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Conformational Dynamics of the Anthrax Lethal Factor Catalytic Center[†]

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ABSTRACT: Anthrax lethal factor (LF) is a zinc-metalloprotease that together with the protective antigen constitutes anthrax lethal toxin, which is the most prominent virulence factor of the anthrax disease. The solution nuclear magnetic resonance and in silico conformational dynamics of the 105 C-terminal residues of the LF catalytic core domain in its apo form are described here. The polypeptide adopts a compact structure even in the absence of the Zn²⁺ cofactor, while the 40 N-terminal residues comprising the metal ligands and residues that participate in substrate and inhibitor recognition exhibit more flexibility than the C-terminal region.

Anthrax is an infectious disease of animals and humans caused by the Gram-positive bacterium Bacillus anthracis. Cutaneous anthrax is rarely lethal for humans, while inhalation of B. anthracis spores is often fatal if the infection is not properly diagnosed and treated with antibiotics during the early stages of infection. After anthrax spores are inhaled, they adhere to alveolar macrophages and then germinate. Bacteria migrate to lymph nodes, where they rapidly multiply and excrete a tripartite exotoxin comprised of protective antigen (PA, 83 kDa), lethal factor (LF, 90 kDa), and calmodulin-activated edema factor adenylyl cyclase (EF, 89 kDa) (1). LF is a highly specific Zn-protease that cleaves members of the mitogen-activated protein kinase kinase family near their amino termini, leading to the inhibition of one or more important signaling pathways (2). According to LF X-ray models (3), the catalytic site and the Zn^{2+} cofactor of the enzyme are found in C-terminal domain IV (residues 552-776). Zn²⁺ is bound to the protein via the consensus sequence of the gluzincin subfamily (HExxH-linker-E, where x is any amino acid) (4). The 105 C-terminal residues of domain IV (LF₆₇₂₋₇₇₆ hereafter) contain this motif and most of the active site of LF, including key residues for substrate recognition and inhibitor binding. The LF₆₇₂₋₇₇₆ construct is expressed in high yield in *Escherichia coli* as a soluble protein (5).

Here, we present the solution nuclear magnetic resonance (NMR) structure and dynamics of LF₆₇₂₋₇₇₆ in its metal-free form. The protein is monomeric in solution, based on gel filtration chromatography and ¹⁵N relaxation data. It adopts a compact structure even in the absence of the metal cofactor (Figure 1), exhibiting remarkable similarities with the catalytic site of full-length LF as seen in X-ray structures, both in its Zn(II)-loaded form and in its Zn(II)-free form (3). The structure

is dominated by six α -helices ($4\alpha 4 - 4\alpha 9$ according to the helix numbering of the crystal structures), with the highly conserved HExxH Zn(II)-binding motif being located in the N-terminal $4\alpha 4$ helix.

The almost complete assignment of apoLF₆₇₂₋₇₇₆ has been deposited in the BioMagResBank (entry 16735). According to circular dichroism (CD) and NMR data (NOEs, $^3J_{\rm HNH\alpha}$, and chemical shift index), the helical content of LF₆₇₂₋₇₇₆ in solution is in the range of 53–58% (\sim 64% in X-ray structures). The solution NMR structure of apoLF₆₇₂₋₇₇₆ was determined from 1181 NOEs, 56 angle constraints, and 28 H-bonds (PDB: 2L0R). The compact fold is based on 209 long-range NOE constraints, many of them between side chains of hydrophobic residues in the LF catalytic site (Figure 1A,B). In contrast, only a few NOEs were identified in the Asp701-Phe714 region (consecutive loops L1 and L2), which links helix $4\alpha4$ to helix $4\alpha5$ (Figure 1B).

The six α -helices of apoLF₆₇₂₋₇₇₆, observed also in LF X-ray models, are arranged with the same topology as in the crystal structure (Figure 1B). The root-mean-square deviation (rmsd) between the solution structure of apoLF₆₇₂₋₇₇₆ and the Zn(II)-LF crystal structure [Protein Data Bank (PDB) entry 1J7N] is 1.26 Å for the backbone and 2.39 Å for all heavy atoms, while a comparison with the apoLF-MEK2 peptide complex (PDB entry 1JKY) (3) shows rmsd values of 1.37 and 2.53 Å, respectively (Figure 1B and Table S3 of the Supporting Information). In the solution structure, helices $4\alpha 4-4\alpha 9$ are rather well-defined through helix diagnostic NOEs and backbone dihedral angle constraints (Figure S3 of the Supporting Information), with the exception of helix $4\alpha 5$ and to a lesser extent helix $4\alpha 9$. The former follows the highly disordered and flexible Asp701—Asp706 and Asn710—Phe714 segments, while the latter is the C-terminal helix. Helix $4\alpha7$ of apoLF₆₇₂₋₇₇₆ is tightly packed among helices $4\alpha 4$, $4\alpha 6$, $4\alpha 8$, and $4\alpha 9$ and forms an angle of 127° with helix $4\alpha 4$ to adopt a so-called "X" topology, typical for the conserved two-helix active site structure of gluzincin proteases (4). These two helices provide the three Zn²⁺-binding residues, i.e., His686 and His690 in helix 4α4 and Glu735 in helix 4α 7. In apoLF₆₇₂₋₇₇₆, these two histidines along with His745 are found to be in the charged state according to H-15N heteronuclear single-quantum coherence (HSQC) NMR spectra optimized for the detection of imidazole ring protons [analogous signals of His749 were too weak to be detected possibly because of exchange broadening (Figure S4 of the Supporting Information)]. Thus, the zinc binding site of apoLF₆₇₂₋₇₇₆ has the same net electrostatic charge of +2 as in Zn(II)-LF. This similarity in the electrostatics of the catalytic site could well explain the remarkable structural similarity between the Zn(II)-LF X-ray structures and the apoLF solution structure presented

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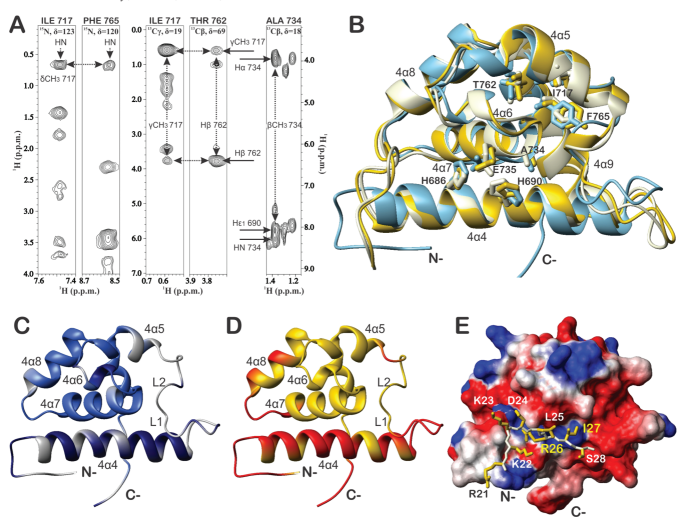


FIGURE 1: (A) Characteristic long-range NOEs defining the hydrophobic core of LF₆₇₂₋₇₇₆. (B) Superimposed LF catalytic core domain structures: X-ray Zn(II)-LF in light gray (PDB entry 1J7N), X-ray apoLF in yellow (PDB entry 1JKY), and NMR apoLF in sky blue (PDB entry 2L0R). (C) Backbone dynamics are illustrated in colors according to S^2 order parameters: royal blue for residues for which $S^2 > 0.6$, and light gray for residues for which S^2 values are not available. (D) Backbone dynamics according to MD. Atomic rms fluctuations of $S^2 > 0.6$, are colored red. (E) LF-MKK3 complex built through homology modeling using the solution structure of apoLF₆₇₂₋₇₇₆ and the MKK3 LF-bound conformation according to ref 7. Molecular surface electrostatic potentials of less than $S^2 > 0.6$ are colored red, potentials of greater than $S^2 > 0.6$ and neutral potentials white (k is the Boltzmann constant). The MKK3 Arg26 (P1) guanidinium group is oriented toward the acidic catalytic site channel and surrounded by glutamates 682, 687, and 735. The carbonyl group of the scissile bond may be oriented toward the positive potential raised by the two charged histidines 686 and 690. This figure was generated with MOLMOL (8).

Glu687 in helix $4\alpha4$ is the conserved glutamate residue in the HExxH motif of all gluzincin proteases (acting as a base to activate the Zn²⁺-bound water during catalysis). The side chains of these residues are solvent-exposed with the exception of His686, which is oriented toward the catalytic pocket, and exhibit rather similar orientations in the solution structure of apoLF₆₇₂₋₇₇₆ and the crystal structures of Zn(II)-loaded (PDB entry 1J7N) and Zn(II)-free (PDB entry 1JKY) LF. The same holds true for the side chains of Tyr728 and Glu739. These two residues along with Glu687 belong to the secondary shell of Zn²⁺, and both experimental (6) and in silico data (7) suggest a significant role of Tyr728 in LF activity. Specifically, computational studies of LF-MKK complexes show that Tyr728 actively participates in substrate harboring through interactions with the carbonyl group of the scissile bond, while in the crystal structure, Tyr728 is hydrogen bonded to a Zn²⁺-coordinated water molecule. Glu739 is also a key residue of the S1 LF catalytic site subpocket and forms a salt bridge with Arg742 in the majority of the NMR models in a manner similar to that reported for the LF crystal structures. The negative charge density in this site (four positions

downstream Zn²⁺-bound Glu735) is rather conserved in the vast majority of gluzincins, and this position is occupied by an aspartate in thermolysin and angiotensin-I converting enzyme. Its role, according to simulated LF-MKK complexes, is to interact efficiently with P1 Arg of LF substrates MKK3b and MKK4 (Figure 1E) determining the orientation of the peptide inside the enzyme's catalytic groove (7). All these residues orient their side chains in the NMR models in a manner similar to that in X-ray structures toward the catalytic channel of the enzyme. Additionally, helices $4\alpha 4$ and $4\alpha 7$ and the helix $4\alpha 4$ —helix $4\alpha 5$ linker (including loops L1 and L2) are rich in acidic residues [Glu682, Glu687, Asp693, Asp694, Asp701, Asp706, Glu733, Glu735, and Glu739 (Figure 1E)]. The carboxyl moieties of these residues are exposed toward the catalytic channel, thus contributing to the substrate subpockets (S4-S1) that accommodate the basic residues of MEKs and MKKs.

The backbone dynamics of apoLF₆₇₂₋₇₇₆ were studied through the analysis of ¹⁵N R_1 and R_2 relaxation rates and heteronuclear ${}^{1}H^{N}{}^{-15}N$ NOEs (Figure S5 of the Supporting Information). The correlation time for isotropic tumbling in solution as

calculated from the R_2/R_1 ratio is 6.8 \pm 0.2 ns, clearly indicating that apoLF₆₇₂₋₇₇₆ is in the monomeric state. Model-free analysis of ¹⁵N relaxation data as implemented in Tensor2 shows distinct dynamic properties for the N- and C-terminal segments of the polypeptide chain. For the N-terminal residues 675-711, the average S^2 order parameter is 0.26 ± 0.13 , whereas S^2 values are much higher for the C-terminal part of apoLF₆₇₂₋₇₇₆ (average value of 0.88 ± 0.17 for residues 717–776) (Figure 1C and Figure S6 of the Supporting Information). Thus, the N-terminal part of the construct is significantly mobile on the subnanosecond time scale, whereas the C-terminal segment appears well-ordered.

The conformational dynamics of apoLF₆₇₂₋₇₇₆ was also probed by a 10 ns unrestrained molecular dynamics (MD) simulation of the average energy-minimized NMR model in explicit water. The MD data on the nanosecond time scale are in good agreement with the ¹⁵N relaxation measurements confirming that the N-terminal part of the protein exhibits higher flexibility than the C-terminal region. In particular, the N-terminal helix 4α4 that is a key component of both the Zn²⁺ binding site and the substrate MKK and/or MEK docking site is intrinsically mobile. Although some helix-typical (i, i+3,4)H-bonds in helix $4\alpha 4$ are lost during the MD, it does not collapse during the simulations or lose contact with helix $4\alpha 7$. This is attributed to several main chain or side chain interactions of helix $4\alpha4$ with helix $4\alpha7$ and loops L1 and L2. Both His686 and His690 of the Zn^{2+} -binding motif in helix $4\alpha 4$ interact through H-bonds of their $N^{\delta 1}$ atoms with the side chains of $4\alpha 7$ Glu735 and $4\alpha 6$ Thr731, and 4α4 Asp693 and L2 Thr709, respectively. His686 and His690 N^{δ 1} atoms are found on average 2.70–2.80 Å from carboxyl oxygen atoms O^{ϵ 1/2} of Glu735 and O^{δ 1/2} of Asp693. For more than 60% of the MD time course, Asp693 $O^{\delta 1/2}$ atoms are found at a favorable distance for H-bonding (2.70-2.90 Å) with the $O^{\gamma 1}$ atom of Thr731 in the enzyme's S1' subpocket and $O^{\gamma 1}$ of Thr709. In agreement with NMR data, these residues are involved in proton-proton interactions detected in ¹³C- and ¹⁵Nedited NOESY spectra and participate in a NOE network between 4\alpha4 His686 and 4\alpha7 residues Ala738, Glu739, and Arg742, surrounding Thr731. In turn, Thr731 protons are found within NOE distance of Glu735. Finally, there are also NOEs between the residues in the $4\alpha4-L2$ H-bond network observed during the MD calculation, namely, between 4α4 His690 and S4' Asp693 and between Asp693 and Leu707 and Val708 preceding L2 Thr709. According to experimental and computational data, Leu707 and Val708 participate in hydrophobic interactions between LF substrates and subpockets S3'-S5' of the enzyme (3, 6, 7).

In conclusion, we have described the solution structure and the dynamical behavior of the monomeric polypeptide that represents the C-terminal region of the LF catalytic core in its metal free state. Despite the absence of the metal cofactor, the NMR model is strikingly similar to the holoenzyme and even exhibits a similar charge density in the catalytic channel, as there is experimental evidence that the zinc binding histidines are in the doubly protonated, charged state. NMR relaxation data in concert with MD simulations support an overall compact fold of the enzyme's catalytic site. However, they reveal a higher

flexibility of helix 4\alpha4 comprising the first Zn²⁺-binding motif and the loops in the sequential stretch of Asp701-Ser711. Additional MD calculations on the NMR model reconstituted with a Zn²⁺ ion and a coordinated H₂O molecule indicate a welldefined helix 4\alpha4 in the holoenzyme during the course of the simulation (10 ns). However, the N-terminus still exhibits a higher conformational plasticity with respect to the rest of the protein (Figure S6C of the Supporting Information). Besides its role in binding the metal cofactor, helix $4\alpha 4$ is crucial for substrate or inhibitor recognition. Despite its flexibility on the picosecond to nanosecond time scale in apoLF, helix $4\alpha 4$ remains in the spatial proximity of helix 4\alpha7 through H-bonds, and the LF catalytic site retains its intrinsic negative charge density. The negatively charged residues in the LF catalytic channel control recognition and binding of MEKs and MKKs, which are characterized by a significant number of positively charged residues close to the scissile bond (Figure 1E). Metal reconstitution studies are underway, and a metal-loaded LF₆₇₂₋₇₇₆ construct could provide additional atomic-level structural, dynamical, and functional insights into LF-substrate or LF-inhibitor binding and interaction.

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SUPPORTING INFORMATION AVAILABLE

Experimental methods, NMR data and structural statistics (Tables S1 and S2), comparison of LF₆₇₂₋₇₇₆ NMR and X-ray models (Table S3), 30 NMR models (Figure S1), NOE classification and rmsd per residue (Figure S2), NOE sequential connectivity map and $^3J_{\rm HNH\alpha}$ values (Figure S3), $^2J_{\rm HN}$ -optimized $^1H^{-15}N$ HSQC spectrum of LF₆₇₂₋₇₇₆ (Figure S4), ^{15}N relaxation data (Figure S5), S² order parameters and MD rms fluctuations (Figure S6), and MD frames (Figure S7). This material is available free of charge via the Internet at http://pubs.acs.org.

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