Title: Galactosyl Transferases in Mycobacterial Cell Wall Synthesis

Running title: Biosynthesis of Mycobacterial Galactan

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ABSTRACT

Two galactosyltransferases can apparently account for the full biosynthesis of the cell wall galactan of mycobacteria. Evidence is presented based on enzymatic incubations with purified natural and synthetic galactofuranose (Gal\(^f\)) acceptors that the recombinant galactofuranosyl transferase, GlfT1, from *Mycobacterium smegmatis*, the *M. tuberculosis* Rv3782 ortholog, known to be involved in the initial steps of galactan formation, harbors dual β-(1→4) and β-(1→5) Gal\(^f\) transferase activities and that the product of the enzyme, decaprenyl-P-P-GlcNAc-Rha-Gal\(^f\)-Gal\(^f\), serves as a direct substrate for full polymerization catalyzed by another bifunctional Gal\(^f\) transferase, the GlfT2, the Rv3808c enzyme.
The mycobacterial cell wall - including the essential, covalently linked complex of peptidoglycan, heteropolymeric arabinogalactan and highly hydrophobic mycolic acids - is responsible for many of the pathophysiological features of members of the *Mycobacterium* genus (9). Several antituberculosis drugs affect the formation of this complex. Isoniazid, ethionamide, thiocarlide and thiacetazone inhibit mycolic acid synthesis (14, 29, 30, 36, 38), ethambutol specifically disrupts the synthesis of arabinan (20, 24, 35, 39), and D-cycloserine, an inhibitor of peptidoglycan synthesis (11), has some clinical utility. However, drug resistance, particularly the multiple (MDR) and extensive (XDR) forms are of pressing public health concern (15, 31), presaging the need for a broader array of targets and drugs affecting both cell wall synthesis and other aspects of mycobacterial metabolism. Yet, our understanding of the synthesis of the mycobacterial cell wall is elementary compared to that of other bacteria.

The initiation point for arabinogalactan biogenesis is the mycobacterial version of the bacterial carrier lipid, bactoprenol, identified as decaprenyl phosphate (C\(_{50}\)-P) (10), and the sequential addition of GlcNAc-P, rhamnose (Rha), and single galactofuranose (Gal\(f\)) units, donated by the appropriate nucleotide sugars (23, 25) (Fig. 1). At some stage the arabinofuranose (Ara\(f\)) units are added, donated, not by a nucleotide sugar, but a lipid carrier, C\(_{50}\)-P-Ara\(f\) (20, 39). Several of the responsible glycosyl transferases taking part in this process have been identified (1, 3, 5, 18, 19, 25, 26, 32, 34).

We recently described the galactofuranosyl transferase, Rv3782, responsible for attaching the first and, perhaps, the second Gal\(f\) unit to the C\(_{50}\)-P-GlcNAc-Rha acceptor (22). Due to its role in the initiation of galactan formation we now name it galactofuranosyl transferase 1 (GlfT1). Previously, yet another galactofuranosyl transferase, Rv3808c (originally called GlfT but now named GlfT2), was recognized and proved to be bifunctional in that it was responsible for the formation of the bulk of the galactan, containing alternating
5- and 6-linked β-GalF units (19, 25, 32). In the present study, we examine the precise roles of GlfT1 and GlfT2 in mycobacterial cell wall galactan synthesis through the application of in vitro reactions with purified natural acceptors, and synthetic products emulating the natural substrates, and the recombinant enzymes expressed in Escherichia coli.

Cloning, expression and activity of the GlfT1 ortholog from *M. smegmatis*.

Attempts to produce a soluble form of the *M. tuberculosis* Rv3782 enzyme in *E. coli* were unsuccessful, apparently due to toxic effects, and the yields of pure active protein from an overproducing strain of *M. smegmatis* were too low to allow further biochemical studies. Instead the *M. smegmatis* ortholog, corresponding to the gene MSMEG_6367 (www.tigr.org), was cloned and expressed in *E. coli*; the two orthologous GlfT1 proteins are similar to the extent of 90.7% and their sequence identity is 76.8% (Clustal X, version 1.8). The *glfT1* genes are located within the conserved arabinogalactan biosynthetic regions in *M. smegmatis* mc²155 and *M. tuberculosis* H37Rv (4), in the center of predicted operons comprised of three genes, where the first one encodes a nucleotide binding protein (MSMEG_6366 and Rv3781 orthologs) and the last one a membrane spanning protein (MSMEG_6369 and Rv3783 orthologs) of the ABC (ATP-Binding Cassette) transport type (6). The *glfT1* gene from *M. smegmatis* mc²155 was amplified based on the oligonucleotide primers, 5'-GCCACCAACATATGACGCACACTGAGGTCGTCTG -3' and 5'-CCCAAGCTTTTCATCGCTGGAACCTTCGCGTC -3', containing Hind III and Nde I restriction sites. The PCR fragment (0.93 kb) was digested, and ligated into the similarly digested pVV2 and pET28a vectors for expression in *M. smegmatis* mc²155 and *E. coli* BL21 (DE3) (13). Production of the MSMEG_6367 protein with the N-terminal 6-histidine fusion tag, was confirmed by SDS-PAGE and Western blot. Assays for GalF transferase activity (22) with the recombinant MSMEG_6367 protein overexpressed in *M. smegmatis* demonstrated increased synthesis of C₅₀-P-P-GlcNAc-Rha-GalF -GalF (glycolipid-4; GL-4) (data not shown),
confirming identical functions for the two orthologous GlfT1 proteins, Rv3782 and MSMEG_6367.

**GlfT1 has dual β-(1→4) and β-(1→5) galactofuranosyl transferase activities.**

Previous studies had shown that GlfT1 from *M. tuberculosis* could catalyze the synthesis of both GL-3 (C₅₀-P-P-GlcNAc-Rha-Galᵢ) and GL-4, attributing both β-(1→4) and β-(1→5) galactofuranosyl transferase activities to the one enzyme (22). However, efforts to directly confirm this dual activity with the purified putative substrates (GL-2 and GL-3) and purified recombinant Rv3782 enzyme were not successful (22). In this present instance, the overproducing strain, *E. coli* BL21(DE3)/pET28a-MSMEG_6367, and control culture with the empty vector, were induced with IPTG, the cells harvested and disintegrated by probe sonication as described (22), and the supernatant from centrifugation was used as a source of GlfT1. The synthetic products, Ac-2, Ac-3 and Ac-4 (Fig. 2), analogs of GL-2, GL-3 and GL-4, served as putative acceptors. In addition to the above-mentioned substrates at 4 mM concentration each, the reaction mixtures contained 6.25 mM NADH, 62.5 µM ATP, UDP-[U-¹⁴C]Galp (278 mCi/mmol; 0.25 µCi), 20 µg of UDP-Galp mutase (25), and 50 µl aliquots of the cell lysate from control or overproducing strain of GlfT1, in a final volume of 80 µl (22). The reactions were stopped by the addition of CHCl₃/CH₃OH (2:1) to yield the organic phase of a biphasic solution (16). Thin layer chromatography (TLC) and autoradiography of the reaction products revealed that both Ac-2 and Ac-3, emulating GL-2 and GL-3, were efficient substrates for recombinant GlfT1, converting them to radiolabelled Ac-3 and Ac-4, respectively (Fig. 3A). However, Ac-4, reflective of GL-4, was not effective as an acceptor of [¹⁴C]Gal under the reaction conditions (results not shown), indicating that GlfT1 could not catalyze subsequent galactan chain extension. Moreover, when the native glycolipids, GL-2, GL-3, GL-4 and GL-5 (isolated as described in ref. 22) were incorporated into similar assays, both GL-2 and GL-3 were readily converted to GL-4 (Fig. 3B); however, as in the case of the
synthetic acceptor Ac-4, GL-4 was not an effective substrate for GlfT1. Similarly, GL-5 did not serve as an acceptor for galactofuranosyl transfer catalyzed by GlfT1 (Fig. 3B).

**GlfT2 catalyzes the formation of GL-5 and further galactan polymerization.**

Following recognition of the role of GlfT1 in GL-3 and GL-4 synthesis and previous evidence that GlfT2, the Rv3808c enzyme, can catalyze the formation of the C$_{50}$-P-P-GlcNAc-Rha-linked galactan polymer with its inherent alternating β-(1→45) Gal$_f$ and β-(1→6) Gal$_f$ units, the question arose as to the initiation of the polymerization events. Specifically, could GL-4 and its synthetic analogue Ac-4 give rise to GL-5/Ac-5 and the subsequent polymer? We investigated this question through enzyme incubations containing GlfT2 from *M. tuberculosis* overexpressed in *E. coli* (32), radiolabeled GL-4 or GL-5 (2000 dpm each) and 0.2 mM cold UDP-Galp instead of the radioactive counterpart. Extraction of the reaction products with CHCl$_3$/CH$_3$OH (2:1) showed consumption of the original substrates GL-4 and GL-5 in the reaction mixtures containing the overexpressed GlfT2 (Fig. 4A). Further extraction with solvents more polar than CHCl$_3$/CH$_3$OH (2:1), namely CHCl$_3$/CH$_3$OH/H$_2$O (10:10:3) (33) and "E-soak" (water/ethanol/diethyl ether/pyridine/ammonium hydroxide, 5:15:5:1:0.017) (2), removed the C$_{50}$-P-P-GlcNAc-Rha-based galactan polymers which were resolved by SDS-PAGE and visualized by autoradiography (Fig. 4B). Clearly GlfT2 catalyzed the conversion of GL-4 to GL-5, and both substrates were suitable for galactan polymerization (Fig. 4B).

When purified GL-2 or GL-3 was used in a similar experiment, galactan build-up did not take place (data not shown). Moreover, Ac-4 and Ac-5, the synthetic equivalents of GL-4 to GL-5, proved to be ready acceptors of [$_{14}$C]Gal provided by UDP-[$_{14}$C]Galp in the standard reaction mixture containing recombinant GlfT2 (222), whereas Ac-2 or Ac-3 were not effective substrates (Table 1). Therefore these data further support the case that GlfT2 catalyzes the addition of the third and subsequent Gal$_f$ residues, but not the first two, to the lipid template in galactan synthesis.
Conclusions

Experimental evidence presented in this paper resolves several questions raised in our previous work (22, 32). Firstly, it shows that the galactosyltransferase, GlfT1, from M. smegmatis, the Rv3782 ortholog, initiates galactan synthesis on the C$_{50}$-P-P-GlcNAc-Rha acceptor and is endowed with dual, β-(1→4) and β-(1→5) Gal$_{f}$ transferase activity. Secondly, the data strongly suggests that the action of GlfT1 is directly followed by that of another bifunctional enzyme GlfT2, the Rv3808c protein, apparently responsible for the majority of galactan polymerization. These enzymes thus represent further examples of a growing number of glycosyl transferases that harbor two distinct transferase activities on a single polypeptide chain (7, 8, 12, 19, 25, 27, 28, 32, 37), belying the early “one enzyme, one linkage” hypothesis (17). Thus it would seem from present evidence that complete synthesis of mycobacterial galactan can be catalyzed by but two enzymes.

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REFERENCES


**Figure legends**

**Figure 1.** Pathway for the biosynthesis of mycobacterial arabinogalactan.

The sequence of reactions is based on identification of GL-1 through GL-5 and the lipid-linked arabinogalactan polymer in cell-free systems containing mycobacterial membranes and cell wall fractions (21-23, 25). GlcNAc-1-P transferase WecA (Rv1302) is proposed to be responsible for step 1 (21); the rhamnosyl transferase WbbL (Rv3265) catalyzes step 2 (26). Subsequent reactions 3 and/or 4 are proposed in this study and previously (21) to be catalyzed by Rv3782. Step 6 represents a series of galactofuranosyl additions catalyzed by Rv3808c. Several arabinosyl transferases involved in reactions under step 7 have been described, such as the Emb proteins (3), AftA (1) and AftB (34).

**Figure 2.** Structures of the synthetic acceptors Ac-2, Ac-3, Ac-4 and Ac-5.

Oligosaccharide portions of Ac-2 to Ac-5 match the structures of the natural intermediates in galactan biosynthesis, namely GL-2, GL-3, GL-4 and GL-5.
Octyl α-L-rhamnopyranosyl-(1→3)-2-acetamido-2-deoxy-α-D-glucopyranoside (Ac-2); octyl β-D-galactofuranosyl-(1→4)-α-L-rhamnopyranosyl-(1→3)-2-acetamido-2-deoxy-α-D-glucopyranoside (Ac-3); octyl β-D-galactofuranosyl-(1→5)-β-D-galactofuranosyl-(1→4)-α-L-rhamnopyranosyl-(1→3)-2-acetamido-2-deoxy-α-D-glucopyranoside (Ac-4); octyl β-D-galactofuranosyl-(1→6)-β-D-galactofuranosyl-(1→5)-β-D-galactofuranosyl-(1→4)-α-L-rhamnopyranosyl-(1→3)-2-acetamido-2-deoxy-α-D-glucopyranoside (Ac-5). Synthesis of these will be described by Lowary et al.

**Figure 3.** Recognition of the dual β-(1→4) and β-(1→5) Galf transferase activities of GlfT1.

The synthetic acceptors Ac-2, Ac-3 (A) and the native substrates, GL-2, GL-3, GL-4 and GL-5 (B) were used as the acceptors of galactofuranosyl residues in the reaction catalyzed by GlfT1 expressed in heterologous host, E. coli. The positions of Ac-2, Ac-3 and Ac-4 substrates on TLC plate were visualized by staining with α-naphtol.

**Figure 4.** Both glycolipids GL-4 and GL-5 serve as substrates for GlfT2.

Natural substrates GL-4 and GL-5 were used as acceptors of galactofuranosyl residues in the reaction catalyzed by GlfT2 expressed in the heterologous host, E. coli. The reaction mixtures were subjected to series of extractions, as described. Panel A: TLC of glycolipid fractions, followed by autoradiography. Panel B: SDS-PAGE of lipid-linked galactan polymer, followed by Western blotting and autoradiography. Lanes labeled “control” represent samples containing host E.coli strain. Lanes labeled “standard” represent fractions extracted from reaction mixture containing membranes and cell wall from M. smegmatis mc²155 and UDP-[U-¹⁴C]Galp as a tracer for production of galactan intermediates (25).
Table 1. Ability of Ac2–Ac5 to serve as acceptors for GlfT2

<table>
<thead>
<tr>
<th>Compound</th>
<th>dpm&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ac2</td>
<td>732</td>
</tr>
<tr>
<td>Ac3</td>
<td>911</td>
</tr>
<tr>
<td>Ac4</td>
<td>18,706</td>
</tr>
<tr>
<td>Ac5</td>
<td>369,378</td>
</tr>
<tr>
<td>no acceptor</td>
<td>985</td>
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</table>

<sup>a</sup>Transfer of radiolabeled Galf to each potential acceptor was measured using recombinant GlfT2 as described previously (31).  
<sup>b</sup>Disintegrations per minute.
Figure 1
Figure 3

A. acceptors (nonradioactive)

B. acceptors (radioactive)
Figure 4

A. acceptors (radioactive)

GL- 4  GL- 5

GL- 3  GL- 4  GL- 5

standard  control  PET-GlfT2  control  PET-GlfT2

B. CHCl₃/CH₃OH/H₂O (10:10:3) extract “E-soak” extract

acceptors (radioactive)

GL- 4  GL- 5  GL- 4  GL- 5

kDa

170  70  30  10  11

standard  control  PET-GlfT2  control  PET-GlfT2  control  PET-GlfT2  control  PET-GlfT2