# MS6-05 Cooperative Motion Analysis of group II chaperonins at single molecule level using nanocrystal and time-resolved diffraction measurement

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Group II chaperonin, found in archaea and in the eukaryotic cytosol, is an indispensable protein that captures a nonnative protein and refolds it to the correct conformation in an ATP dependent manner. ATP-induced structural changes are essential for chaperonin activity and we had reported that the diffracted X-ray tracking (DXT) method could trace ATP induced conformational change of group II chaperonin at single molecule level (Sekiguchi et al., PLoS ONE 2013). In DXT, nanocrystal immobilized on one side of chaperonin-ring is used as motion tracer for structural change of chaperonin as shown in the figure. In this study, we analyzed how ATPase deficient mutant modulate dynamic motion of chaperonin, and cooperativity inter and intra communication of chaperonin ring. We found that although one ring of the asymmetric ring complex lacks ATPase activity, the other wild-type ring undergoes an ATP-dependent conformational change and maintains its protein-folding activity. The results clearly demonstrate that inter-ring communication is dispensable in the reaction cycle of group II chaperonins (Yamamoto et al., JMB 2014), despite the reaction cycle of group I chaperonins, for example GroEL-ES system in E-coli, is controlled by inter-ring allosteric communication.

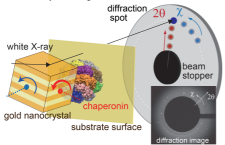


Figure 1. Schematic illustration of motion analysis of chaperonin by diffracted X-ray tracking.

### Keywords: nanocrystal chaperonin dynamics

# MS7 Protein & glycobiology structure determination

Chairs: Jon Agirre, Gerlind Sulzenbacher

## MS7-01 Structures of lytic polysaccharide monooxygenases and their interaction with polysaccharide substrates

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The recent discovery of lytic polysaccharide monooxygenases has revolutionized our understanding of how polysaccharides are degraded in nature and our thoughts on how they could be most efficiently degraded in industrial processes. Cellulose, the main constituent of the plant cell wall, is the most abundant polysaccharide on earth, and as it is present in all plants is of particular of interest for the production of second generation biofuels. Conventional cellulases are glycoside hydrolases, and catalyze the cleavage of the glycosidic linkage by using water molecule in mechanisms involving one or more carboxylate residues. In contrast, lytic polysaccharide monooxygenases (LPMOs) are metalloenzymes which cleave the glycosidic linkage by activating molecular oxygen through a mononuclear copper center held by the so-called histidine brace, thereby mediating oxidation at C1 or C4 (1,2).

Carbohydrate modifying enzymes are classified in the Carbohydrate Active Enzyme Database, CAZY, where LPMOs are now classified as Auxiliary Activities (3) in families AA9 (GH61 prior to discovery of their LPMO activity (4)), AA10 (CBM33 prior to discovery of their LPMO activity), AA11 and AA13. The initially characterized LPMOs had activity on the insoluble β-1,4-linked polysaccharides chitin and cellulose, but LPMOs active on soluble 6-1.4-linked now oligo-/polysaccharides substrates, as well as active on polysaccharides,  $\alpha$ -1.4-linked have heen both characterized biochemically and structurally (5-7).

This talk will provide an overview on the structural biology of LPMOs so far and focus primarily on substrate binding and specificity, in particular the structural differences between  $\alpha$ -1,4 and  $\beta$ -1,4- active LPMOs (5) and the very first LPMO substrate complexes which we recently obtained (7).

The results are largely from the CESBIC consortium collaborative project funded by the ERA-IB initiative, which additionally to the University of Copenhagen included Novozymes A/S, the University of Cambridge, University of York and the CNRS Marseille.

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Keywords: carbohydrate active enzyme; biofuel; lytic polysaccharide monooxygenase; metalloenzyme

## MS7-02 Membrane Enzymes: the Structural basis of Phosphatidylinositol Mannosides Biosynthesis in Mycobacteria

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Membrane enzymes constitute a large class of proteins with critical roles in a variety of cellular processes in all living organisms. They generate a significant amount of structural diversity in biological systems, which are particularly apparent in cell-pathogen interactions. Many of these enzymes are required to access a lipophilic substrate located in the membranes and to catalyze its reaction with a polar, water-soluble compound.<sup>[1]</sup> Here we focus on the membrane enzymes involved in the early steps of the phosphatidylinositol mannosides (PIMs) biosynthetic pathway, essential structural components of the cell envelope of Mycobacterium tuberculosis. Of particular relevance, we demonstrate the occurrence of a conformational switch during the catalytic cycle of the retaining glycosyltransferase PimA, the enzyme that start the pathway, involving both  $\beta$ -strand-to- $\alpha$ -helix and  $\alpha$ -helix-to- $\beta$ -strand transitions. <sup>[2, 3]</sup> These structural changes seem to modulate catalysis and are promoted by interactions of the protein with anionic phospholipids in the membrane surface. Our studies demonstrate that protein-membrane interactions might entail unanticipated structural changes in otherwise well conserved protein architectures, and suggests that similar changes may also play a functional role in other membrane-associated enzymes. Finally, we report the crystal structures of PatA, an essential membrane acyltransferase that transfers a palmitoyl moiety from palmitoyl-CoA to the 6-position of the mannose ring added by PimA, in the presence of its naturally occurring acyl donor palmitate and a nonhydrolyzable palmitoyl-CoA analog. The structures reveal an  $\alpha/\beta$  architecture, with the acyl chain deeply buried into a hydrophobic pocket that runs perpendicular to a long groove where the active site is located. Enzyme catalysis is mediated by an unprecedented charge relay system, which markedly diverges from the canonical HX D motif. Our studies establish the mechanistic basis of substrate/membrane recognition and catalysis for an important family of acyltransferases, providing exciting possibilities for inhibitor design. [4]

## References

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Keywords: membrane enzymes, glycosyltransferase, acyltransferase, glycobiology, mycobacterium