BL-5A, 17A, NW12A, NE3A/2019G548

Crystal structures of enzymes involved in novel lysine biosynthetic pathway using amino-group carrier protein

Takeo TOMITA^{1,2}, Ayako YOSHIDA^{1,2}, Sumire KUROSAWA¹, and Makoto NISHIYAMA^{1,2,*}

¹ Agro-Biotechnology Research Center, The Univ. of Tokyo, Yayoi 1-1-1, Bunkyo-ku, Tokyo, 113-8657, Japan

²CRIIM, The Univ. of Tokyo, Yayoi 1-1-1, Bunkyo-ku, Tokyo 113-8657, Japan

1 Introduction

Lysine biosynthetic pathway is classified into two types; diaminopimelate (DAP) pathway in bacteria and plants, and α-aminoadipate (AAA) pathway in fungi and yeast. We previously found that a bacterium, Thermus thermophilus synthesizes lysine via a novel type of AAA pathway using an amino-group carrier protein called LysW. LysW protects the α-amino group of AAA. LysW is highly acidic protein and our previous studies including crystallographic analyses revealed that LysW functions not only as an aminogroup protecting group but also as a carrier protein by interacting with each enzyme through electrostatic interactions. Therefore, we hereafter call LysW as amino-group carrier protein (AmCP). Since our discovery of AmCP-mediated lysine biosynthesis through AAA in T. thermophilus, evidences have mounted that many microorganisms including thermophilic bacteria and archaea synthesize lysine and also arginine by a similar pathway, indicating that this pathway is one of evolutionary origins of the amino acid biosynthetic pathways.

Recently, we found that a kind of *Streptomyces* used AmCP in the biosynthesis of secondary metabolites which containing non-proteinogenic amino acid (2S,6R)-diamino-(5R,7)-dihydroxyheptanoic acid, DADH [1]. DADH is incorporated into a novel peptide metabolite, vazabitide A, featuring an azabicyclo hexane ring structure, by nonribosomal peptide synthetases and successive modification enzymes.

We also found that AmCP was also used for the biosynthesis of a compound, s56-p1, produced by another strain of *Streptomyces* [2]. s56-p1 contains a unique nitrogen-nitrogen bond (hydrazine moiety) fused with the DADH-derivative moiety. We identified the unprecedented machinery for hydrazine moiety in the strain. Considering that this machinery is widespread across several bacterial phyla, this study highlights the overlooked potential of bacteria to synthesize hydrazine.

Thus, to reveal the structural bases of these unprecedented biosynthetic pathway is important for the further understanding of AmCP-mediated amino acid biosynthetic machinery. Here, we describe the trial to determine the crystal structures of enzymes involved in these pathways to reveal the structural

basis of this unprecedented biosynthetic pathway including AmCP.

2 Experiment

Purification of recombinant proteins

Complex of OrfX and OrfY which are involved in the biosynthesis of vazabitide A, and OrfZ which is in the biosynthesis of s56-p1 were overexpressed in *E. coli* Rosetta2 (DE3) and *E. coli* BL21-Codon-Plus (DE3)-RIL as a host, respectively. In order to apply S-SAD (sulfur single-wavelength anomalous dispersion) method, we overexpressed OrfX/Y in *E. coli* B834 (DE3) as a host. From cell lysates prepared by sonication, Complex of OrfX/Y were purified through Ni²⁺-NTA column and OrfZ was purified through streptactin column, following by gel filtration chromatography (Superdex 200) for crystallization.

Crystallization of Complex of OrfX and OrfY, and OrfZ

The condition for crystallization of Complex of OrfX/Y, and OrfZ were screened with Crystal screen I and II, Wizard classic I, II, and III, and PEG/ION by hanging drop vapor diffusion method at 20 C. Optimization of the crystallization condition was also conducted by modifying the concentration and pH.

3 Results and Discussion

X-ray diffraction analyses of complex of OrfX and OrfY

We found the crystallization condition of complex of OrfX/Y. We carried out X-ray diffraction analyses, and obtained dataset at 1.7 Å resolution. Because these proteins have no similarity to a structural known protein, we applied S-SAD method and determined the phase. The obtained data set was at 1.9 Å resolution, and we determined the crystal structure. Furthermore, we have determined the crystal structure binding with a proposed substrate analog by data set at 2.7 Å resolution. The optimization of the crystallization condition is now in progress to improve the resolution.

X-ray diffraction analysis of OrfZ

We have not found the crystallization condition of OrfZ. Crystallization of OrfZ with different construction is now in progress.

References

- [1] Hasebe, F. et al., Nat. Chem. Biol. **12**, 967 (2016). [2] Matsuda, K., J. Am. Chem. Soc. **140**, 9083 (2018).

^{*} umanis@g.ecc.u-tokyo.ac.jp