MtrP, a putative methyltransferase in Corynebacteria, is required for optimal membrane transport of trehalose mycolates.

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**Abstract**

Pathogenic bacteria of the genera *Mycobacterium* and *Corynebacterium* cause severe human diseases such as tuberculosis (*Mycobacterium tuberculosis*) and diphtheria (*Corynebacterium diphtheriae*). The cells of these species are surrounded by protective cell walls rich in long chain mycolic acids. These fatty acids are conjugated to the disaccharide trehalose on the cytoplasmic side of the bacterial cell membrane. They are then transported across the membrane to the periplasm where they act as donors for other reactions. We have previously shown that transient acetylation of the glycolipid trehalose monohydroxycorynomycolate (hTMCM) enables its efficient transport to the periplasm in *Corynebacterium glutamicum* and that acetylation is mediated by the membrane protein TmaT. Here we show that a putative methyltransferase, encoded at the same genetic locus as TmaT, is also required for optimal hTMCM transport. Deletion of the C.
glutamicum gene NCgl2764 (Rv0224c in M. tuberculosis) abolished acetyltrehalose monocorynomycolate (AcTMCM) synthesis, leading to accumulation of hTMCM in the inner membrane and delaying its conversion to trehalose dihydroxycorynomycolate (h2TDCM). Complementation with NCgl2764 normalized turnover of hTMCM to h2TDCM. In contrast, complementation with NCgl2764 derivatives mutated at residues essential for methyltransferase activity failed to rectify the defect, suggesting that NCgl2764/Rv0224c encodes a methyltransferase, designated here as MtrP. Comprehensive analyses of the individual mtrP and tmaT mutants and of a double mutant revealed strikingly similar changes across several lipid classes compared with wild-type bacteria. These findings indicate that both MtrP and TmaT have non-redundant roles in regulating AcTMCM synthesis, revealing additional complexity in the regulation of trehalose mycolate transport in the Corynebacterineae.

Bacteria of the Corynebacterineae, a suborder of the Actinomycetales, include several species pathogenic to humans including Mycobacterium tuberculosis, Mycobacterium leprae and Corynebacterium diphtheriae. The former latently infects around one-quarter of all humans (~1.7 billion) and causes active tuberculosis (TB) disease which is responsible for 1.6 million deaths annually. In 2017, around 10 million people developed TB, mostly in sub-Saharan Africa and Asia. Drug resistance is a significant problem, with ~0.6 million people developing rifampicin-resistant TB in 2017, 82% of which was multi-drug resistant (MDR-TB) (1). The unusually hydrophobic and distinctive cell wall of these bacteria is a validated drug target, and several first line TB drugs, such as isoniazid and ethambutol, target cell wall biosynthetic pathways. An improved understanding of cell wall synthesis processes is required to reveal and exploit new targets.

The Corynebacterineae cell wall confers intrinsic resistance to antibiotics and host immune responses (2,3). The inner layers comprise peptidoglycan and arabinogalactan (AG) as well as covalently linked very long chain fatty acids, the mycolic acids, that form the bulk of an outer membrane (4). These α-alkyl β-hydroxy fatty acids (70-90 carbons long) are either linked to terminal arabinose residues of the AG polymer or are present as free lipids esterified to trehalose (trehalose monomycolate (TMM) and trehalose dimycolate (TDM)), glucose (GMM) or glycerol (GroMM) (5,6). The Corynebacteria also synthesize mycolic acids, the corynomycolates, that are shorter (C22 to C36) and structurally less complex than the mycobacterial fatty acids (7).

The synthetic pathways and enzymatic machineries that produce these species have been extensively studied and reviewed (4,8-11). In Mycobacteria, two independently-synthesized fatty acids, the short alpha branch (C24-C26) and long meromycolate chain (C50-C60), produced by the single FAS-I enzyme and FAS-II complex (12,13), respectively, are modified then condensed by the cytoplasmic polyketide synthase, Pks13 (14), to form a 2-alkyl 3-keto fatty acid which is then reduced by CmrA to generate the mature, hydroxylated mycolic acid (15). These fatty acids are conjugated to trehalose to form TMM which is then bound and flipped to the periplasm by the integral membrane lipid transporter MmpL3 (16,17), the target of several classes of new antimycobacterial inhibitors (18,19). While structurally diverse classes of MmpL3 inhibitors have been reported, several of these
compounds were subsequently found to indirectly inhibit MmpL3, by disrupting the transmembrane electrochemical proton gradient (20). However, recent studies have identified two compounds, BM212 (a 1,5-diarylpyrrole compound and PIPD1 (a piperidinol-containing molecule), that inhibit MmpL3 via direct interactions, advancing our understanding of the mechanism of action of this transporter (16,21). Periplasmic mycolyltransferases of the Ag85ABC complex (22) subsequently utilize TMM in the periplasmic space as the mycolic acid donor for the synthesis of the mycolated-AG layer, trehalose dimycolate (TDM), or mycolated glucose or glycerol.

Most cell wall components are essential for the growth of pathogenic mycobacteria, hampering efforts to functionally characterize genes involved in cell wall assembly. In contrast, Corynebacteria such as \textit{C. glutamicum} retain viability after loss of conserved cell wall components and are thus excellent model systems for investigating cell wall synthesis in Corynebacterineae (15,23-28). Recently, we identified a new trehalose monocorynomycolate (TMCM) glycolipid species in \textit{C. glutamicum}, which contains an acetyl group on the corynomycolate chain. This species, termed acetyl-trehalose monocorynomycolate (AcTMCM), required expression of the putative acetyltransferase, TmaT (NCgl2759) (Fig. 1). Genetic deletion of \textit{tmaT} significantly reduced the rate of corynomycolate transport across the inner membrane and subsequent conversion to h2TDCM, indicating that AcTMCM is the major substrate transported by \textit{Corynebacterineae} CmpL1/4 (NCgl0228/NCgl2769) (28) (Fig. 1).

Here we show that optimal transport of hTMCM is also regulated by a methyltransferase. Specifically, we show that disruption of a putative methyltransferase gene, NCgl2764, which is located in the \textit{tmaT} locus, also leads to the accumulation of hTMCM on the cytoplasmic site of the inner membrane and defects in the synthesis of h2TDCM. Comprehensive lipidomic analysis of wild type and mutant \textit{C. glutamicum} (29) suggested that NCgl2764 is required for synthesis of AcTMCM. The mutant also accumulated trehalose monoketocorynomycolates (ketoTMCM), had supressed synthesis of phosphatidyglycerol (PG) and alanylated diacylglycerols (Ala-DAG), and lacked a class of acyl-PG-like lipids, mirroring the general lipid phenotype of a \textit{tmaT} mutant. Mutation analysis provided further support for NCgl2764 having methyltransferase activity, revealing unexpected complexity in the regulation of corynomycolate transport across the inner membrane of these bacteria.

Results

\textbf{Identification of the corynebacterial ortholog of \textit{M. tuberculosis} Rv0224c} – We have previously identified a genetic locus in \textit{C. glutamicum} that harbors a number of genes involved in regulating cell wall synthesis. This locus is conserved in all Corynebacterineae (15,25,30), and contains NCgl2759 (\textit{tmaT}, Rv0228) (28), NCgl2760 (Rv0227c) (31) and NCgl2764 (Rv0224c) (Fig. 2). NCgl2764 is functionally uncharacterized but is predicted to encode a protein of 257 residues, sharing 49% identity and 86% similarity with \textit{M. tuberculosis} Rv0224c (Fig. S1). NCgl2764/Rv0224c is annotated as a cytoplasmic methyltransferase based on weak similarity with a range of methyltransferases and the presence of conserved residues essential for activity (see below).
Loss of the NCgl2764 gene affects growth of C. glutamicum – To investigate the function of this gene/protein, a two-step recombination strategy was used to create the NCgl2764 mutant (Fig. 3). Sequences flanking the gene were PCR amplified and cloned into the suicide vector pK18mobsacB (32) which carries a kanamycin resistance gene (aph) and Bacillus subtilis sacB gene conferring sensitivity to sucrose. This plasmid was introduced into C. glutamicum 13032 and a NCgl2764 deletion mutant, designated ΔNCgl2764, isolated and gene deletion confirmed by PCR analysis (Fig. 3A, B). The ΔNCgl2764 mutant formed small colonies relative to the wild-type (WT) parent on agar plates and displayed an extended lag-phase in liquid BHI medium, although ultimately achieved a final cell density comparable to the WT parent strain (Fig. 3C). A similar growth phenotype was observed following loss of tmaT (i.e. ΔNCgl2759) but not other genes (i.e. ΔNCgl2760) in this locus (28,31).

Complementation of C. glutamicum ΔNCgl2764 – Complementation and control strains were created to determine whether the growth defect was due solely to disruption of NCgl2764. To complement the mutant, ΔNCgl2764 was transformed with pSM22:NCgl2764, a plasmid carrying a full-length NCgl2764 gene plus 203 bp of upstream sequence. The empty pSM22 vector (32) was also introduced into ΔNCgl2764 as a control. Growth curves indicated that introduction of pSM22:NCgl2764 into the ΔNCgl2764 mutant restored normal lag phase and growth while the empty pSM22 vector did not (Fig. 3C).

Antibiotic susceptibility testing – Corynebacterial mutants with defects in cell wall synthesis and integrity characteristically display increased sensitivity to antibiotics. To test the condition and permeability of the cell wall in the ΔNCgl2764 mutant, the strains were cultured on agar media and exposed to paper discs impregnated with a variety of antibiotics. The sensitivity of the strains was assessed by the size of zones of inhibition (ZOI) around the discs. For low molecular weight antibiotics (<500g/mol) the ΔNCgl2764 mutant displayed significantly larger ZOIs than the WT parent (P<0.05) and complemented strains (Fig. 4). The increases in zone diameter were particularly apparent for the β-lactam class antibiotics amoxicillin, cefoxitin and imipenem, suggesting increased access of the drugs to the peptidoglycan layer. Increased sensitivity to drugs targeting cytoplasmic processes (RNA and protein synthesis) was also observed. Clustering of the NCgl2764 gene with cell wall biosynthesis genes, together with the altered growth rate and increased antibiotic sensitivity in the ΔNCgl2764 strain, suggested a role for the encoded protein in cell wall biosynthesis.

Loss of NCgl2764 affects synthesis of trehalose corynomycolates – HPTLC analysis of lipid extracts of WT and ΔNCgl2764 bacteria revealed major differences in the abundance of several different cell wall lipid classes (Fig. 5A). In particular, the ΔNCgl2764 mutant accumulated a lipid species during logarithmic growth (15,28) that comigrated with hTMCM on HPTLC and was confirmed to be this glycolipid by MALDI-TOF-MS (Fig. S2). The accumulation of hTMCM in ΔNCgl2764 was associated with concomitant decrease in levels of h2TDCM, relative to WT (Fig. 5A). Levels of both species were restored following complementation of the mutant with pSM22:NCgl2764, but not with empty pSM22. These data suggest that NCgl2764 is required for efficient conversion of hTMCM to h2TDCM.
To determine whether the *M. tuberculosis* ortholog, Rv0224c, could functionally complement the mutant, we constructed a pSM22:Rv0224c plasmid and introduced it into the ΔNCgl2764 mutant. hTMCM and h2TDCM levels in the Rv0224c complemented line were unchanged compared to the original mutant (Fig. S3), suggesting that Rv0224c is unable to substitute for NCgl2764. While the two proteins share significant sequence similarity, including two glycine residues essential for methyltransferase function (see below and Fig. S1), the Rv0224c homologue may lack sequence motifs required for protein:protein interactions essential for activity in *C. glutamicum*.

**Labelling studies** – In order to measure rates of glycolipid turnover in WT and mutant lines, bacteria were pulse labelled with $^{14}$C-acetate, then suspended in BHI medium (chase) and sampled at various time points. Cell wall lipids were then separated by HPTLC and labelled species detected by autoradiography (Fig. 5B). Label was rapidly incorporated into hTMCM and phosphatidylglycerol (PG) species during the pulse and subsequently into h2TDCM in the 5 min chase in WT cells. In contrast, label was not incorporated into h2TDCM in the ΔNCgl2764 strain throughout the 60 min chase period, demonstrating a significant delay in its synthesis. Interestingly, in contrast to the orcinol stained samples (Fig. 5A), no accumulation of $^{14}$C-hTMCM was observed in the ΔNCgl2764 mutant, implying that the high steady state levels of hTMCM (Fig. 5A) reflect reduced rate of turnover of hTMCM rather than excess production of this glycolipid. Interestingly, incorporation of $^{14}$C-label into PG lipids was also reduced in the ΔNCgl2764 mutant, similar to the phenotype of the tmaT mutant (29) and consistent with the lipidomics analysis (see below).

**Altered dynamics of hTMCM turnover and h2TDCM synthesis in a ΔNCgl2764 mutant** – To investigate the dynamics of hTMCM conversion to h2TDCM in the mutant, glycolipids were extracted at specific growth stages and analysed by HPTLC. Cells were cultured in BHI and sampled at OD$_{600}$ 0.5 to 10, and after 24 hours (stationary phase). Samples were extracted and normalized by pellet wet weight prior to HPTLC analysis (Fig. 5C). In the WT and complemented ΔNCgl2764 mutant, h2TDCM levels steadily increased from early log phase through to stationary phase, peaking by OD$_{600}$ ~4. hTMCM levels also peaked at OD$_{600}$ ~4, then reduced to be undetectable by stationary phase. In the ΔNCgl2764 mutant, an accumulation of hTMCM was observed by OD$_{600}$ ~2 (early log phase) before reducing at OD$_{600}$ ~7, and these species were still detectable in stationary phase. h2TDCM synthesis was delayed in the mutant, not reaching maximal levels until OD$_{600}$ ~7. These findings suggested that synthesis of h2TDCM from hTMCM is delayed in the ΔNCgl2764 strain, consistent with a defect in the rate of transport of hTMCM to the periplasmic space in logarithmic phase.

Since hTMCM is known to be the donor of corynomycolates for formation of the AG-linked corynomycolate outer membrane, we also extracted and analyzed covalently bound corynomycolates from the delipidated pellets. No significant differences were observed between WT and the mutant at the various time points (Fig. S4), suggesting that the limited pool of transported corynomycolates are channelled to AG at the expense of h2TDCM synthesis. Unaltered levels of AG-linked corynomycolates were also observed in a tmaT mutant (28).

**Global changes in the cell envelope composition in the C. glutamicum NCgl2764 mutant** – To further investigate whether
hTMCM does indeed accumulate on the cytoplasmic leaflet of the IM in the ∆NCgl2764 mutant, IM and OM lipid extracts of the WT and mutant strains were analyzed by LC-ESI-QTOF-MS in positive ionization mode (29). Lipid species identified by MS/MS from WT and the ∆NCgl2764 mutant are shown in Table 1. After removal of isotopologues, dimer species, isomers and isobars, 139 and 111 lipid species were identified with high confidence in the IM and OM of the NCgl2764 mutant, respectively. Compared to WT bacteria, the IM fraction of the NCgl2764 mutant contained elevated levels of hTMCM, ketoTMCM and hGMM species. Conversely, levels of PG and Ala-DAG species were decreased, while AcTMCM and Acyl-PG like species were either absent or present at very low levels. These changes were also observed in the tmaT (NCgl2759) mutant lipidome (Table 1 and (29)).

As NCgl2764 encodes a putative methyltransferase, we looked for evidence that major TMCM species might be methylated. Methylation of either the lipid or glycan moieties of the major TMCM species carrying C32:0, C34:0, C32:1, C34:2, C26:1 and C28:1 corynomycolic acids would increase their size by 14 Da. Methylation of the glycan headgroup would result in production of fragment ions corresponding to loss of 211 Da (methylated hexose with NH_4^+) and 373 Da (two hexoses with one being methylated, with NH_4^+), respectively, instead of 197 and 359 Da. However, neither of these molecular or fragment ions were detected, suggesting that TMCM is not directly methylated or that only a very small proportion of the steady state pool of TMCM carries this modification. Changes in the relative abundance of individual lipid molecular species in WT and mutant lines were verified by quantitation of the relative ion (lipid) intensities within four replicate samples of each line. Lipids identified in the MS1 analysis are summarized in Fig. 6 (A – F). Both MS1 and MS/MS results confirm the global lipidome changes and an overall shared phenotype between the NCgl2764 and tmaT (29) mutants.

Collectively, these findings suggest that NCgl2764 is required, together with TmaT, for the synthesis of AcTMCM. To investigate whether these genes may function at the same step, we constructed a double mutant by deleting the NCgl2764 gene in the tmaT mutant. The resultant strain grew similarly to the individual mutants (data not shown) and analysis of cell wall glycolipids by HPTLC with orcinol staining revealed an identical lipid profile to the individual NCgl2764 and NCgl2759 knock-out lines (Fig. 7, lane 6). Specifically, the double mutant exhibited a significant accumulation of hTMCM at the expense of h2TDCM and reduced synthesis of PG lipids. These data strongly suggest that NCgl2764 and TmaT regulate the same step or pathway in these bacteria.

**Lipoglycan analysis of a NCgl2764 mutant –**

Our previous studies on genes of this locus revealed that one of them, NCgl2760, was involved in lipoglycan biosynthesis (31). To determine whether loss of NCgl2764 also impacts the synthesis of other cell wall components, lipomannan (LM) and lipoarabinomannan (LAM) were extracted from delipidated cell pellets and analysed by PAGE and LC-TOF-MS. PAGE analysis suggested that the overall abundance and degree of heterogeneity of these lipoglycans were similar in both the WT and ∆NCgl2764 mutant (Fig. S5). Similarly, LC-ESI-TOF-MS analysis using a newly developed profiling method also indicated minimal change in the LM/LAM structure in WT and the ∆NCgl2764 mutant bacteria. The latter approach allows
detection of LM and LAM molecular species that have been assembled on different lipid anchors. In WT bacteria, the glycans chains of detected LM species contain between 1-38 mannose residues, and six different lipid anchors, comprising PI with 2-3 acyl chains (53 molecular species total, 998 – 6997 Da after deconvolution; Fig. S6). Similarly, the LM from the ΔNCgl2764 mutant contained between 1-44 mannose residues and six lipid anchors (61 molecular species, 998 and 8002 Da). Minor differences were observed between WT and mutant bacteria in the degree of polymerization of one of the lipid anchor platforms. The plots in Fig. S6 show the masses (m/z, after deconvolution) of all detected LM/LAM species, from low to high molecular weight, and predicted mannan chain length. Supplemental Fig. S5 of our previous study (31) explains, step by step, the development of Fig. S6.

Mutagenesis of NCgl2764 – To investigate whether the putative methyltransferase activity of NCgl2764/Rv0224c is functionally important, the ΔNCgl2764 mutant was complemented with plasmids encoding NCgl2764 proteins in which specific amino acids in the methyltransferase catalytic site had been mutated. The amino acid sequences of NCgl2764 and its mycobacterial orthologs were compared to two confirmed methyltransferases, MT0146/CbiT and protein arginine methyltransferase 1 (PRMT1; (33)), for which crystal structures have been solved and active site residues identified (34,35). Two glycine residues are conserved in nearly all analyzed S-adenosyl methionine (SAM) dependent methyltransferase sequences and are known to be involved in methyl donor binding (35). These correspond to G69 and G71 in NCgl2764, which are also conserved in Rv0224c and ML2584 (Fig. S1). We hypothesized that substitution of either of those crucial glycine residues with alanine would result in loss of a methyltransferase function, if present, and failure to complement the ΔNCgl2764 phenotype.

The desired mutations were engineered into the pSM22:NCgl2764 plasmid previously used to complement the ΔNCgl2764 strain (see above). Transformation of the deletion mutant with the pSM22:NCgl2764_G69A or pSM22:NCgl2764_G71A constructs, but not the unmodified pSM22:NCgl2764 plasmid encoding the native protein, resulted in strains with the delayed growth phenotype (data not shown) and with glycolipid profiles identical to ΔNCgl2764 (Fig. 7, lanes 1-5). This finding is in agreement with the annotation of NCgl2764 as a methyltransferase, although we cannot discount the possibility that NCgl2764 may have other functions.

Discussion

We have previously identified a conserved gene locus in Mycobacteria and Corynebacteria that is required for cell wall assembly (28,29,31). Most of the genes in this locus are essential in M. tuberculosis, hampering their functional characterization in this pathogen (36,37). In contrast, these genes are not essential in C. glutamicum, reflecting the capacity of this bacterium to tolerate major defects in cell wall synthesis. One of the first genes characterized in this locus, C. glutamicum NCgl2759 (ortholog of M. tuberculosis Rv0228, designated tmaT), encodes an acetyltransferase that acetylates the mycolic acid moiety of hTMCM, facilitating transport of this glycolipid across the inner membrane (28), presumptively by the MmpL3 transporter. Consistent with these findings, TmaT was recently found to be the only protein of the mycolic acid pathway to interact with MmpL3 in a two-hybrid screen (38). It remains unclear whether acetylation is required for recruitment
of AcTMCM to the membrane domains containing the transporter and/or for transporter recognition and transmembrane movement. The next gene in this locus, NCgl2760 (Rv0227c) has been shown to be required for synthesis of mature cell wall lipoglycans, LM and LAM (31) and, surprisingly, Rv0227c also interacts with MmpL3 (38). Although LM/LAM biosynthesis lacks any shared enzymes with the hTMCM pathway, both pathways are likely to be co-ordinately regulated to ensure balanced synthesis of major cell wall components.

In this study, we show that NCgl2764 is also involved in cell wall synthesis. The encoded protein, denoted MtrP, shows similarity to protein methyltransferases. Phenotypic characterization of the ΔNCgl2764 deletion mutant indicated that this protein is involved in trehalose corynomycolate metabolism, rather than LM/LAM synthesis. The mutant was found to share striking cell wall similarities with the ΔtmaT mutant previously characterized by our group. Both mutants failed to produce AcTMCM, accumulated hTMCM and ketoTMCM in the IM fraction, and exhibited markedly reduced kinetics of synthesis of h2TDCM (28). This phenotype is consistent with reduced transport of hTMCM across the inner membrane by C. glutamicum transporters CmpL4 (NCgl0228) and CmpL1 (NCgl2769). The latter has significant sequence similarity to mycobacterial TMM transporter MmpL3, the direct or indirect target of several molecules with antimycobacterial activity (16,20,21,39-41). Deletion of both NCgl2764 and tmaT resulted in phenotype that was essentially identical to that of the single gene knock-out lines, supporting the notion that both proteins act sequentially or on the same step within the same pathway.

In addition to the reduced conversion of hTMCM to h2TDCM in the ΔNCgl2764 strain, a reduced synthesis of PG and Ala-DAG species and the complete absence of Acyl-PG-like lipids was observed. Again, this lipid profile was also observed in our ΔtmaT mutant (29) and likely reflects global changes in the synthesis of other cell wall components to compensate for the accumulation of hTMCM/ketoTMCM in the IM and loss of TDCM in the OM. Changes in LM mannan chain length were also detected in the NCgl2764 mutant but were minor relative to the significant changes in other lipid classes and are probably also a secondary effect.

NCgl2764 and its mycobacterial orthologs are annotated as possible methyltransferases based on sequence similarity with a range of SAM-dependent enzymes. Mutagenesis of two glycine residues equivalent to those in PRMT1 known to bind the methyl donor resulted in loss of NCgl2764 activity. While these analyses support the annotation of NCgl2764 as a methyltransferase, further studies are needed to confirm enzyme activity. We have been unable to find any evidence that TMCM lipids are methylated in WT bacteria, suggesting that NCgl2764 may be involved in regulating the activity of TmaT. This conclusion is supported by the finding that loss of NCgl2764 phenocopies loss of NCgl2759 (TmaT) and the double knockout line. NCgl2764 could either directly methylate TmaT and/or regulate its function through protein:protein interactions. Protein methylation has been shown to modulate the activity of histones, leading to epigenetic effects through repression or activation of gene expression (42). Interestingly, histones can also be acetylated (43), and the histone acetyltransferases responsible are themselves often methylated, usually at arginine residues within glycine/arginine-rich “GAR” sequences.
Alternatively, MtrP and TmaT may form heterodimeric or larger protein complexes. Disruption of one protein could destabilize the other, leading to the similar lipid phenotypes of our mutants. Further studies are required to explore this hypothesis and clarify the relationship between these proteins.

The MtrP ortholog in *M. tuberculosis* H37Rv, Rv0224c, is reported to be essential for in vitro growth based on several high-density mutagenesis and sequencing studies (36,37,46). Although the cell wall defect in the NCgl2764 mutant was not reversed by introduction of a plasmid carrying the Rv0224c gene, the sequence similarity between the two proteins would suggest that Rv0224c may perform a similar function in this pathogen, making it a potential target for studies aimed at developing new antimycobacterial agents or at making bacterial strains more sensitive to existing agents, a phenotype seen in our mutant.

**Experimental Procedures**

*Bacterial strains and culturing conditions* - *Escherichia coli* DH5α was grown in Luria-Bertani (LB) medium at 37°C with aeration. *C. glutamicum* ATCC 13032 was grown in Brain Heart Infusion (BHI) medium (Oxoid) or LBHIS (LB, BHI, sorbitol) at 30°C with aeration. When necessary, ampicillin was added to a final concentration of 100 µg ml⁻¹ and kanamycin at 50 µg ml⁻¹.

*Genetic manipulation of bacteria* - *E. coli* plasmid DNA was isolated using the High Pure plasmid isolation kit (Roche) and *C. glutamicum* genomic DNA was extracted using the Illustra DNA extraction kit (GE Healthcare), according to the manufacturer’s instructions. DNA manipulations and molecular biology techniques were performed as described (28).

**Bioinformatic identification and analysis of NCgl2764** – The corynebacterial ortholog of *M. tuberculosis* Rv0224c was found using the BLASTp (47) algorithm. Mycobacterial DNA and protein sequences were obtained from Mycobrowser (https://mycobrowser.epfl.ch). Amino acid sequence alignments were generated using Clustal Omega (http://www.ebi.ac.uk/Tools/msa/clustalo/).

**Construction of *C. glutamicum* ΔNCgl2764 and a ΔNCgl2764/ΔtmaT double mutant** – The NCgl2764 gene was deleted in wild-type *C. glutamicum* using a two-step allelic replacement strategy (15,25,28,30,31). A 0.9 kb fragment containing sequence from the left side of the NCgl2764 gene was amplified using ProofStart DNA polymerase (Qiagen) with primers NCgl2764leftfor (5’-CCATCCCCGGAAGAACGCCGTGGCGCATCC) and NCgl2764leftrev (5’-CGTATCTAGATCGGGTTTGTTCGT and cloned into the SmaI/XbaI sites (underlined) of pUC18 (48), creating plasmid pUC-NCgl2764left. A 0.9 kb fragment containing sequence from the right side of NCgl2764 was amplified using primers NCgl2764rightfor (5’-AATGTCTAGATGGTTGCCGCTTTTCCCCGC) and NCgl2764rightrev (5’-CGACAAAGCTTGTGCTGACCTAATCCCG) and cloned into the XbaI/HindIII sites (underlined) of pUC18, creating plasmid pUC-NCgl2764right. The right flanking sequence was then excised from pUC-NCgl2764right using XbaI/HindIII and subcloned into XbaI/HindIII-digested pUC-NCgl2764left, fusing the left and right flanking sequences to create plasmid pUC-ΔNCgl2764. The 1.8 kb fused insert was then liberated using SmaI/HindIII and subcloned into SmaI/HindIII.
-digested pK18mobsacB, a suicide plasmid for *C. glutamicum* (49) that contains kanamycin and sucrose selection markers. The resultant plasmid, pK18mobsacB:ΔNCgl2764, was sequenced then electroporated into electrocompetent *C. glutamicum* cells, prepared as previously described (50), using an ECM 630 electroporator (BTX). Clones resulting from single homologous recombination events were selected on kanamycin. These were grown overnight without antibiotic selection then serially diluted and plated onto BHI plates supplemented with 10% (w/v) sucrose to select for a second crossover event. Potential gene knockout mutants (sucrose-resistant, kanamycin-sensitive) were screened by PCR. A ΔNCgl2764/ΔtmaT double mutant was produced by applying the same approach in an existing ΔtmaT mutant strain (28).

**Complementation of ΔNCgl2764** - To complement the ΔNCgl2764 strain with the NCgl2764 gene, a 0.9 kb fragment was PCR amplified using primers NCgl2764-comp_F (5’-GGGTCTAGATGCCATCACCACCATTTCG-3’) and NCgl2764-comp_R (5’-CGCTCTAGAGCGCTTTCTTGTCGTTAACG-3’), digested with XbaI (underlined) and cloned into the unique PvuII site of pSM22 (32), which contains the corynebacterial origin of replication repA and kanamycin resistance gene aphA3. To complement the ΔNCgl2764 strain with the Rv0224c gene from *M. tuberculosis*, a 0.9 kb fragment was PCR amplified using primers Rv0224c_comp-L (5’-TCGGATATCCGATGTCGCAGGTCGACCT-3’) and Rv0224c_comp-R (5’-TCGGATATCCGATGTCGACCT-3’), digested with EcoRV (underlined) and cloned into the unique PvuII site of pSM22. Sequenced complementation plasmids pSM22:NCgl2764, pSM22:Rv0224c and an empty pSM22 control plasmid were electroporated into the *C. glutamicum* ΔNCgl2764 deletion strain, followed by selection on kanamycin supplemented BHI plates.

**Site-directed mutagenesis** – Point mutations were introduced into NCgl2764 using the QuickChange Site-directed Mutagenesis Kit (Stratagene) according to the manufacturer’s instructions. The G69A mutation (underlined) was produced using primers A (5’-CTCGAGTCGCCGCAGGGACACCCGATAC TTCGCC; forward primer) and B (5’-CCGGGTTCGCCGCGACGTCGAAGACG TTCAGG; reverse primer). The G71A mutation (underlined) was produced using primers C (5’-CGACGTCGCAGGACACCCGATAC CGCC; forward primer) and D (5’-CGAAGTATCCGTTGCACCGGACGT CGAGAACCTTT; reverse primer).

**Extraction of cell wall components and HPTLC** – *C. glutamicum* strains were grown to logarythmic phase (OD$_{600nm}$ = 5-7) or stationary phase (OD$_{600nm}$ = 10) in BHI medium (Oxoid). Total lipids were extracted in chloroform/methanol (2:1, v/v) and chloroform/methanol/water (1:2:0.8, v/v) (28). After removal of insoluble material by centrifugation (15 000g, 10 min), extracts were dried under nitrogen and subjected to biphasic partitioning in 1-butanol and water (2:1, v/v). The organic phase was dried, and lipids were resuspended in 1-butanol. Lipid fractions were analyzed by high-performance thin-layer chromatography (HPTLC) using aluminum-backed silica gel sheets (Merck). One-dimensional HPTLCs were developed in chloroform/methanol/13 M ammonium solution/1 M ammonium acetate/water (180:140:9:9:23, v/v). Glycolipids were stained and visualized with orcinol/H$_2$SO$_4$.
Extraction of covalently bound corynomycolates – Dried, delipidated cell pellets were base-treated to cleave AG-bound corynomycolates. Pellets were suspended in 2 ml of 0.1 M KOH in ethanol (30°C, 3 hrs) and samples neutralized with 0.5M acetic acid prior to centrifugation at 3000g for 10 min. Cell pellets were dried by evaporation using a Savant Speed Vac Plus SC11A concentrator then resuspended by sonication in 400 µl of water-saturated 1-butanol and mycolates recovered after phase partitioning in 1-butanol-water (2:1 v/v) and collection of the upper 1-butanol layer.

Pulse-chase radiolabeling – Bacterial strains were grown in BHI to mid-log phase (OD_{600}~5) at which cells were harvested by 10 min centrifugation at 13,000 xg, washed with HEPES-saline (pH 7.4), re-pelleted and weighed. Pellets were resuspended in HEPES-saline to 0.25 g per ml and incubated for 5 min. Aliquots (200 µl) were transferred to new tubes and 2 µCi of ^14C acetate added to each sample (except for the “cold” control) followed by incubation for 5 min. The “pulse” sample was then collected and harvested by brief centrifugation and resuspension of the pellet in 500 µl of chloroform: methanol (2:1 v/v). The “chase” samples were pelleted and the medium replaced with pre-warmed BHI, followed by incubation at 30°C for the designated chase time. Chase samples were collected after 5, 15, 30 and 60 min and harvested by brief centrifugation and resuspension of the pellet in 500 µl of chloroform: methanol (2:1 v/v). Following a 1 h extraction, the supernatants were collected by 5 min centrifugation at 4,000 rpm and the pellets further extracted for 1 h with 500 µl of chloroform:methanol:water (1:2:0.8 v/v). The glycolipid-containing supernatants were collected by 5 min centrifugation at 4,000 rpm and dried under a N₂ stream. The samples were then suspended in 200 µl of water-saturated butanol-1 and 100 µl of butanol-saturated water and the butanol phase collected and dried after vigorous vortexing. The sample was then suspended in 20 µl of butanol-1 and 2 µl was applied to a HPTLC plate and developed as described above.

Extraction and analysis of lipoglycans – LM and LAM were extracted and purified as previously described (31), separated by polyacrylamide gel electrophoresis (PAGE) and stained using a SilverSnap kit (Pierce). Alternatively, purified LM/LAM was suspended in 10 mM ammonium carbonate/bicarbonate buffer (pH 8.16) and 4 µl was injected by direct infusion into an Agilent 6220 Accurate-Mass Time-of-Flight LC/MS system (Agilent Technologies). The run was performed in negative ionization mode, with a mass range of 100–3000 Da. The reference nebulizer was set to 20 psig with a detection window of 100 ppm, a minimum height of 1000 counts, an acquisition rate of 0.63 spectra/s, and an acquisition time of 1589.4 ms/spectrum. The gas temperature was set to 325 °C with a drying gas of 7 L/min, dual ESI 3500 V, Fragmentor 150 V, and Skimmer 65 V. The flow rate was set to 0.25 ml/min, and a solvent consisting of 0.1 M formic acid, acetonitrile (1:1, v/v) was used to wash the lines for 25 min. Data were analyzed using MassHunter and the length of the mannan chain was calculated, assuming the presence of a PI lipid anchor containing C16:1/18:1/16:0 fatty acids. Analyses were performed in triplicate.

Extraction of inner and outer membrane lipids for LC/MS and lipidomics analyses – Inner and outer membrane lipids of C. glutamicum were extracted and analyzed as previously described (29). In brief, four replicates of WT C. glutamicum and mutant strain were grown to exponential phase
(OD$_{600nm} = 2.5 – 3$) in BHI medium. Cells were harvested by centrifugation. The cell pellet was extracted with water-saturated 1-butanol to selectively remove outer membrane lipids. After another centrifugation step, the same pellet was sequentially extracted in chloroform/methanol (2:1, v/v) and chloroform/methanol/water (1:2:0.8, v/v) to extract the remaining lipids of the inner membrane. Both inner and outer membrane lipids were separated on an Agilent 1290 Infinity Quaternary LC System (Agilent Technologies) using a C18 column (Phenomenex Kinetex, 2.6 µm EVO C18 100A) eluted with an isopropanol mobile phase binary solvent system at a flow rate of 260 µl/min. Eluted lipids were analyzed on a 6550 iFunnel Q-TOF LC/MS instrument (Agilent Technologies) in positive ionization mode. Lipids were identified based on their mass-to-charge ratio and fragmentation pattern using a lipid library (29). Statistical analyses were performed using MetaboAnalyst.

**Data availability**

Data described in the manuscript are available from the corresponding author, upon request (paul.crellin@monash.edu).

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**Conflict of interest**

The authors declare that they have no conflicts of interest with the contents of this article.

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**Footnotes**

The abbreviations used in the text are: hTMCM, trehalose monohydroxycorynomycolate; h2TDCM, trehalose dihydroxycorynomycolate; AcTMCM, acetyltrehalose monocorynomycolate; ketoTMCM, monoketocorynomycolate; TMM, trehalose monomycolate; TDM, trehalose dimycolate; GMM, glucose monomycolate; GroMM, glycerol monomycolate; TmaT, TMCM mycolyl acetyltransferase; AG, arabinoxylol; LAM, lipoarabinomannan; LM, lipomannan; HPTLC, high-performance thin-layer chromatography; WT, wild-type; PAGE, polyacrylamide gel electrophoresis; ZOI, zone of inhibition; PG, phosphatidylglycerol; Ala-PG, alanlated-phosphatidylglycerol; CL, cardiolipin; SAM, S-adenosyl methionine; Gl-X, mannosyl-glucuronic acid diacylgllycerol; AcPIM2, Man2-acyl-PI.
Table 1. Detected lipid (sub)classes and lipid species of the IM and OM fraction of the *C. glutamicum* WT, *tmaT*(NCgl2759) and NCgl2764 mutants via LC-MS/MS ESI TOF in positive ionization mode (CE30).

| Lipid (Sub)Class | Wild-type | ΔNCgl2759 | ΔNCgl2764 |
|------------------|-----------|-----------|-----------|
|                  | IM        | OM        | IM        | OM        |
| 1                | hTMCM     | 18        | 20        | 26        | 15        | 23        | 18        |
| 2                | AcTMCM    | 3         | 8         | 0         | 0         | 0         | 0         |
| 3                | keto-TMCM  | 3         | 0         | 10        | 3         | 8         | 5         |
| 4                | Acyl-hTMCM| 7         | 5         | 12        | 5         | 9         | 7         |
| 5                | Acyl-AcTMCM| 1        | 0         | 0         | 0         | 0         | 0         |
| 6                | h2TDCM    | 28        | 28        | 30        | 25        | 27        | 29        |
| 7                | Ac1-hTDCM | 3         | 3         | 3         | 1         | 0         | 0         |
| 8                | hGMM      | 5         | 11        | 8         | 7         | 8         | 8         |
| 9                | DAG       | 7         | 4         | 8         | 4         | 7         | 5         |
| 10               | Ala-DAG   | 11        | 5         | 9         | 2         | 7         | 1         |
| 11               | CDP-DAG   | 1         | 0         | 1         | 0         | 1         | 1         |
| 12               | TAG       | 10        | 10        | 11        | 11        | 11        | 10        |
| 13               | PG        | 11        | 5         | 5         | 2         | 6         | 5         |
| 14               | Acyl-PG   | 3         | 0         | 1         | 0         | 1         | 0         |
| 15               | Ala-PG    | 2         | 1         | 1         | 0         | 1         | 1         |
| 16               | PA        | 2         | 0         | 0         | 0         | 1         | 1         |
| 17               | CL        | 16        | 3         | 14        | 4         | 10        | 5         |
| 18               | PI        | 4         | 3         | 3         | 2         | 4         | 5         |
| 19               | PIM1      | 1         | 1         | 1         | 0         | 1         | 1         |
| 20               | PIM2      | 1         | 1         | 1         | 1         | 1         | 1         |
| 21               | AcPIM2    | 4         | 1         | 5         | 1         | 4         | 1         |
| 22               | AcPIM3    | 1         | 0         | 1         | 0         | 1         | 1         |
| 23               | AcPIM4    | 1         | 0         | 1         | 0         | 1         | 0         |
| 24               | GI-A      | 6         | 2         | 5         | 1         | 3         | 2         |
| 25               | GI-X      | 6         | 1         | 4         | 1         | 3         | 2         |
| 26               | GI-Y      | 1         | 0         | 1         | 1         | 0         | 1         |
| 27               | GI-Z      | 1         | 0         | 1         | 0         | 1         | 1         |
| 28               | Acyl-PG-like | 4    | 3         | 0         | 0         | 0         | 0         |
| **SUM:**         | **161**   | **115**   | **162**   | **86**    | **139**   | **111**   |
Figure 1. Pathway of trehalose corynomycolate synthesis and transport in *C. glutamicum*. Products of the FAS-I and FAS-II pathways are condensed then attached to trehalose, producing keto-TMCM in the inner leaflet of the IM. Reduction by CmrA produces hTMCM which may be acetylated to AcTMCM by the integral membrane acetyltransferase TmaT, facilitating transport by CmpL1/4. It is hypothesized that an unidentified deacetylase removes the acetyl group after transport, reconstituting hTMCM in the outer leaflet of the IM. Corynomycolyltransferases then produce h2TDCM from two hTMCM molecules or transfer the corynomycolates to acceptors on AG. Other acylation variants of hTMCM, AcTMCM and hTDCM have been identified (29) but are not shown here for clarity. Results of the current study place NCgl2764 (MtrP) at the same step as TmaT in the pathway.
Figure 2. NCgl2764 is encoded within a genetic locus highly conserved in Corynebacterineae. The \textit{tmaT} locus of Corynebacterineae. Likely ortholog genes in the four species are shown using the same colour. The focus of the current study NCgl2764 (boxed) and its likely mycobacterial orthologs are in bold. \textit{MSMEG\_0310} is reported to be a pseudogene. Previously studied genes are NCgl2759 (\textit{tmaT}; (28)) and NCgl2760 (31) while the remaining genes remain uncharacterized.
Figure 3. Disruption strategy and growth analysis for the ΔNCgl2764 mutant. (A) Diagram showing the arrangement of genes in the NCgl2764 region of WT C. glutamicum (above) and ΔNCgl2764 mutant (below). Small horizontal arrows indicate the binding sites for the four primers used to construct the ΔNCgl2761 mutant (a, NCgl2764leftfor; b, NCgl2764leftrev; c, NCgl2764leftrightfor; d, NCgl2764rightrev) and screen for the gene knockout (e, NCgl2764-comp_F; f, NCgl2764-comp_R). Expected PCR product sizes using the e/f primer combination are indicated in kb (kilobase pairs) (B) PCR amplification of genomic DNA of C. glutamicum WT (lane 2) and the ΔNCgl2764 mutant (lane 3) using primers e and f. Positions of Sigma-Aldrich DNA markers III (lane 1) are indicated in kb. (C) Growth curves of WT C. glutamicum, the ΔNCgl2764 mutant and complementation strains in liquid BHI medium. Each strain was grown to saturation, then diluted 1:100 in fresh BHI medium. Triplicate cultures were sampled at the times indicated to determine the optical density (OD) at a wavelength of 600nm. The curves for the WT and ΔNCgl2764 + pSM22:NCgl2764 (complementation) strains overlap as do the ΔNCgl2764 and ΔNCgl2764 + empty pSM22 strains, indicating that adding back the NCgl2764 gene reverses the mild growth defect in the mutant.
Figure 4. Antibiotic susceptibility of a *C. glutamicum* ΔNCgl2764 mutant. Antibiotic sensitivity was measured by determining the ZOI around antibiotic-soaked discs placed on BHI plates spread with the indicated strains.
**Figure 5.** Altered lipid composition and dynamics of turnover in a *C. glutamicum ΔNCgl2764* mutant. 
A) A ΔNCgl2764 mutant accumulates hTMCM during active growth. HPTLC analyses of cell wall glycolipids extracted and purified from cells growing in logarithmic phase (OD$_{600}$=3). Lipids were separated by HPTLC then visualized by orcinol-sulfuric acid staining Lane 1, WT; lane 2, ΔNCgl2764 mutant; lane 3, ΔNCgl2764 containing pSM22:NCgl2764; lane 4, ΔNCgl2764 containing empty pSM22. B) Pulse/chase labelling reveals delayed synthesis of h2TDCM in a ΔNCgl2764 mutant. Autoradiogram of a HPTLC analysis of $^{14}$C acetate labelled lipids extracted during a pulse/chase experiment. The experiment reveals the dynamics of *de novo* synthesis of hTMCM/h2TDCM. In WT *C. glutamicum*, the label appears in h2TDCM within 5 minutes post-labelling. In contrast, no labelled h2TDCM was detected in the ΔNCgl2764 samples even after 60 minutes, consistent with a significantly slower rate of h2TDCM synthesis in this strain. C) Delayed conversion of hTMCM to h2TDCM in a ΔNCgl2764 mutant. Glycolipids were extracted from cells harvested at OD$_{600}$ of 0.5, 2, 4, 7, 10 and from a stationary phase culture (~24 hours growth). HPTLC analysis of glycolipids showed that the accumulation of hTMCM peaks during mid-log growth phase in the analyzed strains. In samples from the ΔNCgl2764 strain, hTMCMs accumulate to a higher level and do so earlier than in the WT or the complemented mutant. The dynamics of h2TDCM synthesis are inverted relative to hTMCM and reach their highest point in the stationary phase where no hTMCM is detected. The identities of glycolipids were based on previously published reports. Gl-X, mannosyl-glucuronic acid diacylglycerol; AcPIM2, Man$_2$-acyl-PI.
Figure 6. Comparison of the relative abundances of IM and OM lipids in C. glutamicum WT and the NCgl2764 mutant. The relative abundance of 142 species, identified by LC-MS/MS profiling, in the IM and OM fractions. Lipid abundances (based on ion intensities) represent the mean value of four replicates. The relative abundance of DAG and TAG based lipid classes (A), PG and CL-based lipid classes (B), PI/PIM and Gl glycolipid classes (C), hTMCM species (D), other trehalose and glucose corynomycolates (E) and h2TDCM species (F) are shown. Compared with WT bacteria (left), the IM fraction of the NCgl2764 mutant (right) lacked detectable AcTMCM and Acyl-PG-like species, and had decreased amounts of PG, Acyl-PG, Ala-DAG, and h2TDCM, while the OM fraction was deficient in Acyl-PG, decreased in PG, hTMCM, AcTMCM and h2TDCM, and increased in hGMM. The lipid phenotype of the NCgl2764 mutant shares the same major defects with a tmaT mutant.
Figure 7. Glycolipid analyses of NCgl2764 mutants and a ΔNCgl2764/ΔtmaT double mutant. HPTLC analysis and orcinol/H$_2$SO$_4$ staining of glycolipids extracted from WT C. glutamicum (lane 1), ΔNCgl2764 (lane 2), ΔNCgl2764 + pSM22:NCgl2764 (lane 3), ΔNCgl2764 + pSM22:NCgl2764 carrying G69A substitution (lane 4), ΔNCgl2764 + pSM22:NCgl2764 carrying G71A substitution (lane 5) and a ΔNCgl2764/ΔtmaT double mutant (lane 6).
MtrP, a putative methyltransferase in Corynebacteria, is required for optimal membrane transport of trehalose mycolates
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