Structural studies on Laz, a promiscuous anticancer Neisserial protein

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Azurin and Laz (lipidated azurin) are 2 bacterial proteins with anticancer, anti-viral and anti-parasitic activities. Azurin, isolated from the bacterium Pseudomonas aeruginosa, termed Paz, demonstrates anticancer activity against a range of cancers but not against brain tumors. In contrast, Laz is produced by members of Gonococci/Meningococci, including Neisseria meningitides which can cross the blood-brain barrier to infect brain meninges. It has been previously reported that Laz has an additional 39 amino acid moiety, called an H.8 epitope, in the N-terminal part of the azurin moiety that allows Laz to cross the entry barrier to brain tumors such as glioblastomas. Exactly, how the H.8 epitope helps the azurin moiety of Laz to cross the entry barriers to attack glioblastoma cells is unknown. In this paper, we describe the structural features of the H.8 moiety in Laz using X-ray crystallography and demonstrate that while the azurin moiety of Laz adopts a β-sandwich fold with 2 β-sheets arranged in the Greek key motif, the H.8 epitope was present as a disordered structure outside the Greek key motif. Structures of Paz and H.8 epitope-deficient Laz are well superimposed. The structural flexibility of the H.8 motif in Laz explains the extracellular location of Laz in Neisseria where it can bind the key components of brain tumor cells to disrupt their tight junctions and allow entry of Laz inside the tumors to exert cytotoxicity.

Introduction

Azurin, a blue copper-containing protein belonging to the cupredoxin superfamily, is localized in the periplasm of gram-negative bacteria and is involved in electron transport during respiration.1 Although structural and functional characteristics of bacterial azurin as a redox protein have been well documented,2 we found the following additional function of the protein: clinical isolates of Pseudomonas aeruginosa secrete azurin extracellularly in response to the presence of various human cancer cells (e.g., melanoma and breast cancer cells).3 Azurin exhibits significant cytotoxicity against cancer cells, while little cytotoxicity is observed against normal cells.4 The protein can preferentially enter cancer cells and bind to the tumor suppressor protein p53.5 The complex formed by azurin and p53 as well as its electron transfer partner cytochrome c induces apoptosis in cancer cells through cell cycle arrest at the G1 phase or caspase-mediated mitochondrial cytochrome c release.6,7 Azurin is therefore expected to be a potential drug for cancer therapy,8,9

The precursor form of azurin (148 residues) produced by P. aeruginosa is converted to a mature form (Paz, 128 residues) through the release of a signal peptide (20 residues) across the inner membrane (Fig. 1).10 Site-directed mutagenesis and deletion mutations indicate that the presence of a copper ion and redox activity was not essential for the cytotoxicity of Paz.11 The internal sequence of Paz (residue nos. 50–77 in the mature form, termed p28) functions as a putative protein transduction domain responsible for azurin’s penetration into cancer cells12 through endocytotic and nonendocytotic mechanisms.13 Based on crystal structures, bacterial azurins show structural similarity to ephrinB2, a ligand for the receptor tyrosine kinase EphB2.14,15 Cell signaling through Eph/ephrin is involved in cancer progression. The chemically synthesized C-terminal domain of Paz (residue nos. 96–113 in the mature form), which is similar to ephrinB2 at the GH loop region responsible for receptor binding,15 inhibits the growth of various cancer cells.14 Indeed, it has been shown that azurin represses ephrinB2-mediated autophosphorylation of the EphB2 tyrosine residue, thereby interfering with upstream cell signaling and contributing to cancer cell growth inhibition.14

In addition to azurin, rusticyanin, another bacterial cupredoxin, causes apoptosis in human cancer cells.16 Neisserial azurin-like protein, a “lipobox”-dependent lipid-modified azurin (Laz) bound to the outer membrane,17 also enters brain tumors such as glioblastomas and exhibits significant toxicity against the cancer cells,18 suggesting that Laz must cross the entry barrier to reach brain tumor cells. Laz includes a 39-amino acid residue region called the H.8 epitope at the N terminus prior to the azurin domain. In our previous studies with Laz, the H.8 epitope-deficient Laz are well superimposed. The structural flexibility of the H.8 motif in Laz explains the extracellular location of Laz in Neisseria where it can bind the key components of brain tumor cells to disrupt their tight junctions and allow entry of Laz inside the tumors to exert cytotoxicity.

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essential for disrupting the entry barrier to provide access to brain tumor cells. To obtain clues on the entry of Laz into brain essential for disrupting the entry barrier to provide access to brain tumor cells.

**Results**

**Comparison of primary structure between Laz and Paz**

Laz consists of 183 amino acid residues with a molecular weight of 18532 (Fig. 1). Similar to other bacterial outer membrane-bound lipoproteins, Laz includes a conserved 4-amino acid sequence termed lipobox [LVI][ASTVI][GAS]C at the N terminus. The thiol group of the cysteine residue in the lipobox sequence [LVI][ASTVI][GAS]C at the N terminus was fold10- higher than that of copper ion. Inductively coupled plasma atomic emission spectrometry analysis revealed that the protein contained metal ions such as zinc and copper. The content of zinc ion was fold10- higher than that of copper ion.

**Structure determination**

Stick-shaped crystals of Laz were found in a droplet consisting of 1.5 M ammonium sulfate, 0.1 M Tris-HCl (pH 8.5), and 20% glycerol (Fig. 3, inset). Diffraction images of the crystal were collected at up to 1.90 Å resolution (Fig. 3). The structure of Laz was solved by molecular replacement, using Paz as a reference model. Statistics for data collection and structure refinement are shown in Table 1. The N-terminal sequence (40 amino acid residues, Asn58–Asp183) is in good agreement with that determined by mass spectrometry (16832). The difference in molecular weight between SDS-PAGE and mass spectrometry is possibly due to the significantly low pI of the H.8 epitope. Inductively coupled plasma atomic emission spectrometry analysis revealed that the protein contained metal ions such as zinc and copper. The content of zinc ion was fold10- higher than that of copper ion.

Thus, X-ray crystallography of Laz was performed to clarify the structure–function relationship of the H.8 epitope involved in disrupting the entry barrier to brain tumor cells.

**Purification and characterization of Laz**

Lipidation is known to hinder crystallization of lipoproteins. Thus, the Laz mutant with a pectate lyase B (pelB) signal peptide instead of the native lipobox peptide was purified and characterized because this mutant can also enter glioblastoma cells and exhibit cytotoxicity. In the mutant, the signal peptide was fused to the H.8 epitope with the N-terminal cysteine residue (Cys18) substituted with alanine (Ala18) to avoid lipidation (Fig. 1). Laz purified from the recombinant E. coli cells showed a single protein band of 22 kDa on the SDS-PAGE gel (Fig. 2, inset). Positive-mode matrix-assisted laser desorption/ionization time-of-flight mass spectrometry demonstrated that the molecular weight of the purified protein was 16832.
Cα atoms), indicating that the structures of molecules A and B are basically identical to each other. On the basis of theoretical curves in the plot calculated according to Luzzati,21 the absolute positional error was estimated to be 0.214 Å/C at 1.90 Å resolution. Ramachandran plot analysis22 in which the stereochemical correctness of the backbone structure is indicated by (ϕ, ψ) torsion angles showed that 93.1% of nonglycine residues lie within the most favored regions and 6.9% of nonglycine residues lie in the additionally allowed regions.

Overall structure

The overall structure (Fig. 4A) and topology of secondary structure elements (Fig. 4B) indicate that the H.8 epitope-deficient Laz (C-terminal domain of Laz, Laz-C) adopts a β-sandwich fold as a basic scaffold. Five additional α-helices (H1, residues 96–101; H2, 111–120; H3, 121–126; H4, 153–158; and H5, 170–174) are included in the domain. The β-sandwich fold consists of 2 β-sheets arranged in the Greek key motif. One sheet (SA) is composed of 3 strands (SA1, 60–65; SA2, 85–91; and SA3, 147–152) and the other (SB) has 5 strands (SB1, 75–79; SB2, 176–182; SB3, 162–165; SB4, 105–108; and SB5, 135–138).

Structural homologs of Laz-C were searched in Protein Data Bank (PDB) using the DALI program.23 Bacterial azurin proteins were found to exhibit significant structural homology with Laz-C (Table 2). The overall structure of Laz-C is most similar to that of Alcaligenes xylosoxidans azurin (PDB code, 1RKR; Z = 24.7) with rmsd of 1.0 Å for 126 Cα. Laz-C is also structurally similar to Paz (Z = 24.5). Superimposition of the crystal structures of Laz-C and Paz (PDB code, 1AG0) is shown in Fig. 5A. Because a large number of bacterial azurins and plant plastocyanins with the Greek key motif are categorized into the plastocyanin/azurin-like family in the cupredoxin superfamily on the SCOP database (http://scop.mrc-lmb.cam.ac.uk/scop/), Laz-C is considered to be structurally classified into this family.

Metal-binding site

A density map for metal ions was observed in each protein molecule, suggesting that Laz contained one molecule of metal ion per protein molecule. Although inductively coupled plasma atomic emission spectrometry showed that 2 ions such as zinc and copper are included in Laz, the protein was suggested to contain a zinc or copper ion. Based on the content of metal ions, Laz used in this study is considered to incorporate a zinc rather than a copper ion. This is possibly due to an insufficient supply of copper ions during culturing of the bacteria, purification of the protein, or both. The four atoms, O of Gly101, Nδ of His102, Sγ of Cys166, and Nδ of His171, are coordinated to the zinc ion, and the coordination geometry comprises a distorted tetrahedron (Fig. 5B, left). The distance between the zinc ion and these atoms ranges from 1.94 to 2.30 Å (average, 2.15 Å).

The copper ion in Paz (Cu-Paz) is known to be bound to 5 atoms, O of Gly65, Nδ of His66, Sγ of Cys132, Nδ of His137, and Sδ of Met141 (Fig. 5B, right, purple).24 These 5 residues (G/H/C/H/M) involved in binding to the copper ion are also completely conserved in Laz (Fig. 1), although Met175 in Laz-C is located slightly
farther from the zinc ion (3.50 Å) (Fig. 5B, left). Among a large number of azurin-like crystal structures determined to date, a zinc ion-bound Paz (Zn-Paz) expressed in recombinant E. coli cells has been isolated and structurally characterized.\(^25\) In the crystal structure of Zn-Paz (PDB code, 1E67), the distance between the zinc ion and S6 of Met41 is 3.31 Å (Fig. 5B, right, blue), while the corresponding distance is 2.97 Å in Cu-Paz (PDB code, 3AZU) (Fig. 5B, right, purple), indicating that Laz-C is structurally similar to Zn-Paz.

**Table 1.** Data collection and refinement statistics

| Space group  | P6\(_5\) |
|--------------|-----------|
| Unit cell parameters (Å) | \(a = b = 77.5, c = 94.6\) |

| Data collection | |
|-----------------|------------------|
| Wavelength (Å)  | 1.0000 |
| Resolution limit (Å) | 50.0–1.90 (1.97–1.90)\(^a\) |
| Total reflections | 76,372 |
| Unique reflections | 25,438 |
| Redundancy | 3.1 (2.9) |
| Completeness (%) | 96.9 (92.8) |
| \(f/sigma (f)\) | 9.6 (1.54) |
| \(R_{merge}\) (%) | 6.0 (39.7) |

| Refinement | 252 (126 × 2) amino acid residues, 161 water molecules, 2 zinc ions, 2 sulfate ions, 2 glycerol molecules |
|-----------------|--------------------------------------------------|
| Resolution limit (Å) | 31.6–1.90 (1.95–1.90) |
| Used reflections | 23,363 (1,637) |
| Completeness (%) | 97.0 (92.5) |
| Average B factor (Å\(^2\)) | 23.8, 24.6 |
| Protein (molecule A, B) | 36.0 |
| Copper ion | 20.2 |
| Sulfate ion | 33.7 |
| Glycerol molecules | 33.0 |
| \(R\) factor (%) | 19.3 (25.5) |
| \(R_{free}\) (%) | 22.5 (29.5) |

| Root-mean-square deviations | |
|------------------------|------------------|
| Bond (Å)  | 0.008 |
| Angle (°) | 1.15 |

| Ramachandran plot (%) | |
|-----------------------|------------------|
| Most favored regions | 93.1 |
| Additional allowed regions | 6.9 |
| Generously allowed regions | 0.0 |

\(^a\)Data on highest shells are given in parentheses.

**Discussion**

Because of disorder, the structure of the H.8 epitope could not be determined. Recently, natively unfolded or intrinsically disordered proteins (IDPs) or regions have been found mainly in eukaryotic cells.\(^26,27\) In addition, the sigma factor inhibitor FlgM in bacteria has been identified as an IDP.\(^28\) A large number of IDPs are classified as a superfamily of macromolecule (DNA, RNA, or protein)-associated proteins such as transcriptional factors, RNA-binding proteins, and cell cycle regulators.\(^27\) A typical IDP is the transcriptional factor leucine zipper protein GCN4.\(^29\) The basic DNA-binding region in GCN4 is unfolded in the absence of DNA, while a helical structure is inductively formed in the region of DNA binding. This is called the coupled binding and folding process.\(^26\)

Through bioinformatics approaches, 5 common features of IDPs are postulated:\(^26,27\) (i) Short sequences are repeated in IDPs. (ii) Little sequence homology is observed in disordered regions among IDPs. (iii) Disordered regions are located at the N or C terminus of IDPs. (iv) IDPs are primarily produced by eukaryotes. (v) IDPs are frequently localized in the nucleus and rarely in mitochondria. In case of Laz, the N-terminal H.8 epitope with a 5 residue-repeated sequence (AAEAP, a typical but somewhat variable sequence) is attached to the azurin domain (Fig. 1). No proteins or peptides were found to be homologous to the H.8 epitope in the protein databases. Several programs for identification of intrinsically disordered regions in IDPs are available on the web. The H.8 epitope of Laz was predicted to be intrinsically disordered by all the typical programs, PONDR-FIT,\(^30\) DISOPRED 2,\(^31\) and DisEMBL\(^32\) (Figs. 1 and 6). Although the target molecules that bind to Laz have not been identified in the entry barrier to brain tumors, the intrinsically disordered H.8 epitope may inducibly fold into a rigid structure by binding to target molecules.

Why is the H.8 epitope of Laz intrinsically disordered rather than being integrated with the folded region of Laz-C? It should be emphasized that the H.8 epitope of Laz seems to have 2 major functions in which a loosely associated H.8 epitope may be required. First, the H.8 epitope has been shown to be involved in facilitating the entry of Laz-C (the 127 amino acid azurin moiety) into glioblastomas\(^18\) as well as its passage through the blood–brain barrier.\(^33\) Thus, the presence of the H.8 epitope allows the Laz-C moiety to enter glioblastoma or other brain tumors when Neisseria such as N. meningitides are present in the brain meninges. Because P. aeruginosa does not normally cause infections in the brain, it has no particular need to equip its anticancer weapon Paz with a blood–brain barrier-crossing capability. The other hypothetical function of the H.8 epitope is to allow N. meningitides to protect its weapon Laz-C from being attacked by the immune system following the release of Laz-C from the bacterial cells for entry into the tumor. Interestingly, the H.8 epitope has been shown\(^34\) to induce formation of blocking antibodies that bind complements or other antibodies generated by the host to remove the invading bacteria thereby inactivating the host-generated antibodies. Such induction of blocking antibodies by the H.8 epitope is believed to prevent

**Table 2.** Structure-based homology with Laz

| Z score | Description | rmsd\(^a\) | Lali\(^b\) | PDB ID |
|---------|-------------|----------|----------|--------|
| 24.7    | Azoligens xylosoxidans azurin | 1.0     | 126      | 1RKR   |
| 24.7    | Pseudomonas putida azurin | 0.8     | 125      | 1NWP   |
| 24.5    | Azoligens denitrificans azurin | 1.0     | 126      | 1AZB   |
| 24.5    | Pseudomonas aeruginosa azurin | 0.9     | 125      | 1R1C   |

\(^a\)Root-mean-square deviations.

\(^b\)Total number of equivalence residues.
complement-mediated bactericidal action and provide a survival advantage to group B meningococci. It is, however, equally possible that such blocking antibodies protect Laz from immune attack when released for entry into brain tumors. Normally, azurin (Paz) demonstrates low immunogenicity because of its structural similarity with immunoglobulins. Both azurin and immunoglobulins, while having low sequence identity, demonstrate structural features with β-sandwich packing. Both these proteins exhibit tyrosine residues in a motif called the “tyrosine corner,” which plays a significant role in the structural stability and folding of the Greek key β-barrel structure exhibited by azurin and immunoglobulins. This low immunogenicity of azurin (Laz-C) is further enhanced by the presence of the intrinsically disordered H.8 epitope that elicits blocking antibodies to protect Laz from immune attack.

In conclusion, this is the first report on the structure of Neisseria azurin (Laz), which is cytotoxic to brain and other tumor cells. Laz is composed of an N-terminal intrinsically disordered H.8 epitope and a rigid azurin domain with the Greek key motif. In contrast, the P. aeruginosa azurin Paz, which lacks the H.8 epitope, is deficient in the ability to enter brain tumor cells and consequently demonstrates significantly lower cytotoxicity against such cells. It should be noted, however, that as mentioned earlier, Paz, unlike Laz, is deficient in entry in glioblastoma cells, while H.8-Paz, where the H.8 epitope was cloned in the N-terminal of Paz, had significant cytotoxicity against glioblastoma cells, confirming the presence of the H.8 epitope as critical for entry in brain tumors. It should also be noted that Paz, Laz and H.8-Paz had not only anticancer activity, but strong anti-viral activity against the AIDS-causing HIV-1 virus and strong anti-parasitic activity against parasites such as the malarial parasite Plasmodium falciparum and the toxoplasmosis-causing parasite Toxoplasma gondii. The potential clinical importance of such bacterial proteins as Paz and Laz is reflected in the fact that a 28 amino acid peptide fragment from azurin (azurin 50–77), termed p28, has shown very little toxicity in 15 stage IV cancer patients in a phase I clinical trial, but significant beneficial effect including partial and complete regression of the drug-resistant tumors in several patients. Another phase I trial with p28, currently in progress in pediatric brain tumor patients in 11 hospitals in the US.
with a linear gradient of saturated ammonium sulfate (30–0%, 300 ml), and a 6.6-ml fraction was collected every 10 min. The Laz fractions, eluted with 20–10% saturated ammonium sulfate, were combined and dialyzed for 3 h against 20 mM Tris-HCl (pH 7.5). The dialysate was applied to a SuperQ-Toyopearl 650S (Tosoh) column (1 cm × 10 cm) previously equilibrated with 20 mM Tris-HCl (pH 7.5). The absorbed proteins were eluted with a linear gradient of NaCl (0–0.5 M) in 20 mM Tris-HCl (pH 7.5, 30 ml), and a 1-ml fraction was collected every 1 min. The Laz fractions, eluted with 0.2–0.25 M NaCl, were combined and applied to a HiLoad 16/60 Superdex 75 pg (GE healthcare) column (1.6 cm × 60 cm) previously equilibrated with 20 mM Tris-HCl (pH 7.5) containing 0.15 M NaCl. Laz was eluted with the same buffer (120 ml), and a 2-ml fraction was collected every 2 min. The Laz fractions were combined and dialyzed for 3 h against 20 mM Tris-HCl (pH 7.5). The dialysate was used as the purified Laz. The homogeneity of the purified protein was confirmed by SDS-PAGE. The protein content was determined by measuring the absorbance at 280 nm using a cuvette with a path length of 1 cm, assuming that $E_{280} = 0.177$ (Laz) corresponds to 1 mg/ml.

**Analytical methods**

The N-terminal amino acid sequence of Laz was determined by the Edman degradation method using a Procise 492 protein sequencer (Applied Biosystems). The molecular weight of Laz was determined using a matrix-assisted laser desorption/ionization time-of-flight mass spectrometer (AXIMA Performance; Shimadzu); sinapinic acid was used as the matrix. Metal ions included in Laz were examined using an inductively coupled plasma atomic emission spectrometer (ICPS-8100; Shimadzu).

**Crystallization and X-ray diffraction**

Laz (98 mg/ml) was crystallized at 20°C using the sitting drop vapor diffusion method. The mother liquor consisting of 1.5 M ammonium sulfate, 0.1 M Tris-HCl (pH 8.5), and 20% glycerol (100 µl) was used as a reservoir solution, and 1 µl of the Laz solution was mixed with 1 µl of the reservoir solution to form the drop. The Laz crystal was removed from the drop solution using a mounted nylon loop (Hampton Research) and then placed directly into a cold nitrogen gas stream at −173°C. X-ray diffraction images of the crystal were collected at −173°C under a nitrogen gas stream using a Jupiter 210 charge-coupled device detector and synchrotron radiation at the BL-38B1 station of SPring-8 (Japan). The diffraction data for the native crystal were collected at 1.90 Å resolution and processed using the HKL2000 program package. Data collection statistics from the crystal are summarized in Table 1.

**Structure determination and refinement**

The crystal structure of Laz was solved by molecular replacement using the Molrep program in the CCP4 program package; the Paz structure (PDB code, 1AG0) was used as a reference model. The Coot program was used for manual modification of the initial model. Initial rigid body refinement and several rounds of restrained refinement against the dataset were...
performed using the Refmac5 program.\(^44\) Water molecules were incorporated where the difference in density exceeded 3.0\(\sigma\) above the mean and the 2\(F_o-F_c\) map showed a density of over 1.0\(\sigma\). At this stage, metal ions were included in the calculation and refinement continued until convergence at maximum resolution. Protein models were superimposed and their rmset was determined using the LSQKAB program,\(^45\) a part of CCP4. Final model quality was determined using the PROCHECK program.\(^46\) Ribbon plots were prepared using the PyMOL program.\(^47\) Coordinates used in this study were taken from PDB, Research Collaboratory for Structural Bioinformatics.\(^48\)

Protein structure accession number

The atomic coordinates and structure factors (PDB code, in PDB, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

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Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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