X-ray Crystallographic Analysis of MinC N-Terminal Domain from *Escherichia coli*

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Abstract—MinC plays an important role in bacterial cell division system by inhibiting FtsZ assembly. However, the molecular mechanism of the action is poorly understood. *E. coli* MinC Nterminus domain was purified and crystallized using 1.4 M sodium citrate pH 6.5 as a precipitant. X-ray diffraction data was collected and processed to 2.3 Å from a native crystal. The crystal belonged to space group $P2_12_12_1$, with the unit cell parameters a = 52.7, b = 54.0, c = 64.7 Å. Assuming the presence of two molecules in the asymmetric unit, the Matthews coefficient value is 1.94 ų Da⁻¹, which corresponds to a solvent content of 36.5%. The overall structure of MinC^N is observed as a dimer form through anti-parallel β-strand interaction.

Keywords—MinC, Cell division, Crystallization.

I. INTRODUCTION

N bacteria, cytokinesis is controlled by cytokinetic ring, Lcalled as the Z-ring, which recruits other cell-division proteins [1]-[3]. Z-ring consists of FtsZ filaments and is normally located at the midcell in cell division by attaching to the membrane through membrane-associated proteins ZipA and FtsA. However, Z-ring is not limited to midcell and also occurs near the cell polar region in the absence of the min system [4]. Successful cell division in Escherichia coli requires the mutual cooperative action of the Min proteins, MinC, MinD and MinE [5]-[6]. MinC interacts with MinD to become a division inhibitor that block polar division by destabilizing FtsZ polymers at polar region while oscillating from one polar region to the other polar region in a cell [7]-[8]. MinD concentrates MinC at the membrane, which enhances the inhibitory activity of MinC up to 50-fold [9, 10]. However, MinE inhibits MinCD by dissociating the MinCD complex near the midcell division site to prevent the disassembly of FtsZ polymers [11]-[12]. Based on these behaviors between Min proteins, Z-ring is stably positioned at the midcell division site and a cell could be equally divided into two daughter cells.

MinC is composed of two functional domains, which are both of N-terminal domain (MinC^N) that blocks FtsZ assembly and C-terminal domain (MinC^C) that interacts with MinD and binds to FtsZ to antagonize the bundling of FtsZ filaments as well [13]-[15]. Thus, the studies on MinC affecting FtsZ assembly have been actively reported. Nevertheless, the molecular mechanism of MinC that inhibits FtsZ assembly hasn't been understood yet. Currently, only two MinC structures have been known. Reported crystal structures of full

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length MinC from T. maritima (PDB code 1hf2) and MinC^N from S. typhimurium (PDB code 3ghf) show different dimerization mode. To provide structural evidence underlying dimer interaction by the MinC^N from E. coli as a model system, we purified and crystallized the dimeric MinC^N from E. coli.

II. MATERIALS AND METHODS

Expression and purification

The recombinant gene was cloned from E. coli (ATCC No.700926D-5) genomic DNA by PCR technique using the primer pair (5'-GTC AAC GGA TCC ATG TCA AAC ACG CC-3' and reverse primer 5'-CCC TTC CTC GAG TCA CGC TTT TTC CTT-3'), where underlined letters indicate nucleotide that were introduced to generate the restriction sites and the end codon is described as bold letters. The DNA was inserted into the modified pET-28a vector using BamHI/XhoI site. The recombinant plasmid was transformed into E. coli strain BL21 (DE3) for overexpression of protein. The cells were grown in LB medium using 50 µg/ml kanamycin at 310 K until OD₆₀₀ reached approximately 0.7 and expression of the recombinant protein was induced by the addition of 0.5 mM Isopropyl β-D-1-thiogalactopyranoside (IPTG) followed by 9 h incubation at the same temperature. The cells were obtained by centrifugating at 4,500g for 15 min at 277 K. The cell pellet was resuspended with buffer A containing (50 mM sodium phosphate pH 8.0, 300 mM NaCl, 5 mM imidazole) and lysed by sonication. The crude lysate was centrifuged at 16,000g for 50 min at 277 K. The supernatant was loaded onto a Ni-NTA resin (Peptron) that had been previously equilibrated and washed with buffer A. The protein was eluted with buffer B containing (50 mM sodium phosphate pH 8.0, 300 mM NaCl, 300 mM imidazole). The eluate was concentrated using Centriprep YM-3 (Millipore) and size exclusion chromatography was performed with Superdex 200 16/60 column (GE Healthcare, USA) that had been equilibrated with buffer C containing (20 mM Tris-HCl pH 8.0, 150 mM NaCl, 1 mM DTT). The eluates were finally concentrated to 14 mg/ml with Centriprep YM-3 (Millipore) for crystallization.

Crystallization and X-ray analysis

Initial crystallization trial was performed with Crystal screen I and II (Hampton Research, USA). Crystallization conditions were further refined by the hanging-drop vapor-diffusion method at 294 K, mixing 1 μ l of protein solution with 1 μ l of reservoir solution.

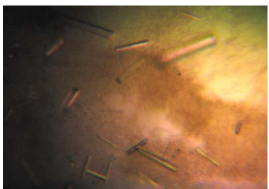


Fig. 1 Native crystals of MinC^N from *E. coli* for X-ray data-collection. The maximum dimensions of crystal are approximately $0.05 \times 0.05 \times 0.3$ mm.

The best crystals were observed after 3 days in reservoir solution consists of 1.4 M sodium citrate pH 6.5 (Figure 1). The crystal was immersed into paratone as a cryoprotectant before flash-freezing in liquid nitrogen for X-ray diffraction experiment. X-ray diffraction data was collected on beamline 4A (MXW) of the Pohang Accelerator Laboratory (Pohang, South Korea). The crystal-to-detector distance was set to 300 mm and a total of 300 images were recorded with an oscillation angle of 1.0°, an exposure time of 10 seconds per image and a wavelength of 1.0000 Å. The data set(Table 1) was processed and scaled using *HKL*-2000 [16].

Structure determination and refinement

The native crystal has space group $P2_12_12_1$ and unit cell parameter a=52.7, b=54.0, c=64.7 Å. Assuming the presence of two molecules in the asymmetric unit, the calculated Matthews coefficient value is $1.94 \text{ Å}^3 \text{ Da}^{-1}$, which corresponds to a solvent content of 36.5% [17]. The crystal structure of $E.\ coli\ \text{MinC}^{\text{N}}$ was solved by the molecular-replacement method using the program PHASER [18] with that of $Salmonella\ typhimurium\ \text{MinC}$ (PDB code 3ghf) as a search model. To Refine the structure of $E.\ coli\ \text{MinC}^{\text{N}}$, Many cycles of manual rebuilding using the program COOT[19] and refinement using the program CNS[20] and Refmac 5[21] were performed (Table 1).

III. RESULTS AND DISCUSSION

Overall structure of MinC^N

The structure of MinC N-terminal domain has been solve by MR method using that of *Salmonella typhimurium* MinC (PDB code 3ghf) as a search model. The final R value is refined to an $R_{work} = 24\%$, $R_{free} = 27.4\%$. After determining the structure, it has been shown that the overall structure is a dimer form that is associated with each other through antiparallel β -strand interaction, which is correlated with that size-exclusion chromatography showed that MinC^N forms a stable dimer in buffer solution (data not shown). Monomer of MinC^N has three α -helices which are composed of α 1 (residues 25-43), α 2 (residues 61-71), α 3 (residues 84-93) and a three-stranded β sheet that consist of β 1 (residues 6-20), β 2 (residues 47-51), β 3 (residues 77-79).

Putative polymer model structure of MinC

Bacterial cell has an unque cell division system to do identical cell division. Z-ring is located to mid-site to perfom properly the cell division. But, Z- ring also can move to polar zones. MinC inhibits Z-ring near polar zones to prevent the result in a cell. In that point, MinC plays a important role in Z-ring mid-position. As mentioned above, the structure of MinC^N has been revealed as a dimer. Based on the aspect of MinC^N interactions, we could think that individual MinC molecules are strongly associated with each other through Nterminal domain when MinC polymer should be formed (Figure 3). And it would be thought that dimer of MinC^N is a effective form to fufill optimally because MinC^N would carry out the work by grabbing interfaces between subunits when MinC should attack the longitudinal bond between FtsZ subunits to perform its role. Additionally, it is associated with each other through anti-parallel B-strand interaction, which means mighty connection. Therefore, it would be stably maintained as dimer form regardless of that the interface between MinC^N are opened or closed even though plenty of motility occurs to inhibit Z-ring.

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 $\label{eq:table I} Table \ I$ Data collection and refinement statistics for \textit{E.coli} MinC^N

	N.
Dataset	$MinC^N$
X-ray source	PAL-4A
Wavelength (Å)	1.0000
Space group	$P2_12_12_1$
Unit cell dimensions (Å)	a =52.7, b=54.0, c=64.7
Resolution (Å) ^a	50-2.30 [2.34-2.30]
Observed reflections	99925
Unique reflections	8799
Multiplicity	11.3 [10.2]
Completeness (%)	99.8 [98.8]
R_{merge}^{b} (%)	6.5 [41.5]
$I/\sigma(I)$	13.8 [4.6]
Refinement	
$R_{\text{work}}^{\text{c}}$ total (%)	24
$R_{\rm free}$ d total (%)	27.4
R.m.s. bond length (Å)	0.008
R.m.s. bond angle (°)	1.7
Average B value (Å2)	52.1

^a Values in brackets are for the highest resolution shell.

^dRfree calculated with 10% of all reflections excluded from refinement stages using high resolution data.

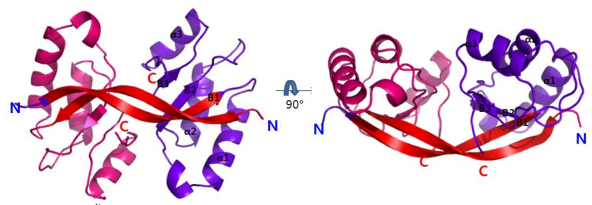
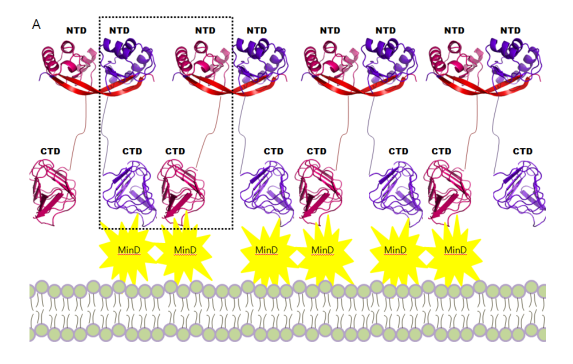


Fig. 2 Structure of MinC^N from E. coli. N-terminal domains are connected to each other through anti-parallel β -strand interaction. Each monomer consists of three helices and a three-stranded β sheet.

 $^{{}^}bR_{\text{merge}} = \sum_{hkl} \sum_i |I_i(hkl) - \langle I(hkl) \rangle | / \sum_{hkl} \sum_i I_i(hkl)$, where $I_i(hkl)$ is the intensity of the *i*th observation of reflection *hkl* and $\langle I(hkl) \rangle$ is the average intensity of reflection *hkl*.

^c Rwork = $\Sigma \mid |Fo| - |Fc| \mid /\Sigma |Fo|$



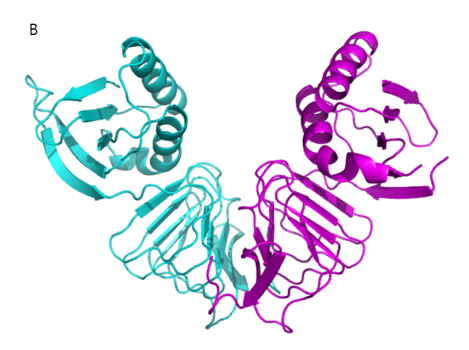


Fig. 3 (A) Putative polymer structure model of MinC from *E. coli*. black square area means dimer form of putative MinC structure. CTD region of MinC is modeled from *T. maritima* MinC. (B) Structure of MinC(PDB code 1hf2) from *T. maritima*