermostability of NylC. Furthermore, we investigated the possibility of using the thermostable NylC mutant to degrade polymeric nylon-6 at high temperatures, which is expected to increase the reactivity of the polymer.

EXPERIMENTAL PROCEDURES

DNA Preparation and Site-directed Mutagenesis

The plasmids pSKFC4 (NylC_{p2}), pSKRC4 (NylC_A), and pSKKC4 (NylC_K) contained 1.1-kb genes flanked by BamHI and PstI restriction sites, which were cloned into the expression vector pBluescript II SK(+) (Stratagene, La Jolla, CA.) (5). *Escherichia coli* JM109-competent cells were prepared by the CaCl₂ method (35) and stored at –80 °C before use. To introduce the D36A, G111S, D122G, H130Y, L137A, V225M, and E263Q mutations into the NylC_{p2} sequence, site-directed mutagenesis was performed using the PrimeSTAR mutagenesis kit (Takara Bio Inc.) with the primers listed in supplemental Table S1. The plasmids that contained the 1.1-kb fragments with mutated NylC_{p2} were isolated from transformed *E. coli* JM109 cells.

DNA sequencing confirmed that the desired mutations were introduced into the wild-type *nylC*_{p2} sequence.

Cultivation, Enzyme Purification, and Enzyme Assay

Cultivation and purification from *E. coli* clones were performed as reported previously (34). In the NylC activity assays, the enzyme solution (0.1 ml) was mixed with an Ahx cyclic oligomer solution (0.9 ml, 4 mg ml^{–1} Ahx cyclic oligomer in 20 mM phosphate buffer, pH 7.3, 10% glycerol (buffer A)) and incubated at 30 °C (standard assay condition). An increase in the concentration of the amino group was determined using trinitrobenzene sulfonic acid (5). Kinetic studies were performed under standard assay conditions, with the exception of the different Ahx cyclic oligomer concentrations used.

Nylon Degradation Tests

Nylon-6 that was mechanically disintegrated to a powder was a generous gift from Toyobo Co. (Tsuruga, Japan). To analyze the particle sizes, the nylon sample was magnified 80-fold with a scanning electron microscope (Hitachi, model TM-1000). The diameter (maximum length) of 157 particles observed in five electron microphotographs was measured. The diameters ranged from 68 to 720 μ m (average \pm S.D., 270 \pm 140 μ m). To increase the reactivity of the nylon sample, the nylon-6 powder (10 mg) was autoclaved in plastic tubes (Eppendorf Co.) containing buffer A (180 μ l) at 120 °C for 20 min prior to the initiation of the enzyme reaction. Hydrolytic reactions were initiated by the addition of the thermostable NylC_{p2}-G¹²²Y¹³⁰A³⁶Q²⁶³ mutant enzyme (1 mg ml^{–1}, 20 μ l) to each tube and incubated at 60 °C for 2 h. The gas cluster secondary ion mass spectrometry (SIMS) equipment (developed in the project “Development of System and Technology for Advanced Measurement and Analysis,” Japan Science and Technology Agency, 2006–2010) was used to analyze the reaction products (36). Enzyme reactions were performed in triplicate. The reaction products (containing both the insoluble and the soluble fractions) from the first tube were spotted onto a silicon plate (1-cm² square plate). Mass-to-charge ratio (*m/z*) ranges of 5,000–35,000 and 0–1,500 were analyzed. The solid fractions (~10 mg) that were recovered from the reaction products (the second tube) were washed with distilled water, lyophilized, and dissolved in trifluoroethanol (0.2 ml). Subsequently, a fraction (0.02 ml) was spotted onto a silicon plate (1-cm² square plate). The *m/z* range of 1,500–15,000 was analyzed. The soluble fraction (1 μ l) from the third tube was spotted onto a thin layer plate and developed by a solvent mixture containing 1-propanol:water:ethyl acetate:ammonia (24:12:4:1.3). The degradation products were detected by spraying the plate with a 0.2% ninhydrin solution (in *n*-butyl alcohol saturated with water) (5).

Crystallographic Analysis

Crystallization and Diffraction Data Collection—For analysis of NylC_A, plate-like crystals (0.8 \times 0.4 \times 0.3 mm) were obtained by the sitting-drop vapor diffusion method. Droplets were prepared by mixing 2 μ l of purified NylC_A solution (10 mg ml^{–1} protein in buffer A) and 2 μ l of reservoir solution (1.0 M sodium citrate as a precipitant in 0.1 M HEPES buffer (pH 7.5), 0.2 M NaCl) and were equilibrated against 100 μ l of reservoir

TABLE 1
Data collection statistics and refinement statistics

Values in parentheses are for the outer resolution shell.

	Native (NylC _A)	K ₂ PtCl ₄ derivative (NylC _A)
Data collection statistics		
Space group	I222	I222
Unit cell (Å)		
<i>a</i>	155.86	155.99
<i>b</i>	214.45	214.72
<i>c</i>	478.80	477.82
Wavelength (Å)	1.0000	1.0000
Resolution (outer shell) (Å)	30-2.00 (2.07-2.00)	30-2.20 (2.28-2.20)
Total reflections	1,789,588	2,786,883
Unique reflections (outer shell)	501,977 (38,621)	398,011 (36,644)
Completeness (outer shell) (%)	94.3 (73.0)	99.0 (91.7)
<i>R</i> _{merge} (outer shell) (%)	11.1 (25.4)	6.4 (46.2)
<i>I</i> / <i>σ</i> (<i>I</i>) (outer shell)	10.6 (2.32)	26.1 (2.87)
Multiplicity	3.6 (2.0)	7.0 (4.8)
Refinement statistics		
Resolution range (outer shell) (Å)	50-2.00 (2.07-2.00)	
<i>R</i> _{work} ^a (outer shell) (%)	22.3 (31.9)	
<i>R</i> _{free} ^a (outer shell) (%)	25.2 (33.5)	
No. of protein atoms	37,819	
No. of water molecules	3,006	
r.m.s. deviations from ideal values		
r.m.s. bond distances (Å)	0.006	
r.m.s. bond angles (°)	0.9	
Dihedral angles (°)	22.3	
Improper angles (°)	0.74	
Ramachandran plot		
Favored (%)	96.2	
Allowed (%)	99.9	
Outliers	4	

$$^a R = \frac{\sum_{hkl} |F_{\text{obs}}| - k|F_{\text{calc}}|}{\sum_{hkl} F_{\text{obs}}}^{-1}; k = \text{scaling factor.}$$

solution at 10 °C for 1 week. The crystal belonged to the space group I222, with unit cell parameters *a* = 155.86 Å, *b* = 214.45 Å, and *c* = 478.80 Å (Table 1).

For native crystals, the crystals were soaked for 24 h in a cryoprotectant solution (1.0 M sodium citrate, 0.1 M HEPES (pH 7.5), 0.2 M NaCl, 25% glycerol). Heavy atom derivatives of NylC_A were prepared by soaking the crystals for 72 h in cryoprotectant solution containing 5 mM K₂PtCl₄. Cryo-cooling was performed by blowing cold nitrogen steam onto the crystals at 100 K.

The diffraction data sets were collected at the SPring-8 (Hyogo, Japan) beamline BL38B1 equipped with a Rigaku Jupiter CCD detector system. The following parameters were chosen for data collection: wavelength, 1.0000 Å; crystal to detector distance, 180 mm; oscillation range per image, 0.5° (for native and platinum derivative). Indexing, integration, and the scaling of reflections were performed using the HKL2000 program package (37). Diffraction data were collected from native NylC_A crystals and from the K₂PtCl₄-derivative to resolutions of 2.00 Å and 2.20 Å, respectively (Table 1).

Phase Determination, Model Building, and Crystallographic Refinement—The NylC structure was determined by the single wavelength anomalous diffraction method using the K₂PtCl₄ derivative data. The platinum substructure for the derivative crystal was solved at a resolution of 2.2 Å by the program BnP (38) using the anomalous signal of the platinum atoms. Initial phase parameters were determined using the program SHARP (39). The electron density map was automatically traced with ARP/wARP. Models for the untraced regions, with the exception of the amino acids at positions 1–18 and 261–266, were constructed by manual model building using XFIT (40). Rigid body refinement was performed using the coordinates of the

initial model to fit the unit cell of the NylC_A crystal followed by positional and *B*-factor refinement with the program CNS (41). Non-crystallographic symmetry restraints were applied throughout refinement with the exception of the residues 25–40 and 131–137 in which significant structural differences among chains were detected.

After several cycles of manual model rebuilding by XFIT, *R*-factor and *R*_{free} were determined to be 18.1 and 19.5%. No residue was found in the outlying regions. The results of the crystal structure analysis are summarized in Table 1. The figures of the three-dimensional structural models were generated with the program MolFeat (version 3.6, FiatLux Co., Tokyo, Japan).

CD Analysis

Circular dichroism (CD) spectra at far-UV wavelengths (200–250 nm) were measured using a spectropolarimeter (JASCO, model J-720WI). A cuvette with a path length of 1 mm was used for far-UV CD measurements. The results are expressed as the mean residue molar ellipticity, $[\theta]$, which is defined as $[\theta] = 100 (\theta_{\text{obs}} - \theta_{\text{back}}) l^{-1} c^{-1}$, where θ_{obs} is the observed ellipticity in degrees, θ_{back} is the observed ellipticity in degrees in the absence of enzyme (as background), *c* is the molar concentration of the residue, and *l* is the length of the light path (in centimeters). The temperature was controlled at 25 °C with a JASCO PTC-348WI Peltier system. For thermal transition experiments, the temperature was shifted from 25 to 95 °C with a JASCO PTC-348WI Peltier system at 1 °C min⁻¹, and CD measurements were performed at 220 nm. The protein concentration used in the far-UV CD measurements was 0.1 mg ml⁻¹.

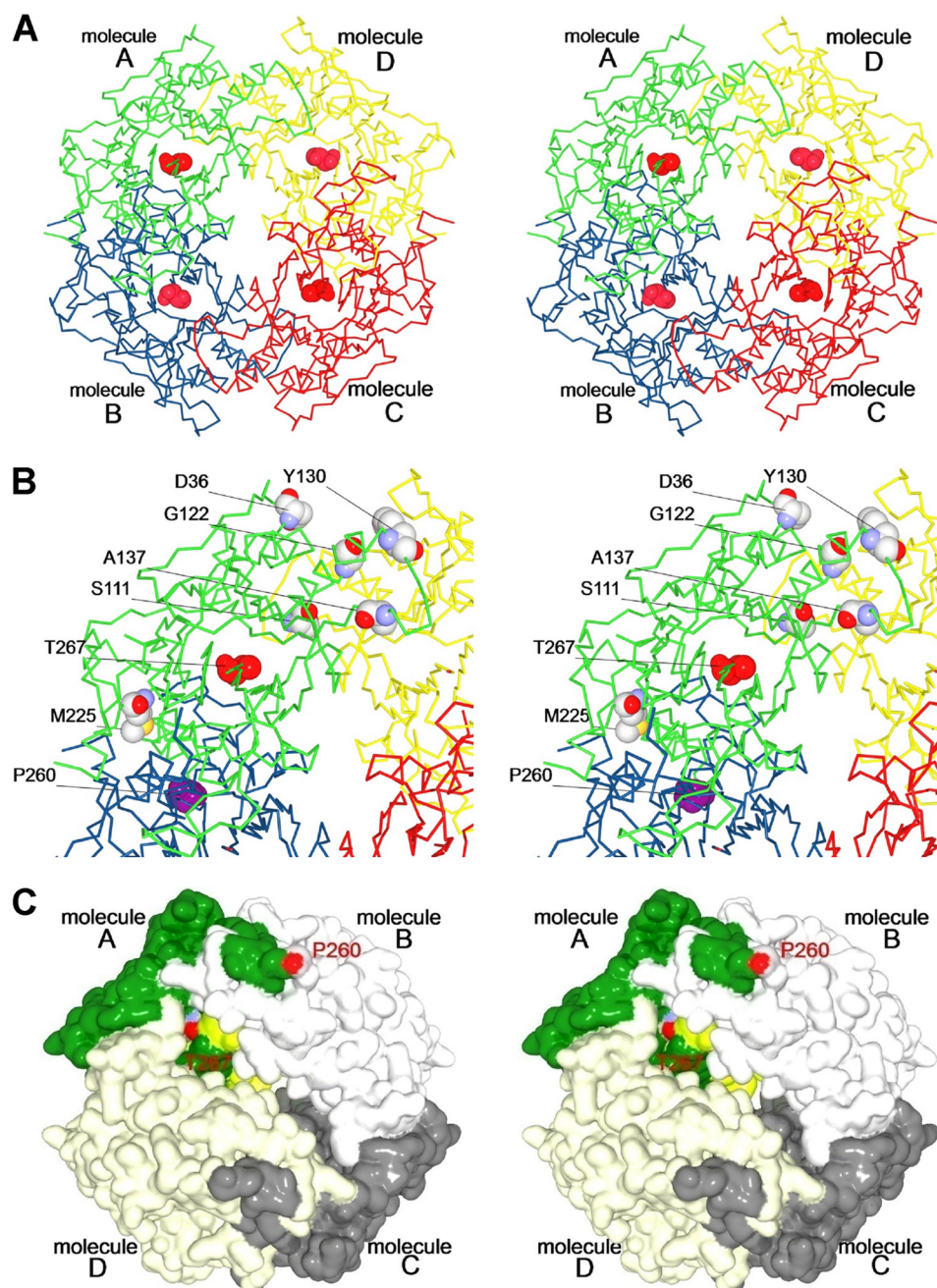


FIGURE 1. **Stereo view of quaternary structure of NylC_A.** A, the quaternary structure is shown, with different colors highlighting the individual molecules A (green), B (blue), C (red), and D (yellow). The catalytic residue Thr-267 (the N terminus of the β-subunit) is shown as a space-filling model (red). B, an enlarged view of molecule A and its interfaces with the adjacent molecules B and D. Six residues selected for mutagenesis (Asp-36 (Ala-36 in NylC_K), Ser-111, Gly-122, Tyr-130, Ala-137, and Met-225) are shown as space-filling models. Because of the poor electron density distribution of the C-terminal region in the α-subunit, including Glu-263 (Gln-263 in NylC_K), the adjacent Pro-260 is shown as a space-filling model (magenta). C, surface structure of NylC_A. The α-subunit and β-subunit in a single heterodimer (molecule A) are highlighted in dark green and light green, respectively. The other three heterodimers are shown in different shades of gray.

RESULTS AND DISCUSSION

Quaternary Structure

The asymmetric unit contains 15 molecules (molecules A–O), each composed of an α- and a β-subunit. One molecule (O) is part of a tetramer in which the four heterodimers are related by crystallographic 222 symmetry. Two molecules in dimer (M/N) form another tetramer that is related by a crystallographic two-fold axis, and the rest of the 12 molecules comprise three tetramers (A/B/C/D, E/F/G/H, and I/J/K/L) that are related by non-crystallographic 222 symmetries. The root mean square (r.m.s.) deviations

of the superimposed C_α atoms for all 15 molecules were calculated to be within the range of 0.13–0.38 Å by secondary structure matching (supplemental Table S2) (42), demonstrating that the overall structures are almost identical. Based on these results, we concluded that NylC_A adopts a doughnut-shaped quaternary structure in which four heterodimer molecules are mutually related by a perpendicular crystallographic and/or non-crystallographic two-fold axis (Fig. 1). Contacts at the A/B and C/D interfaces (3,121 Å²) were observed to be more extensive than those at the B/C and A/D interfaces (1,451 Å²).

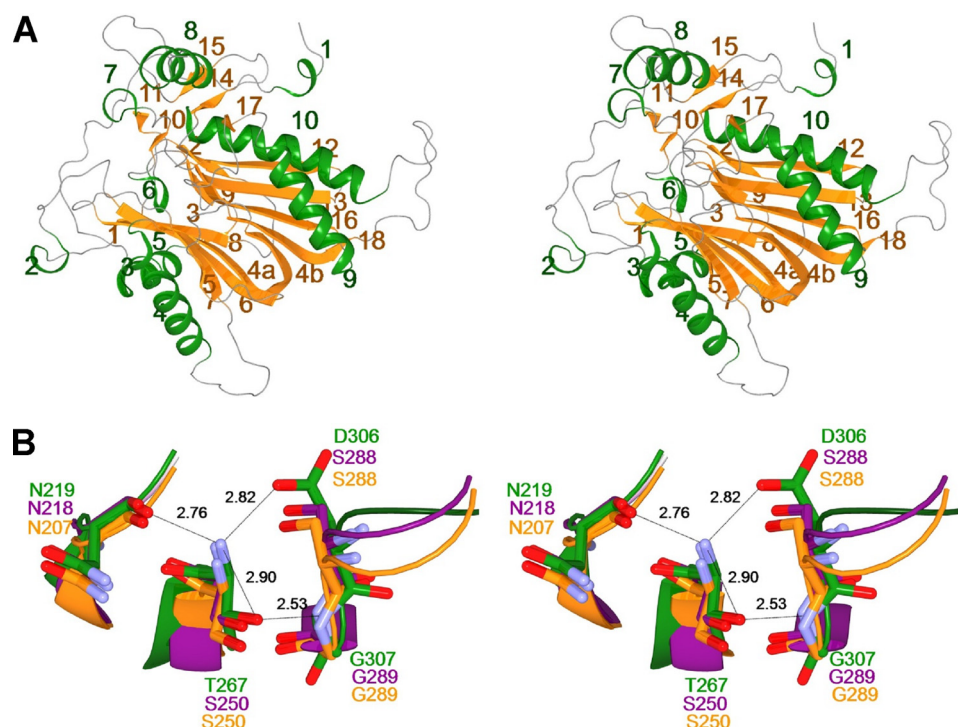


FIGURE 2. **Stereo view of subunit structure and catalytic center of NylC_A.** *A*, the overall structure of the heterodimer (molecule A) is shown as a *ribbon diagram*. Ten helices (H1–H10) and eighteen β -strands (S1–S18) are colored in *green* and *orange*, respectively. H1–H6 and H8–H10 are α -helices. H7 is a 3_{10} helix. *B*, the structure around the catalytic residue Thr-267 of NylC_A (*green*) is superimposed onto the structure of DmpA (PDB ID code, 1B65; *magenta*) and BapA (PDB ID code, 3N33; *orange*). Possible hydrogen bonds in NylC_A are indicated as *dotted lines* with the distances listed in *angstroms*.

Subunit Structure and Function Relationship with Other N-tn Family Enzymes

Each heterodimer contains 10 helices, H1–H10, which are α -helices with the exception of H7 (3_{10} helix), and 18 β -strands. The two subunits (α and β) obtained by intracleavage at Asn-266/Thr-267 are folded into a single domain, generating a stacked $\alpha\beta\beta\alpha$ core structure (Fig. 2A, supplemental Fig. S2). Namely, the central antiparallel β -sheet composed of five β -strands (β 3, β 4a, β 5, β 6, β 7) and another antiparallel β -sheet composed of five β -strands (β 4b, β 12, β 13, β 16, β 18) are packed against each other and connected by a common long β -strand (β 4). These two sheets are flanked on one side by the α -helices H4 and H5 and on the other by α -helices H9 and H10. The $\alpha\beta\beta\alpha$ fold is typically conserved throughout the N-tn hydrolase superfamily (16–33). However, our data demonstrate that NylC differs from most N-tn hydrolases in the directionality and connectivity of its secondary structure elements.

To search for proteins that are evolutionary related to NylC, we performed a homology search based on the NylC_A structure for PDB using the DALI program (43). The following four proteins (including one hypothetical protein) that exhibited high z-scores (z-score >14) were identified: L-aminopeptidase D-Ala-esterase/amidase from *Ochrobactrum anthropi* (DmpA; z-score = 26.3; PDB ID code: 1B65) (18), hypothetical D-aminopeptidase (z-score = 25.9; PDB ID code, 2DRH), β -peptidyl amino-peptidase from *Sphingosinella xenopeptidilytica* (BapA; z-score = 25.2; PDB ID code: 3N33) (20), and ornithine acetyltransferase from *Streptomyces clavuligerus* (OAT; z-score = 14.1; PDB ID code: 1VZ6) (21) (supplemental Table

S3). In contrast, the DALI z-scores of the other proteins in the PDB were determined to be below 8.

DmpA is an aminopeptidase that hydrolyzes peptides, with a preference for N-terminal residues in an L-configuration (L-Ala-Gly-Gly) and also amide or ester derivatives of D-Ala (19). BapA hydrolyzes β -oligopeptides and mixed β/α -oligopeptides with a β -amino acid residue at the N terminus (20). OAT is involved in the arginine biosynthetic pathway because it catalyzes the transfer of an acetyl group from N-acetylornithine to glutamate (21, 22). In contrast, NylC_{p2} hydrolyzes Ahx cyclic and linear oligomers (degree of polymerization >3) but has no detectable activity with the 66 peptides tested (supplemental Table S4), including D,L-Ala-Gly-Gly (14, 15). Thus, NylC exhibits distinct substrate specificity from that of DmpA/BapA/OAT.

DmpA/OAT are considered to have a unique fold (designated as DOM-fold), which is different from that of most N-tn hydrolases (17). Based on fundamental topology differences, Cheng and Grishin (17) proposed that the functional similarities between DmpA/OAT and the other N-tn hydrolases result from convergent evolution rather than from having a common evolutionary origin. The fold of NylC is similar to that of DmpA/BapA, and the structurally superimposable regions comprise 259 residues (DmpA) and 264 residues (BapA) (supplemental Table S3). The r.m.s. deviations of the superimposed C α atoms were calculated to be 2.0 Å (DmpA) and 2.0 Å (BapA) by secondary structure matching. However, the following major structural differences were found among the family enzymes.

In contrast to the single domain structure of NylC/DmpA/BapA, OAT is composed of two domains (supplemental Fig. S3).

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NylC is superimposable with OAT domain 1 located at positions 1–258. OAT domain 2 is located at the C-terminal region and is composed of 124 residues that have a successive order of secondary structure (β 10-H9-H10-H11- β 11- β 12-H12- β 13- β 14-H13). DmpA and BapA are longer than NylC at the C-terminal region of the β -subunit by 34 and 30 residues, respectively (supplemental Fig. S4).

Helix H8, which is located at the C-terminal region of the α -subunit, is unique in NylC. H8 and its linked loop region is flipped out to the surface of the adjacent molecules upon autoprocessing (Fig. 1C). In DmpA/OAT, the corresponding helix is absent. Furthermore, in BapA, the corresponding region (positions 236–245), which contains β 14 and a loop, is folded within the same subunit molecule without flipping out to the adjacent molecule.

The N-terminal 26 residues, including H1, generate unique structure specifically found in NylC. Moreover, in OAT, N-terminal 48 residues, including H1-H2- β 1- β 2, are absent (supplemental Fig. S4).

A loop flanked by β -strand β 18 and helix H10 in NylC (positions 317–333) spatially share different positions in Dmp (positions 299–318) and in BapA (positions 299–316). In OAT, the corresponding loop is absent, and β 9 and H18 are combined with a single residue (Val-235) at the junction (supplemental Fig. S4).

In NylC, a long β -strand (β 4a, β 4b) constitutes a part of the central two β -sheets (Fig. 2A). However, in DmpA/BapA/OAT, the corresponding β -strand is divided into two β -strands (e.g. β 2, β 3 in DmpA), which are connected by a short loop and constitute each β -sheet (supplemental Fig. S4).

Residues Responsible for Autoprocessing and Catalytic Function

In DmpA, autoprocessing occurs between Gly-249 and Ser-250. Additionally, the newly generated N-terminal Ser-250 of the β -subunit plays the roles of both the nucleophile (hydroxyl group) and the general base (α -amino group) in catalytic reactions (18). In NylC, Thr-267 participates in a hydrogen-bonding network with Asn-219, Asp-306, and Gly-307 (backbone-N). The positions of these four residues are spatially similar to those of Ser-250, Asn-218, Ser-288, and Gly-289 (backbone-N) in DmpA, respectively (Fig. 2B). The catalytic nucleophile (Ser-250) and the surrounding residues (Asn-207, Ser-288, and Gly-289 (backbone-N)) are also conserved in BapA. These results indicate that Thr-267 is most likely responsible for the catalytic function of NylC. Assuming that auto-proteolysis of the NylC precursor proceeds in manner similar to that proposed for other N-tn hydrolases (23–27), the reaction would be initiated by nucleophilic attack of Thr-267-O_γ to Asn-266-C_α carbon, generating a tetrahedral intermediate. This intermediate would rearrange into an ester intermediate (N–O acyl shift) that is subsequently hydrolyzed by an adjacent water molecule, producing the active enzyme.

Structural Alterations Induced by Autoprocessing

The poor electron density distribution in the carboxyl-terminal region of the α -subunit (positions 261–267) of the active NylC_A enzyme has prohibited the determination of a structural

model. The C-terminal region of the α -subunit has not been identified by x-ray crystallographic analyses of DmpA/BapA/OAT (supplemental Fig. S4). In a three-dimensional model of NylC_A, the distance between Pro-260-C_α and Thr-267-C_α is estimated to be 32.6 Å (supplemental Fig. S2). The large distance suggests that the local structural alteration that occurs is accompanied by autoprocessing. Namely, the terminal region of the α -subunit, including helix H8 and Pro-260, at each subunit interface flips out toward the adjacent molecules (Fig. 1C). We speculate that the structural alteration is important for generating the catalytic center responsible for the hydrolysis of incoming substrates (6-aminohexanoate-oligomers). Structural alterations induced by autoprocessing have been reported for other N-tn hydrolases (25).

Effect of Amino Acid Substitutions on Protein Stability

Thermal denaturation experiments using CD showed that the T_m of NylC_A and NylC_K are 60 and 67 °C, respectively (Fig. 3, Table 2). These values are 8–15 °C higher than the T_m of NylC_{p2} (52 °C). We have suggested that at least one among the five alterations (G111S, D122G, H130Y, L137A, V225M) in NylC_A contributes to the increase in the thermostability of NylC_{p2}. Moreover, at least one of the 10 alterations (D36A, A41V, M50T, I60V, A62S, T230G, V231I, V257L, E263Q, G354A) in NylC_K is estimated to contribute to the further increase in thermostability (supplemental Fig. S1) (5). The three-dimensional structure of NylC_A shows that most amino acid residues that differ among the three NylC enzymes were found to be located at the interfaces between the subunits (Fig. 1B). To examine the effect of the amino acid substitutions on the thermostability of proteins, we individually replaced the residues in NylC_{p2} with the corresponding NylC_A or NylC_K sequences.

A single amino acid mutation of either D122G or H130Y, which are located at the A/D interface, increased the T_m of NylC_{p2} by 24 and 11 °C, respectively (Fig. 3, Table 2). However a single substitution of either G111S or L137A at the same interface decreases the stability by ~10 °C. Even more drastic effects were observed with the combinations of these mutations. D122G/H130Y double mutations in NylC_{p2} enhanced the thermostability to 81 °C. Far-UV CD spectra showed very little change, even at 75 °C, for the G¹²²Y¹³⁰ mutant, whereas the parental NylC_{p2} and NylC_A enzymes denatured at 75 °C (supplemental Fig. S5). The addition of two further mutations, D36A/E263Q derived from NylC_K, to the G¹²²Y¹³⁰ mutant enhanced the T_m to 88 °C (G¹²²Y¹³⁰A³⁶Q²⁶³ mutant; Fig. 3). Notably, we observed that more than 90% of the enzyme activity was retained even after incubation of the G¹²²Y¹³⁰A³⁶Q²⁶³ mutant for 30 min at 70 °C. In contrast, an L137A mutation drastically decreased thermostability in the G¹²²Y¹³⁰ and S¹¹¹G¹²²Y¹³⁰ mutants by 27 °C. Similarly, a V225M single mutation decreased the protein stability and/or the expression level in cells because no protein was detected in the cell extracts that were prepared by expressing the M²²⁵-mutant NylC_{p2}. However, the stabilization effect is altered by the combination of other amino acid alterations close to Met-225 because the V225M mutation, in the context of the S¹¹¹G¹²²Y¹³⁰A¹³⁷ quadruple mutant, recapitulates the sequence of NylC_A and

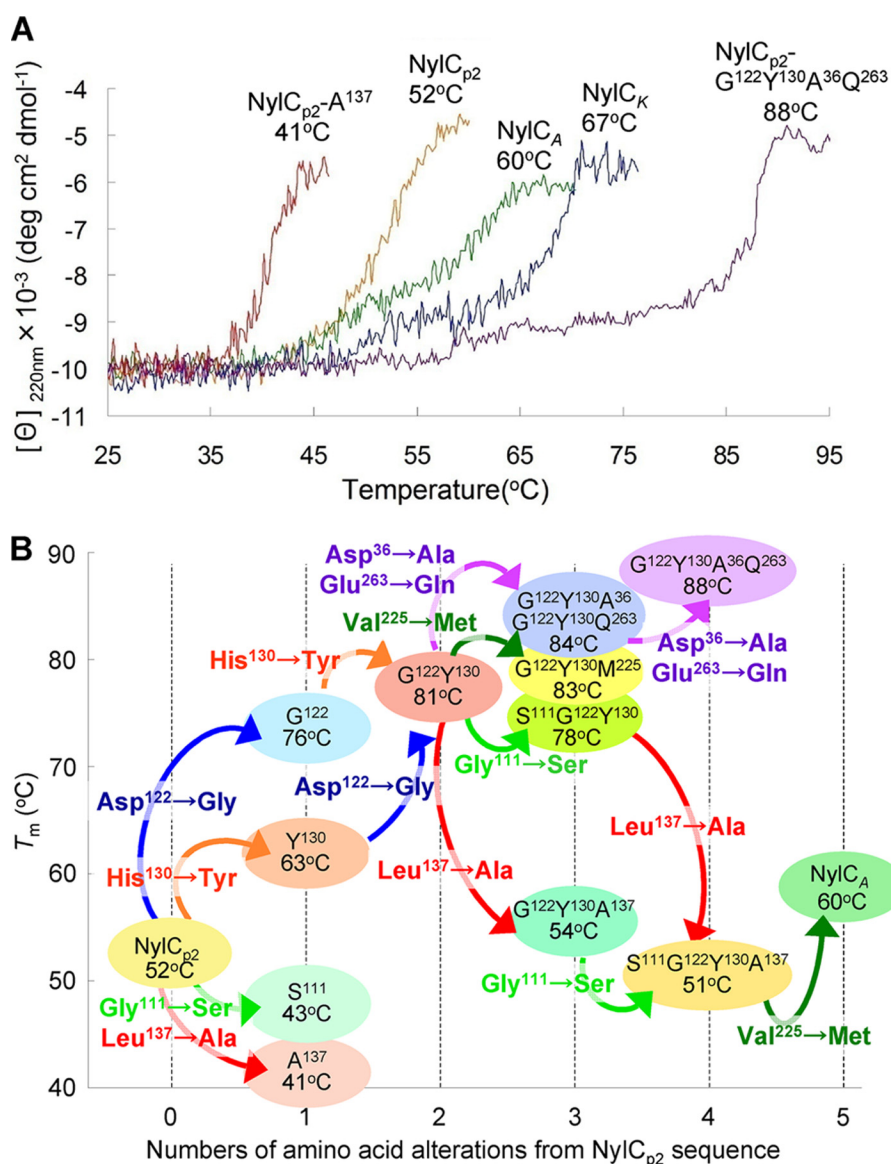


FIGURE 3. **Thermostability of NylC mutants.** A, thermal transition curves of the various NylC mutant enzymes. CD measurements were performed at 220 nm from 25 to 95 °C (1 °C min⁻¹). The results are expressed as the mean residue molar ellipticity $[\theta]$. Protein concentrations of 0.1 mg ml⁻¹ were used. *deg*, degree. B, the cumulative effects of amino acid mutations on the melting temperatures of denaturation (T_m) are shown.

improves its stability by 9 °C. Additionally, the V225M mutation in the G¹²²Y¹³⁰ mutant improved the stability by 2 °C (see G¹²²Y¹³⁰M²²⁵). These results demonstrate that subunit interactions that generate the quaternary structure drastically affect the thermostability of the enzyme (47 °C by five mutations; Fig. 3, Table 2).

Molecular Basis of Protein Stabilization

Based on the three-dimensional structure of NylC_A, we estimate that the following structural effects occurring at the subunit interfaces A/D (D122G, H130Y, L137A, G111S, and D36A) and A/B (V225M and E263Q) can cause changes in the thermostability of NylC_{p2}.

D122G—In NylC_A, Lys-159-NH₃⁺ (in molecule A) is located 2.79 Å apart from Glu-115-COO⁻ (in molecule D) (Fig. 4A). Therefore, the electrostatic effect between the two residues will enhance subunit binding around Gly-122 located on helix H4. However, in NylC_{p2}, the amino acid residue at posi-

tion 122 is replaced with Asp, and the close proximity of the acidic residue Asp-122 (close to Glu-115) will reduce this stabilization effect.

H130Y and L137A—Tyr-130 (in molecule A) is located at a loop region (Arg-127–Ala-135) between helix H4 and β-strand β7 (supplemental Fig. S4). Tyr-130-O_η forms a hydrogen bond with Glu-126-O_ε located at the end of helix H4 (Fig. 4B). Mutating Tyr-130 to His may destroy the hydrogen bonding and destabilize the loop. Therefore, a H130Y substitution in NylC_{p2} will contribute to an increase in thermostability by stabilizing the loop region. In contrast, L137A substitution (in NylC_{p2}, G¹²²Y¹³⁰ and S¹¹¹G¹²²Y¹³⁰) occurring at the same loop region resulted in the drastic decrease in the protein stability (Fig. 3B). Therefore, hydrophobic effect by Leu-137 should be involved in the stabilization of the loop region. Ala-137/Leu-137 affects the subunit interaction cooperatively with Gly-111/Ser-111 (Fig. 4C).

TABLE 2
Effect of amino acid substitutions on thermostability

Enzyme	Amino acid substitutions in NylC _{p2} sequence	T _m °C
NylC _{p2}		52
NylC _A	G111S/D122G/H130Y/L137A/V225M	60
NylC _K	D36A/A41V/M50T/I60V/A62S G111S/D122G/H130Y/L137A/V225M T230G/V231I/V257L/E263Q/G354A	67
Single mutant		
NylC _{p2} -S ¹¹¹	G111S	43
NylC _{p2} -G ¹²²	D122G	76
NylC _{p2} -Y ¹³⁰	H130Y	63
NylC _{p2} -A ¹³⁷	L137A	41
NylC _{p2} -M ²²⁵	V225M	Not expressed
Double mutant		
G ¹²² Y ¹³⁰	D122G/H130Y	81
Triple mutant		
G ¹²² Y ¹³⁰ A ³⁶	D122G/H130Y/D36A	84
G ¹²² Y ¹³⁰ Q ²⁶³	D122G/H130Y/E263Q	84
G ¹²² Y ¹³⁰ M ²²⁵	D122G/H130Y/V225M	83
S ¹¹¹ G ¹²² Y ¹³⁰	G111S/D122G/H130Y	78
G ¹²² Y ¹³⁰ A ¹³⁷	D122G/H130Y/L137A	54
Quadruple mutant		
G ¹²² Y ¹³⁰ A ³⁶ Q ²⁶³	D122G/H130Y/D36A/E263Q	88
S ¹¹¹ G ¹²² Y ¹³⁰ A ¹³⁷	G111S/D122G/H130Y/L137A	51

G111S—Hydrophilic Ser-111-O_γ (in molecule A) is surrounded by four hydrophobic residues (Tyr-98, Ala-137, Leu-139 (in molecule D) and Tyr-112 (in molecule A)) (Fig. 4C). Therefore, substitution from Ser-111 to Gly-111 (aliphatic small residue) should improve the hydrophobic stabilization effect at the A/D interface. In contrast, its reverse G111S substitution (in NylC_{p2}, G¹²²Y¹³⁰ and G¹²²Y¹³⁰A¹³⁷) actually reduces the thermostability (Fig. 3B).

D36A—Asp-36 (in molecule D) is located at helix H2, which is close to Glu-126 (in molecule A) (4.40 Å), whereas Asp-36 is replaced with Ala-36 in NylC_K (Fig. 4B). Therefore, the D36A mutation will reduce electrostatic repulsion with Glu-126. Actually, D36A substitution (in G¹²²Y¹³⁰ and G¹²²Y¹³⁰Q²⁶³) improves the thermostability (Fig. 3B).

V225M—Met-225 (in molecule A) does not directly contact with any adjacent subunits (Leu-9-C_{δ1} in molecule B is the nearest position; 5.29 Å). Met-225 is close to Gln-299 in the same subunit, which interacts with Arg-296-N_{η2} (3.01 Å) (in molecule B) (Fig. 4D). Moreover, the position of Gln-299 should be stabilized by interaction with His-245-N_{ε2} (2.81 Å). Therefore, it is likely that amino acid substitution at position 225 indirectly affects the subunit interaction. Actually, V225M substitution (in G¹²²Y¹³⁰ and S¹¹¹G¹²²Y¹³⁰A¹³⁷) improves the thermostability (Fig. 3B), probably by generating a new interaction between Met-S_δ and Gln-299-N_{ε1} (3.56 Å) (Fig. 4D). However, it should be also noted that V225M substitution decreases the expression level and/or stability of NylC_{p2} in the cell, as described above. Therefore, the total stabilization effects should be dependent on the combined interactions with the surrounding residues.

E263Q—Glu-263 is located at the terminal region of the α-subunit (Val-261–Asn-266), which had a poor electron density distribution in the x-ray diffraction study of NylC_A (Fig. 1). In NylC_K, the acidic amino acid residue Glu-263 is replaced with neutral Gln, suggesting that altering the electrostatic environment induced by the E263Q mutation assists in improving subunit interactions.

Enzymatic Hydrolysis of Polymeric Nylon-6

Although the melting point of nylon-6 (220–225 °C) is much higher than the thermostability of the enzyme, performing the reaction at a high temperature should have the following advantages for the hydrolysis of nylon. 1) Hydrogen bonding between the polymer chains is partially weakened, allowing the enzyme to attack the polymer chains that are exposed to the solvent; and 2) the release of the cleaved fragments from the solid phase generates new sites for the subsequent reaction.

The mutated sites in the G¹²²Y¹³⁰A³⁶Q²⁶³ quadruple mutant are located at the interface with other subunits and apart from catalytic Thr-267 (e.g. 26.6 Å to Gly-122 at C_α). Kinetic studies of the G¹²²Y¹³⁰A³⁶Q²⁶³ mutant with the Ahx cyclic oligomer determined that the *k*_{cat} and *K*_m values are 2.8 ± 0.11 s⁻¹ and 0.72 ± 0.06 mg ml⁻¹, respectively. When compared with the kinetic parameters of wild-type NylC_{p2} (*k*_{cat} = 6.5 ± 0.29 s⁻¹; *K*_m = 3.7 ± 0.27 mg ml⁻¹) (5), these mutations decreased the turnover of the product but improved the affinity of the enzyme for the substrate. As a result, the calculated *k*_{cat}/*K*_m of the G¹²²Y¹³⁰A³⁶Q²⁶³ mutant (3.9 s⁻¹ ml mg⁻¹) is 2-fold higher than that of NylC_{p2} (1.8 s⁻¹ ml mg⁻¹). Based on the high thermostability and increasing catalytic activity of the G¹²²Y¹³⁰A³⁶Q²⁶³ mutant, we considered this mutant to be suitable for testing the hydrolysis of polymeric nylon-6.

To increase the efficiency of nylon degradation, pellets of nylon-6 plastic were mechanically disintegrated to powder (diameter = 0.27 ± 0.14 mm, see “Experimental Procedures”). The powder was resuspended in buffer A (50 mg ml⁻¹), and the suspension was pretreated at 120 °C for 20 min. The enzyme reaction was performed using the G¹²²Y¹³⁰A³⁶Q²⁶³ mutant (0.1 mg ml⁻¹) at 60 °C for 2 h.

SIMS measures the time of flight of the secondary ions generated by the bombardment of a sample with primary ion beams and sensitively detects the alterations in mass size occurring at the surface of solid particles (36, 44). We used gas cluster SIMS equipment, in which the primary ions are argon cluster

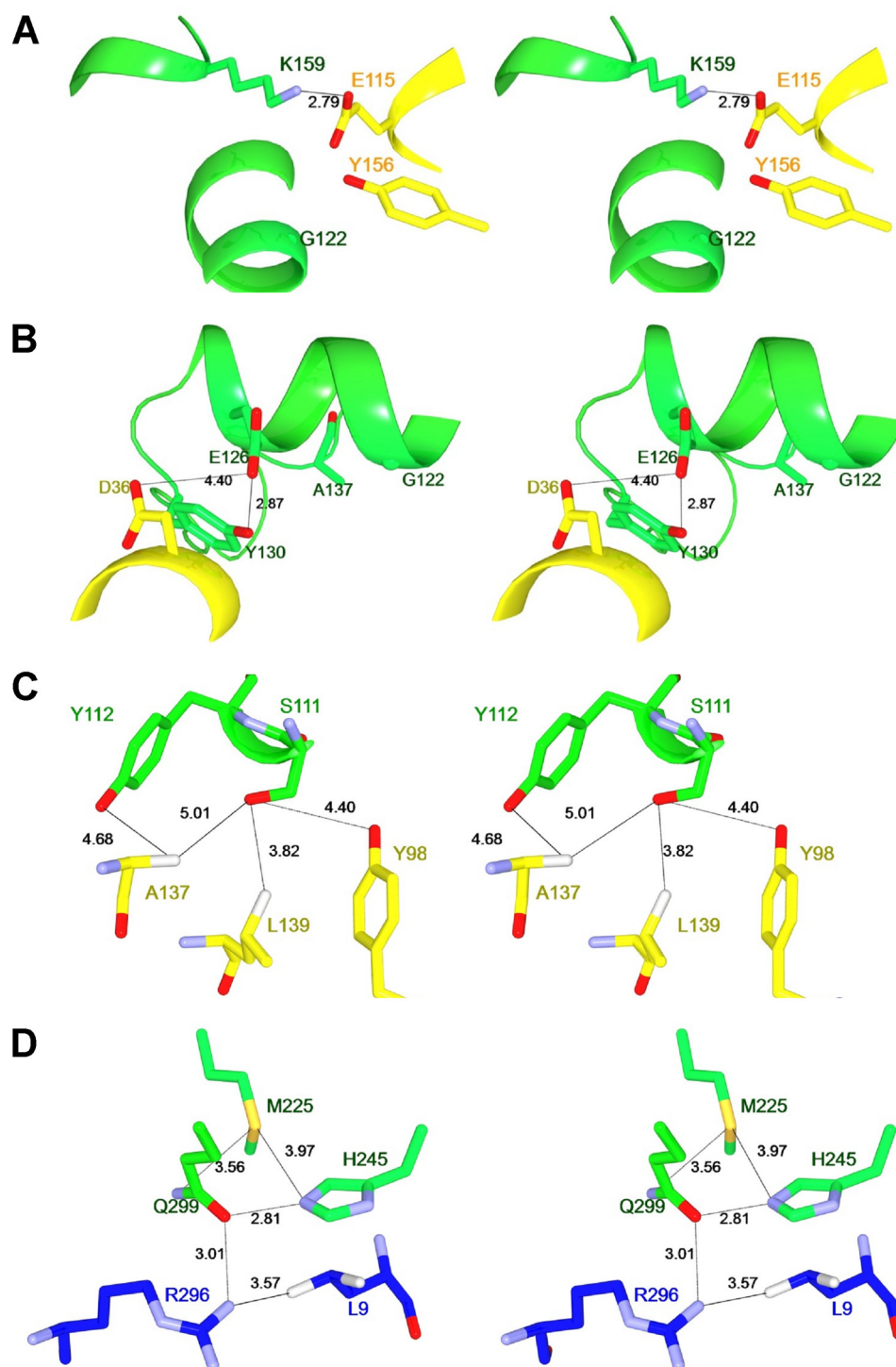


FIGURE 4. **Possible interactions at the subunit interfaces.** A–D, the possible interactions at the subunit interfaces around Gly-122 (A), Asp-36/Tyr-130 (B), Ser-111/Ala-137 (C), and Met-225 (D) are shown. Molecule A, molecule B, and molecule D in the quaternary structure are colored in green, blue, and yellow, respectively. Possible hydrogen bonds and contacts between two atoms are indicated as dotted lines with the distances listed in angstroms.

ions with a kinetic energy per atom controlled to be lower than 4 eV. This low energy of primary ions reduces the internal cleavage of polymer molecules, increasing the detection of the intact ions (36, 44). We observed a wide peak with an m/z range of 10,000–25,000 (top peak: 14,500) for untreated nylon-6, but the major peak was shifted to a smaller range with an m/z range of 8,000–23,000 (top peak: 13,000) for the reaction products (Fig. 5A). Moreover, SIMS analyses of the reaction products

(dissolved in trifluoroethanol) revealed a new peak corresponding to an m/z range of 1,500–3,000 (top peak: 2,000; Fig. 5B). The presence of the specific fragments in the solid fraction demonstrated that NylC hydrolyzes nylon, but the fragments that were produced were still bound to polymer chains through hydrogen bonding (Fig. 6). These fragments corresponded to oligomers with 13–25 monomeric units, assuming an electric charge (z) = 1. In contrast, smaller fragments (<10 monomeric

Enzymatic Hydrolysis of Nylon

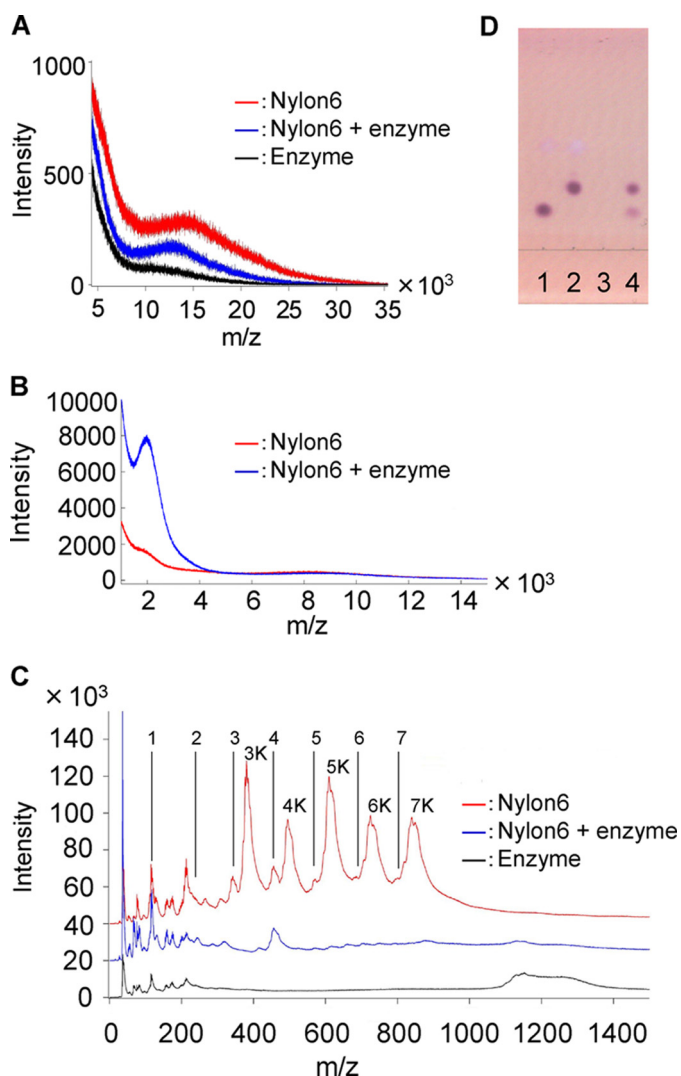


FIGURE 5. Nylon degradation tests using argon cluster SIMS and TLC. Nylon-6 powder (10 mg) was pretreated in triplicate in buffer A (180 μ l) at 120 $^{\circ}$ C for 20 min. The NylC_{p22}-G¹²²Y¹³⁰A³⁶Q²⁶³ mutant (1 mg ml⁻¹, 20 μ l) was then added and incubated at 60 $^{\circ}$ C for 2 h. The experiment was performed in triplicate. **A**, the reaction products (both the soluble and the insoluble fractions) were spotted onto a silicon plate (1 cm²). The m/z range of 5,000–35,000 was analyzed. *Red line*, pretreated nylon-6 at 120 $^{\circ}$ C in buffer A; *blue line*, enzyme-treated nylon-6; *black line*, enzyme alone. **B**, the solid fraction was washed with distilled water, lyophilized, and dissolved in trifluoroethanol (0.2 ml), and a fraction (0.02 ml) was spotted onto a silicon plate (1 cm²). The m/z range of 1,500–15,000 was analyzed. **C**, the reaction products (both the soluble and the insoluble fractions) were spotted onto a silicon plate (1 cm²). The m/z range of 0–1,500 was analyzed. The positions of Ahx monomer to heptamer (marked by 1–7; m/z = 113, 226, 339, 452, 565, 678, 791) and those of the potassium-bound forms (marked by 3K–7K) are shown. **D**, the soluble fractions (1 μ l) were spotted onto a thin layer plate and developed, and the degradation products were detected by the ninhydrin reaction (5). *Slot 1*, 6-aminohexanoate; *slot 2*, 6-aminohexanoate-dimer; *slot 3*, pretreated nylon-6 at 120 $^{\circ}$ C in buffer A; and *slot 4*, enzyme-treated nylon-6.

units) released from the solid fraction should be readily hydrolyzed to dimers and monomers (Fig. 5C). Consistent with this hypothesis, TLC analysis showed that the dimers and monomers are detected in the soluble fractions (Fig. 5D). Therefore, we confirmed that the dimers are converted to monomers by a subsequent NylB reaction.

The original nylon sample exhibited five major peaks (Fig. 5C, 3K–7K), which are estimated to be potassium ion-bound forms, $[\text{HN}(\text{CH}_2)_5\text{CO}]_n\text{K}^+$ for Ahx trimer to heptamer (m/z =

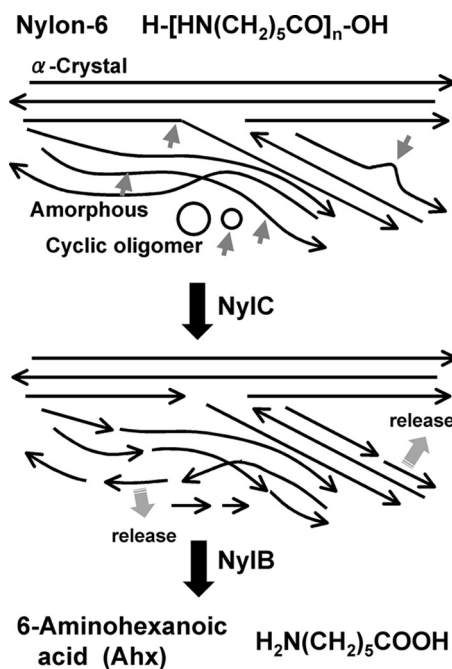


FIGURE 6. Mode of nylon-6 degradation. The polymer chains are stabilized by hydrogen bonds with adjacent chains aligned in the reverse orientation (α -Crystal) (1). Some chains constitute amorphous regions. The *circles* indicate the cyclic oligomers attached to the polymers. *Long arrows* indicate the direction of the polymer chains. *Short arrows* indicate cleavage by the enzyme. The 6-aminohexanoate-oligomers were converted to 6-aminohexanoate by the subsequent NylB reaction.

378, 491, 604, 717, and 830), because the sample includes potassium ions in the reaction mixture and the cation-bound forms generally display a higher intensity than the unbound forms in the SIMS analysis (36). However, these peaks disappeared in the NylC-treated sample. Additionally, TLC analysis of the untreated nylon did not detect any spots by ninhydrin reaction. However, after NylC treatment, spots indicative of dimers and monomers were produced (Fig. 5D). From these results, we conclude that the Ahx cyclic oligomers were attached to the nylon sample and that these compounds were hydrolyzed by NylC (Fig. 6).

Potential Use in Industrial and Environmental Applications

Because SIMS analyses of the reaction products demonstrate that the thermostable NylC hydrolyzes polymeric nylon-6, we propose that NylC should be designated as nylon hydrolase (or nylonase) (EC 3.5.-.-). Notably, there is potential for the use of nylon hydrolysis in industrial and environmental applications, although the catalytic function and thermostability of NylC, as well as the pretreatment conditions of nylons for efficient hydrolysis, warrant further improvement.

Tools for Evaluation of Biodegradability of Polyamides

Ordinary biodegradability tests of polymers are performed in activated sludge (for 1 month) or in soil (for 4 months). Because the extent of hydrolysis of the polyamides by NylC is related to the biodegradability of the polyamide, an NylC reaction followed by SIMS analysis is appropriate for the prescreening of “biodegradable polyamides.”

Improvement of Surface Structures of Nylon Fibers

Suitable methods to improve the surface structures of nylon remain poorly developed. With NylC, the partial enzymatic hydrolysis of nylon surfaces can be used to change the smoothness of nylon fibers.

Recycling of Nylons

Monomers (Ahx) obtained from nylon oligomers are aerobically metabolized in nylon oligomer-degrading strains (2–6). Suitable fermentation or biotransformation processes could enable the conversion of Ahx to other metabolites, such as organic acids or alcohols. Alternatively, Ahx may be a reusable reagent for the production of nylon-6 after conversion to ϵ -caprolactam by intramolecular dehydration. Therefore, the development of these processes could enable the recycling of nylons and decrease the environmental waste caused by the accumulation of man-made compounds.

Acknowledgment—Nylon-6 was a generous gift from Toyobo Co. Ltd. (Tsuruga, Japan).

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