Biosynthesis of mycobacterial methylmannose polysaccharides requires a unique 1-O-methyltransferase specific for 3-O-methylated mannosides

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Mycobacteria are priority pathogens in terms of drug resistance worldwide and efforts aimed at deciphering their unique metabolic pathways and unveiling new targets for innovative drugs should be intensified. In particular, nontuberculous mycobacteria (NTM) are environmental organisms increasingly associated to opportunistic infections [1] and known to produce methylmannose polysaccharides (MMP). MMP have been implicated in the metabolism of precursors of cell envelope lipids crucial for stress resistance and pathogenesis. Although the functions of MMP remain to be confirmed experimentally, their tight interactions with fatty acids are intrinsically associated to unique and extensive methylation patterns, resulting from the action of hitherto uncharacterized methyltransferases.

In this work, we identified and characterized biochemically a novel mycobacterial methyltransferase (MeT1) that specifically blocks the non-reducing end of a MMP precursor. We crystallized and determined the first X-ray structure of the SAM-dependent MeT1 from M. haisiacum in complex with magnesium and its exhausted cofactor, SAH. In particular, the three high-resolution 3D structures (in space groups P321 and C221; PDB entries 6H40, 6G7D and 6G80) in combination with SAXS data (SASBDB entry SASDDJ6) unveiled a dimeric arrangement of the enzyme in solution and a highly flexible lid important for its catalytic cycle. This structural information, together with molecular docking simulations, allowed the elucidation of the enzyme’s reaction mechanism, furthering our knowledge of MMP biosynthesis and providing important tools to dissect the role of MMP in NTM physiology and resilience [2].


Keywords: Mycobacterium; polymethylated polysaccharides; methyltransferase; S-adenosyl-L-methionine; 3D structure

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