

## X-ray structure of L-lactate oxidase from *Enterococcus hirae* in complex with a product “D-lactate form ligand”

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### 1 Introduction

L-Lactate oxidase (LOx) belongs to a family of flavin mononucleotide (FMN)-dependent  $\alpha$ -hydroxyacid oxidizing flavoproteins. To date, the structures of LOx and mutagenesis studies have been reported, and its substrate recognition mechanism has also been elucidated, however, its substrate/product inhibition mechanisms are yet to be elucidated. In this study, we determined the X-ray structure of LOx derived from *Enterococcus hirae* (EhLOx) in complex with “D-lactate form ligand”, which was covalently bonded with Tyr211. This structure is the direct evidence to support a new hypothesis of the product-inhibition mechanism of LOx [1].

### 2 Experiment

The recombinant EhLOx was used for crystallization. Crystals were obtained in a droplet containing a mixture of 0.8  $\mu$ l protein solution (10–12 mg/ml in 20 mM potassium phosphate buffer pH 7.0) and 0.8  $\mu$ l reservoir solution (0.1 M Tris-HCl, pH 8.0, 30% (w/v) Polyethylene glycol monomethyl ether 2,000) in a well containing 50  $\mu$ l reservoir solution using the sitting-drop method at 293 K. To obtain the structure in complex with substrate, protein solution containing 0.4 M sodium pyruvate was used in the above crystallization condition.

X-ray diffraction data were collected on the PF BL-5A in the KEK, and processed using the programs XDS and the CCP4 suite. The structure was determined by molecular replacement with the program MOLREP using the structure of LOx from *Aerococcus viridans* (PDB ID 4YL2).

### 3 Results and Discussion

The structure of EhLOx in complex with D-lactate form ligand was determined at 1.7  $\text{\AA}$  resolution. EhLOx forms a homotetramer (Fig. 1). Each monomer of EhLOx has  $(\beta/\alpha)_8$  TIM-barrel fold with FMN, same as other  $\alpha$ -hydroxyacid oxidizing flavoproteins. In the complex structure of EhLOx, the whole flexible loop region (187–216) was ordered. The simulated-annealing omit maps (sa-omit maps) contoured at  $4\sigma$  revealed the presence of the strong and continuous electron density maps from C $\alpha$  of the ligand to the hydroxyl group of Tyr211 (Fig. 2).

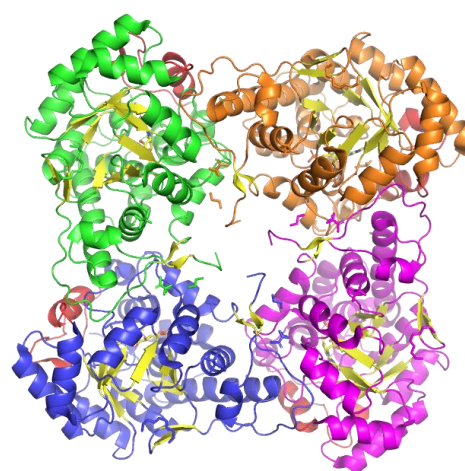


Fig. 1: Overall structure of homotetrameric EhLOx in complex with D-lactate form ligand. The flexible loop regions (187–218) are colored in red and all  $\beta$ -strands are colored in yellow.

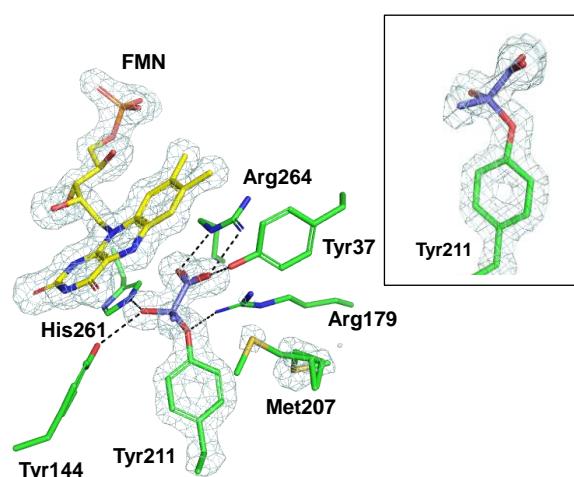


Fig. 2: Structure of EhLOx active site in complex with D-lactate form ligand (purple stick).

This observation revealed that C $\alpha$  of the ligand was mainly bonded to Tyr211 in this complex structure, and the bound ligand does not fit an  $sp^2$  configuration of pyruvate, but instead adapts  $sp^3$  configuration of a C $\alpha$  of D-lactate. Considering the additional covalent bond, the bound ligand was refined as “D-lactate” and hence called the ligand the “D-lactate form ligand.” In this binding mode, the nucleophilic  $sp^3$  N5 of reduced FMN cannot attack on  $\alpha$ -proton of D-lactate form ligand because it orients to methyl group. Therefore, the observed D-lactate form ligand could bind in the active site as an inhibitor in EhLOx.

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#### Reference

[1] Hiraka, K. et al., Protein Science. 31, e4434 (2022).

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