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A vitamin B₁₂ transporter in *Mycobacterium tuberculosis*

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1. Summary

Vitamin B₁₂-dependent enzymes function in core biochemical pathways in Mycobacterium tuberculosis, an obligate pathogen whose metabolism in vivo is poorly understood. Although M. tuberculosis can access vitamin B₁₂ in vitro, it is uncertain whether the organism is able to scavenge B₁₂ during host infection. This question is crucial to predictions of metabolic function, but its resolution is complicated by the absence in the M. tuberculosis genome of a direct homologue of BtuFCD, the only bacterial B₁₂ transport system described to date. We applied genome-wide transposon mutagenesis to identify M. tuberculosis mutants defective in their ability to use exogenous B₁₂. A small proportion of these mapped to Rv1314c, identifying the putative PduO-type ATP: co(I)rrinoid adenosyltransferase as essential for B_{12} assimilation. Most notably, however, insertions in Rv1819c dominated the mutant pool, revealing an unexpected function in B₁₂ acquisition for an ATP-binding cassette (ABC)-type protein previously investigated as the mycobacterial BacA homologue. Moreover, targeted deletion of Rv1819c eliminated the ability of M. tuberculosis to transport B₁₂ and related corrinoids in vitro. Our results establish an alternative to the canonical BtuCD-type system for B_{12} uptake in M. tuberculosis, and elucidate a role in B_{12} metabolism for an ABC protein implicated in chronic mycobacterial infection.

2. Introduction

The genome of *Mycobacterium tuberculosis*, obligate human pathogen and causative agent of tuberculosis, encodes three B_{12} -dependent enzymes. Previous work in our laboratory has established that both the methylmalonyl-coenzyme A (CoA) mutase, MutAB [1], and the *metH*-encoded methionine synthase [2] are functional, and require B_{12} for activity. *Mycobacterium tuberculosis* also possesses a predicted pathway for B_{12} biosynthesis [3], but appears not to produce the cofactor *in vitro* [1,2] or in macrophages [4]. Nevertheless, the bacillus can use exogenous vitamin B_{12} and encodes a B_{12} -responsive riboswitch that suppresses transcription of the alternative, B_{12} -independent methionine synthase, *metE*, in B_{12} -replete conditions [2]. These observations imply a role for the cofactor

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Figure 1. Structure of vitamin B_{12} and B_{12} -derived cofactors.

in M. tuberculosis pathogenesis. However, it is uncertain whether B_{12} is available during infection, and which mycobacterial genes are required for its uptake and assimilation.

Vitamin B₁₂ and B₁₂ derivatives are members of the cobalamin group of corrinoid macrocycles [5]. Cobalamins are structurally complex, comprising a defining tetrapyrrole framework with a centrally chelated cobalt ion held in place by a lower axial base, dimethylbenzimidazole and an upper ligand that determines the cofactor form (figure 1). The cyano group in vitamin B₁₂ (cyanocobalamin, CNCbl) must be replaced by deoxyadenosine and methyl ligands, respectively, during conversion to the biologically active cofactors: adenosylcobalamin (AdoCbl or coenzyme B₁₂), which is required by methylmalonyl-CoA mutase, and methylcobalamin (MeCbl), which serves as an intermediary in the synthesis of methionine from homocysteine and methyltetrahydrofolate [6]. The reactivity of B₁₂ cofactors derives from the cobalt-coordinated organic ligands [7] and, together with the size of the cobalamin core, underlies the need for multi-component systems to mediate controlled translocation and delivery of B_{12} across the cell membrane to its target enzyme [8].

Although bioinformatic analyses have predicted alternative vitamin transporters [9], BtuCD-BtuF remains the only confirmed bacterial B₁₂ transport system identified to date [10]. The Escherichia coli model is the best characterized: a high-affinity corrinoid transporter, BtuB, operates with the TonB-ExbBD complex to traffic B₁₂ across the outer membrane into the periplasm [11] where it is captured by the btuF-encoded B₁₂-binding protein and delivered to the ATPbinding cassette (ABC) importer, BtuCD, which spans the cytoplasmic membrane [12]. Mycobacterium tuberculosis is characterized by a notoriously complex cell envelope comprising a cytoplasmic membrane and an external cell wall [13]. However, despite its demonstrated ability to use exogenous B_{12} [2,4], the proteins involved in mycobacterial B_{12} transport and assimilation are unknown: M. tuberculosis is included in the small number of B₁₂-using bacteria that lack a candidate BtuFCD-type B_{12} transport system [3,9,14] as well as an identifiable homologue of TonB [15].

In this study, we used random mutagenesis to identify genes whose disruption abrogated the ability of M. tuberculosis to use exogenous vitamin B_{12} in vitro. Our results establish an essential role in B_{12} uptake for Rv1819c, a predicted ABC protein implicated in chronic infection $in\ vivo\ [16]$, thereby revealing an alternative to the well-characterized BtuCD system for B_{12} transport.

3. Material and methods

3.1. Bacterial strains and growth conditions

Strains, plasmids and oligonucleotides are described in the electronic supplementary material, table S1. *Mycobacterium tuberculosis* was grown on Middlebrook 7H10 (Difco) supplemented with 0.5 per cent glycerol and Middlebrook OADC enrichment (Difco) or in Middlebrook OADC and 0.05 per cent Tween 80 or 0.05 per cent tyloxapol, as required. For propionate utilization experiments, 7H9 broth was supplemented with 0.5 per cent bovine serum albumin fraction V (Sigma), 0.085 per cent NaCl and 0.1 per cent (w/v) sodium propionate, as described [1]. Hygromycin (hyg), kanamycin (kan) and gentamicin (gent) were used at 50, 25 and 2.5 μ g ml⁻¹, respectively, CNCbl and AdoCbl at 10 μ g ml⁻¹, (CN)₂Cbi at 1 μ M and 3-nitropropionate (3NP) at 0.1 mM.

3.2. Construction of transposon mutant library

A library of transposon (Tn) mutants was constructed in M. tuberculosis H37Rv $\Delta metH$, using the MycoMarT7 phage as described [17]. For the primary screen, transductants were plated across multiple 7H10 plates containing $20~\mu g \, ml^{-1}$ kan and $10~\mu g \, ml^{-1}$ CNCbl at a density of 20~000 colony forming units (CFU) per plate. The secondary screen was performed in duplicate in microtitre plate format and, for each Tn mutant, comprised four parallel wells containing 0.1 per cent propionate plus $20~\mu g \, ml^{-1}$ kan as base medium in each well: the first well constituted a growth control and contained only the base medium; in well 2, $10~\mu g \, ml^{-1}$ CNCbl was added to the base medium; in well 3, the base medium was supplemented with 0.1~mM 3NP; and in well 4, 0.1~mM 3NP and $10~\mu g \, ml^{-1}$ CNCbl were added.

3.3. Identification of transposon insertion sites

A combination of Tn-linker [18] and rescue cloning [19] strategies was applied to identify Tn insertion sites using the oligonucleotides in the electronic supplementary material, table S1.

3.4. Construction of mutant strains of *Mycobacterium tuberculosis*

Mycobacterium tuberculosis mutants were constructed using suicide plasmids described in electronic supplementary material, table S1. Genetic complementation used tweety-based vectors [20].

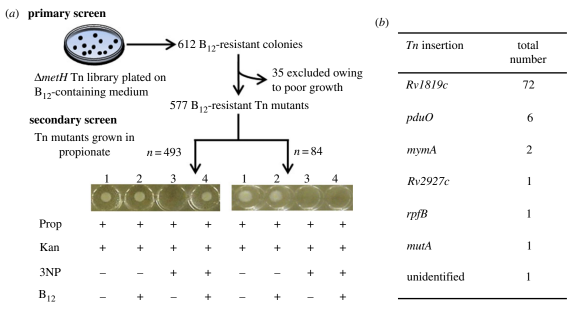


Figure 2. Identification of genes required for B_{12} transport and assimilation. (*a*) Schematic of the screening cascade. The Δ *metH* Tn library was plated on selective medium containing 10 μ g ml⁻¹ CNCbl. 612 ' B_{12} -resistant' clones were isolated and regrown in standard liquid medium, eliminating 35 mutants owing to poor (n=14) or absent (n=21) growth. A secondary screen tested the B_{12} uptake ability of the remaining 577 insertion mutants in a four-well microtitre assay using 0.1% propionate (Prop) plus 20 μ g ml⁻¹ kanamycin as base medium (well 1) supplemented with 10 μ g ml⁻¹ CNCbl (well 2), 3NP (well 3) and 3NP plus 10 μ g ml⁻¹ CNCbl (well 4). A total of 84 mutants failed to grow in well 4, suggesting impaired B_{12} uptake. Each determination was performed in duplicate, and the results confirmed in batch culture. (*b*) Insertion mutants with disrupted B_{12} uptake ability.

3.5. DNA sequencing

Mycobacterium tuberculosis genomic DNA was sequenced using an Illumina GenomeAnalyzer II, as described previously [21].

3.6. Homology modelling

The initial detection of crystal structures related to Rv1819c was performed using HHsearch [22] and COMA [23]. The Rv1819c model was then generated using a previously described iterative approach [24,25]. Briefly, both the set of structural templates and corresponding alignments were refined until the resulting model stopped improving and the visual inspection revealed no significant flaws.

4. Results

4.1. A forward genetic screen identifies B_{12} uptake mutants

We showed previously that deletion of the B₁₂-dependent methionine synthase, MetH, renders *M. tuberculosis* sensitive to vitamin B₁₂ during growth on solid medium [2]. This phenotype depends on the function of a B₁₂ riboswitch that is located immediately upstream of *metE*, the gene encoding an alternative, B₁₂-independent methionine synthase in *M. tuberculosis*. In wild-type *M. tuberculosis*, exogenous B₁₂ suppresses transcription of *metE* by binding to the riboswitch [2], possibly ensuring efficient B₁₂-dependent methionine synthesis by MetH. In the *metH* deletion mutant, however, riboswitch-mediated suppression of *metE* in response to B₁₂ effectively results in the complete shutdown of methionine synthase activity, thereby eliminating production of an

essential amino acid and so inhibiting bacillary growth [2]. This effect is most profoundly manifest on solid medium, where exposure to $10 \,\mu \mathrm{g \, ml}^{-1}$ CNCbl results in a $3 \log_{10}$ fold reduction in viable CFU of $\Delta metH$ knockout mutants [2]. Here, we exploited the observed B_{12} sensitivity of metHmutants in a genetic screen designed to elucidate a potential B₁₂ transport system in M. tuberculosis (figure 2). To this end, we constructed an unmarked metH knockout of the laboratory strain, M. tuberculosis H37RvJO [21] (electronic supplementary material, figure S1a) and confirmed that it phenocopied the previously described hygromycin (hyg)marked $\Delta metH$ (BB) deletion mutant [2] during growth on B₁₂-containing solid medium (see the electronic supplementary material, figure S1b). The unmarked $\Delta metH$ knockout was used as background strain in which to generate a Tn mutant library using the MycoMarT7 phage [17] that carries a kan resistance marker and inserts randomly at TA dinucleotides [19]. In the primary screen, the library of insertion mutants was plated on solid medium containing kan and CNCbl to enable the identification of genes whose disruption alleviated the growth defect of the metH mutant (figure 2a). In total, 612 individual clones were isolated, each of which was picked and regrown in standard liquid medium; of these, 35 grew poorly or not at all and were eliminated, leaving 577 $^{\prime}B_{12}$ -resistant $^{\prime}$ insertion mutants for further analysis.

Previously, in characterizing the $\Delta metH$ (BB) mutant, we noted the high frequency at which suppressor mutants arose spontaneously on B₁₂-containing solid medium, with single-nucleotide polymorphisms (SNPs) in the B₁₂ riboswitch located upstream of metE accounting for approximately 10–20 per cent of these [2]. In the current screen, we used dual selection on kan and CNCbl in order to limit the potentially confounding effects of spontaneous riboswitch mutations: according to these criteria, growth on CNCbl plus kan would require successful transduction with the kan-resistant Tn as well as disruption—

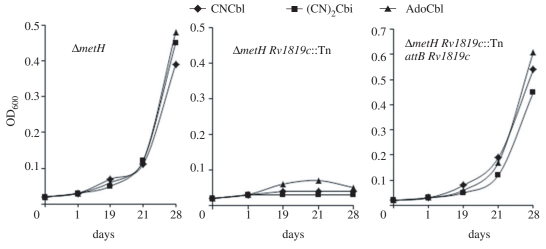


Figure 3. Disruption of Rv1819c eliminates the ability of M. tuberculosis $\Delta metH$ to use corrinoids for growth in 0.1% propionate-containing 3NP. Data are from a representative experiment performed in duplicate.

spontaneous or Tn-mediated—of B₁₂-dependent growth inhibition. Nevertheless, we predicted that a significant proportion of B₁₂-resistant mutants might contain Tn insertions in the riboswitch motif. So, in order to minimize the impact of disruptions to the B₁₂ riboswitch, we applied a secondary screen (figure 2a) to determine the capacity of the insertion mutants to assimilate exogenous CNCbl for growth in liquid medium containing propionate in the presence of 3NP, an inhibitor of the key methylcitrate cycle enzyme, isocitrate lyase [26]. Two prior observations informed the design of this screen: (i) the inhibitory effect of genetic ($\Delta prpDC$) or chemical (3NP) abrogation of methylcitrate cycle enzymes during growth in liquid medium containing propionate can be alleviated by supplementing the culture with CNCbl, thereby enabling M. tuberculosis to use propionate as a carbon source via the methylmalonyl pathway that includes the B₁₂-dependent methylmalonyl-CoA mutase, MutAB [1]; (ii) for reasons that are not clear, B₁₂-mediated growth inhibition is less effective in liquid versus solid medium—that is, the $\Delta metH$ mutant can grow in B₁₂-supplemented liquid medium (see the electronic supplementary material, figure S1c).

The secondary screen therefore assessed the ability of all 577 Tn mutants to use exogenous CNCbl for growth in liquid medium containing propionate in the presence of 3NP (figure 2a). The majority of Tn mutants (n=493) phenocopied the parental $\Delta metH$ strain in this assay, and were eliminated as candidate B_{12} uptake mutants. In contrast, the remaining 84 Tn mutants were unable to grow in well 4, suggesting impaired ability to use exogenous B_{12} for methylmalonyl pathway-dependent propionate catabolism. To verify these results, 43 of the 84 mutants were selected at random for phenotypic confirmation of disrupted B_{12} uptake in batch culture (data not shown) and on B_{12} -containing solid medium (see the electronic supplementary material, figure S2a).

4.2. Disruption of *Rv1819c* eliminates the ability of *Mycobacterium tuberculosis* to use exogenous B₁₂ *in vitro*

Insertions in *Rv1819c* accounted for 72 of the 84 Tn mutants (figure 2*b*; electronic supplementary material,

figure S2a-c) and, moreover, mapped throughout the 1920 bp gene (electronic supplementary material, figure S2d). This result strongly suggested a role in B₁₂ uptake for a predicted ABC transport protein previously identified as the putative M. tuberculosis homologue of BacA [16,27], a protein of cryptic function implicated in chronic infection in multiple host-pathogen models [25]. In their study of M. tuberculosis Rv1819c, Domenech et al. [16] constructed a deletion mutant by allelic exchange mutagenesis (see the electronic supplementary material, figure S2b). We assessed the ability of this mutant—referred to as $\Delta bacA::hyg$ by Domenech et al. [16]—to use exogenous B₁₂ for MutABdependent growth in propionate (electronic supplementary material, figure S3a). Consistent with the inferred role of Rv1819c in B₁₂ uptake, the ΔbacA::hyg strain grew very poorly in propionate plus 3NP supplemented with B₁₂, reproducing the phenotype of the *AmetH Rv1819c*::Tn mutants (figure 3). By contrast, the complemented derivative carrying a full-length copy of Rv1819c at the attB site, referred to as ΔbacA::pKLMt5 in the original study [16], was able to use B_{12} for growth (see the electronic supplementary material, figure S3a). Similarly, integration of full-length Rv1819c at attB restored the B₁₂-sensitive phenotype of a randomly selected $\Delta metH$ Rv1819c::Tn mutant during growth on solid medium supplemented with CNCbl (see the electronic supplementary material, figure S3b), and reversed the inability of the same mutant to use B₁₂ for growth in propionate-containing liquid medium supplemented with 3NP (figure 3), confirming the essentiality of Rv1819c in this assay.

It was noticeable in the propionate utilization experiment (see the electronic supplementary material, figure S3a) that the $\Delta bacA::hyg$ mutant started to replicate after two to three weeks of apparent growth arrest, possibly indicating the emergence of suppressor mutants. To circumvent this complication, we deleted the prpDC locus [28] in this strain, thereby negating the need to use 3NP to eliminate methylcitrate pathway function [1]. In contrast to the single prpDC deletion mutant, the double $\Delta bacA::hyg$ $\Delta prpDC$ knockout exhibited no growth at all in propionate over the 28-day time course (see the electronic supplementary material, figure S3c), even when supplemented with CNCbl, strongly suggesting that Rv1819c is required for the assimilation of exogenous B₁₂ to enable methylmalonyl-CoA pathway function.

4.3. Spontaneous B_{12} -resistant mutants carrying non-synonymous single-nucleotide polymorphisms in Rv1819c

We reported previously that SNPs in the metE-associated B₁₂ riboswitch accounted for 10-20 per cent of all B₁₂-resistant mutants isolated after plating the *AmetH* (BB) knockout on medium containing CNCbl, whereas the remaining B₁₂-resistance mutations were unknown [2]. To investigate the possibility that mutations in Rv1819c might account for B₁₂ resistance in those clones lacking riboswitch mutations, we plated the $\Delta metH$ (BB) strain on medium containing CNCbl and sequenced the riboswitch region and Rv1819c locus in 10 spontaneous B₁₂-resistant mutants. Consistent with previous results [2], two isolates carried independent mutations in the highly conserved B12-box motif within the metE riboswitch [29], namely $C \rightarrow T$ transversions at positions -155 and −163 relative to the *metE* start codon, respectively. Notably, four other B₁₂-resistant mutants had wild-type riboswitch sequences, but contained non-synonymous SNPs in Rv1819c (see the electronic supplementary material, table S2), supporting the inferred role of Rv1819c in B₁₂ uptake. To eliminate the possibility that an additional, unidentified mutation (or mutations) might account for the observed phenotype, we sequenced the genome of a representative Rv1819c point mutant, SP09 (see the electronic supplementary material, table S2). The parental, B_{12} -sensitive strain, $\Delta metH$ (BB), was differentiated from the laboratory strain, H37RvJO [21], only in the targeted deletion of metH sequence. Moreover, the Rv1819c mutation constituted the sole polymorphism separating SP09 from its $\Delta metH$ (BB) parent and, importantly, complementation with wild-type Rv1819c at the attB locus restored B₁₂ sensitivity to both SP09 and SP18 (see the electronic supplementary material, figure S4).

In the primary Tn screen (figure 2a), 'B₁₂-resistant' mutants had been selected on kan and CNCbl in order to limit the potentially confounding effects of spontaneous riboswitch mutations. To verify the utility of this approach, we analysed the insertion sites in a random selection of 20 of the $493 \ \Delta metH$ Tn mutants subsequently eliminated in the secondary screen owing to their inability to use exogenous B₁₂ for growth in propionate. All 20 mutants contained insertions in the B₁₂ riboswitch region directly upstream of metE (data not shown), confirming that disrupted riboswitch function represents a major mechanism for loss of B₁₂ regulation in strains which carry an intact Rv1819c gene.

4.4. Rv1819c is essential for corrinoid transport in *Mycobacterium tuberculosis*

Mycobacterium tuberculosis is predicted to encode a complete pathway for B_{12} biosynthesis, including enzymes required for the conversion of the B_{12} precursor, cobinamide, to AdoCbl through the addition of dimethylbenzimidazole and deoxyadenosine ligands [3]. The *E. coli* corrinoid transporter, BtuFCD, mediates uptake of cobinamide as well as CNCbl and AdoCbl [30], suggesting that Rv1819c might fulfil a corresponding role in *M. tuberculosis*. In support of this idea, cobinamide—provided as the dicyanide salt, (CN)₂Cbi—was unable to complement the growth defect of Δ metH Rv1819c::Tn mutants in propionate in the presence

of 3NP, mimicking similar observations with AdoCbl and CNCbl (figure 3). Insertions in *Rv1819c* also alleviated the growth inhibitory effect of AdoCbl, CNCbl and (CN)₂Cbi on the *metH* knockout mutant on solid medium, a phenotype that was reversed upon complementation with wild-type *Rv1819c* (see the electronic supplementary material, figure S5). In combination, these results confirmed the essentiality of Rv1819c for corrinoid transport in *M. tuberculosis*.

4.5. Impaired vitamin B_{12} uptake in spontaneous bleomycin-resistant *Rv1819c* mutants

Domenech et al. [16] showed that deletion of Rv1819c decreased the susceptibility of M. tuberculosis to the glycopeptide antibiotic, bleomycin, a phenotype commonly associated with BacA function [31-33]. We determined the minimum inhibitory concentration (MIC) of bleomycin against a selected Rv1819c::Tn mutant (electronic supplementary material, figure S6a) as well as the spontaneous B₁₂-resistant mutants, SP09 and SP18 (see the electronic supplementary material, figure S6b), and observed values comparable to that reported for ΔbacA::hyg [16]. To explore further the overlap between B₁₂ uptake and bleomycin susceptibility, we isolated spontaneous bleomycin-resistant (Bleo^R) mutants in two different genetic backgrounds, $\Delta prpDC$ and the unmarked metH knockout, by plating the strains on solid medium containing $3 \mu g \text{ ml}^{-1}$ bleomycin ($10 \times \text{MIC}$). Five Bleo^R mutants each of the $\Delta prpDC$ and $\Delta metH$ knockouts were selected at random, and shown to be defective in their ability to use B_{12} for MutAB-dependent growth in propionate (see the electronic supplementary material, figure S7a). Moreover, the spontaneous $Bleo^R$ mutants of $\Delta metH$ were resistant to CNCbl during growth on solid medium (see the electronic supplementary material, figure S7b). All five Bleo^R mutants derived from the $\Delta prpDC$ strain carried nonsense mutations in Rv1819c, whereas missense mutations in Rv1819c were identified in four of five spontaneous $Bleo^R \Delta met H$ mutants (see the electronic supplementary material, table S2). Moreover, complementation with full-length Rv1819c reversed the inability of the spontaneous Rv1819c point mutants of $\Delta prpDC$ to catabolize propionate in liquid medium supplemented with CNCbl (figure 4), and restored the bleomycin susceptibility of SP09 to wild-type levels (see the electronic supplementary material, figure S6c).

4.6. *Rv1819c* encodes an ATP-binding cassette-type transporter

Rv1819c was previously included in a group of 'BacA-related' proteins identified on the basis of their similarity to the highly conserved BacA and SbmA proteins of *Sinorhizobium* and *E. coli*, respectively [27]. Unlike BacA/SbmA orthologues, however, which are predicted to require an interaction with a separate cytoplasmic protein for function, *Rv1819c* encodes both transmembrane (TMD) and nucleotide-binding (NBD) domains of an ABC transport protein on a single polypeptide. Sequence similarity analyses using only the TMD located *M. tuberculosis* Rv1819c in a cluster distinct from BacA/SbmA (see the electronic supplementary material, figure S8a). Moreover, these analyses indicated that Rv1819c was more closely related to ABC proteins other than BacA/SbmA in both *E. coli* and *Sinorhizobium*, namely YddA [34] and ExsE [35], respectively. The Rv1819c NBD similarly identified YddA

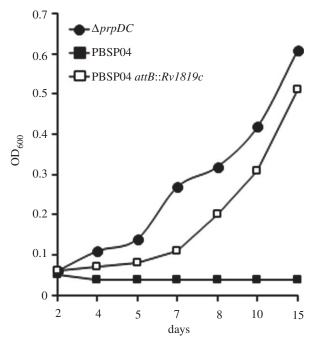


Figure 4. Impaired B_{12} uptake in a spontaneous bleomycin-resistant (Bleo^R) *prpDC* mutant, PBSP04, carrying a SNP in *Rv1819c* (see the electronic supplementary material, table S2). Strains were grown in 0.1% propionate supplemented with 3NP and CNCbl. Data are from a representative experiment performed in duplicate.

and ExsE as close homologues in an equivalent similarity search (see the electronic supplementary material, figure S8b), together with the recently described human ABC-type B_{12} transporter, ABCD4 [36].

We built a homology model of Rv1819c based on the crystal structures of two polyspecific ABC exporters, Staphylococcus aureus Sav1866 [37] and Salmonella typhimurium MsbA [38]. Consistent with known ABC protein architecture [39], Rv1819c is predicted to form a homodimer (figure 5), with each subunit comprising an N-terminal TMD fused to a highly conserved NBD that features all the motifs characteristic of functional ABC transporters (see the electronic supplementary material, figure S9a). Unlike Sav1866 and MsbA, though, the TMD domain of Rv1819c possesses an extra N-terminal region which is predicted to contain an additional transmembrane helix (see the electronic supplementary material, figure S9b). Proteomic analyses in the closely related M. bovis BCG suggest that this region is present in the mature protein [40], and therefore is not a signal peptide. However, in the absence of a close structural template containing seven transmembrane helices, we omitted the first 65 N-terminal residues in building the Rv1819c model. The predicted structure nevertheless provides a useful framework for the interpretation of experimental data. Notably, all three SNPs which resulted in substituted amino acids in the spontaneous B₁₂-resistant and Bleo^R mutants (see the electronic supplementary material, table S2) affect residues located in conserved regions of Rv1819c (figure 5). While the structural consequences of the P349T and G411D mutations require further investigation, L442S affects a conserved position in the putative nucleotidebinding pocket formed by two interacting ABC domains. In Sav1866, the corresponding residue, Ile356, makes a van der Waals contact with the sugar moiety of the bound ADP [37], and so supports the inferred association between a distorted pocket and crippled protein function.

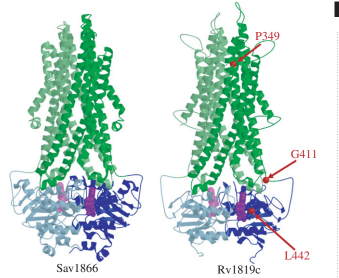


Figure 5. Rv1819c encodes an ABC-type transporter. Computational model of *M. tuberculosis* Rv1819c (amino acid residues 66–639) compared with the x-ray structure of *Staphylococcus aureus* Sav1866 (PDB code: 2HYD) [37]. The model is based on the crystal structures of Sav1866 and the ABC lipid flippase, MsbA, from *Salmonella typhimurium* (PDB code: 3B60) [38], both of which contain transmembrane- (green) and nucleotide-binding (blue) domains fused into a single polypeptide chain that interacts to form a homo-dimer in the active protein. The two subunits in both structures are denoted by the different colour intensities. ADP molecules bound to each subunit are shown in purple and light purple, respectively. Residues substituted in spontaneous *Rv1819c* mutants are indicated with red arrows.

4.7. A PduO-type adenosyltransferase is required for assimilation of vitamin B_{12}

CNCbl must be adenosylated to generate the active cofactor, AdoCbl [41] (figure 6a). The genome of M. tuberculosis is predicted to encode both CobO (Rv2849c) and PduO (Rv1314c) ATP:co(I)rrinoid adenosyltransferases [3], non-homologous enzymes that catalyse this reaction in other bacteria [42,43]. It was notable, therefore, that six putative B₁₂ uptake mutants (figure 2) contained Tn insertions in pduO (see the electronic supplementary material, figure S2d), because this suggested that an impaired ability to convert exogenous CNCbl to the cofactor form could confer B₁₂ resistance in the primary screen, as well as eliminate the ability of M. tuberculosis to use B₁₂ for growth in propionate. To confirm the role of PduOdependent adenosylation in these phenotypes, we evaluated the abilities of the $\Delta met H pdu O$::Tn mutants to assimilate different corrinoids for growth in propionate in the presence of 3NP (figure 6b). The mutants were unable to use either cyano form, CNCbl or (CN)₂Cbi, both of which are adenosylated in the biosynthetic pathway to AdoCbl (figure 6a). In contrast, supplementation with AdoCbl itself restored growth in this assay (figure 6b), suggesting bypass of PduO function. In combination, these observations implicate PduO as sole active adenosyltransferase in M. tuberculosis during growth in vitro.

5. Discussion

Our results identify Rv1819c as sole corrinoid transporter in *M. tuberculosis* under standard *in vitro* conditions and, moreover, establish the capacity of the organism to scavenge

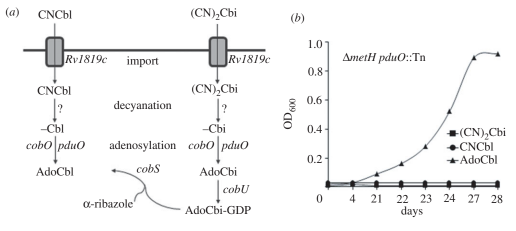


Figure 6. PduO is essential for B_{12} salvage and assimilation. (a) Predicted steps in late-stage AdoCbl biosynthesis and salvage in *M. tuberculosis*. Cobinamide is provided *in vitro* as a dicyanide salt, $(CN)_2$ Cbi. (b) The Δ metH pduO::Tn mutant cannot use CNCbl or $(CN)_2$ Cbi for growth in 0.1% propionate-containing 3NP. Data are from a representative experiment performed in duplicate.

corrinoids. The association of Rv1819c with B₁₂ uptake is unexpected, particularly given previous studies suggesting that Rv1819c might function in ATP-dependent peptide transport [16,44]. Moreover, the properties enabling polyspecific translocation of compounds such as bleomycin, B₁₂ and antimicrobial peptides which lack obvious structural similarity remain unclear [45]. In Gram-positive organisms, ABC-mediated importers function together with a high-affinity substrate-binding protein (SBP) that is anchored to the extracytoplasmic membrane [46]. Although M. tuberculosis possesses in excess of 30 ABC transporters, as well as 15 putative SBPs [47], we failed to identify a candidate B₁₂-binding protein, raising the possibility that Rv1819c-mediated B_{12} import occurs in the absence of a specific SBP or that multiple proteins fulfil this role [48]. In most bacteria, the components of the ABC transporters involved in the uptake of ferric siderophores, haem and vitamin B₁₂ are closely related [49]. However, our screen identified Rv1819c as sole transport candidate, excluding the possibility that other ABC proteins might perform overlapping functions in mycobacterial B₁₂ transport, at least under in vitro conditions. Instead, in associating Rv1819c with B₁₂ uptake, our results add to the expanding complement of atypical mycobacterial nutrient acquisition systems. For example, a novel pathway was recently elucidated that enables the scavenging of haem [50]—a tetrapyrrole which, like B_{12} , is derived from δ -aminolaevulinic acid via a uroporphyrinogen III intermediate [5]. In that system, uptake is mediated by the combined activity of a haem-binding protein and the MmpL family members MmpL3 and MmpL11—predicted RND-type efflux pumps which have been associated with multiple cellular functions [51]. In addition to haem import, recent evidence suggests that MmpL3 fulfils an essential role in exporting trehalose monomycolate across the cell membrane for incorporation into cell wall mycolic acids [52,53], and it has also been implicated in the susceptibility of M. tuberculosis to diverse small molecules [54,55]. It is tempting, therefore, to consider the analogy with Rv1819c-itself a predicted export protein that has now been implicated in the uptake of antimicrobial peptides [16,44] and vitamin B₁₂, and might also play a role in cell wall biogenesis [16].

Rv1819c has been extensively investigated as *M. tuberculosis* BacA [16,44]. Unlike BacA/SbmA orthologues, however, deletion of *Rv1819c* does not render *M. tuberculosis*

hypersusceptible to other antimicrobial drugs and cell disrupting agents [16]. Moreover, our structural model of M. tuberculosis Rv1819c predicts an ABC transporter comprising both TMD and NBD within a single polypeptide. This distinguishes the mycobacterial protein from BacA proteins in Brucella and other intracellular pathogens [32] that contain the TMD only and, importantly, is supported by sequence analyses that situate Rv1819c in a separate cluster from the BacA subfamily even when based on TMD sequence alone. The M. tuberculosis protein also differs from BacA proteins in its potential role in pathogenesis. While the essentiality of the mycobacterial protein for the maintenance of chronic infection in vivo [16] is reminiscent of BacA-like phenotype, closer inspection of the comparative in vivo infection dynamics of different 'bacA' mutants suggests divergent function: for example, in contrast to the Brucella and Sinorhizobium deletion mutants [27,32], the M. tuberculosis Rv1819c knockout is not impaired in its ability to establish an infection [16]. It is tempting, therefore, to consider the virulence defect of the Rv1819c deletion mutant in the light of recent studies describing the accumulation during chronic infection of cholesterol-rich lipid bodies inside foamy macrophages and their infecting bacilli [56,57]. That is, Rv1819c might function to ensure adequate supply of host-derived corrinoids for the B₁₂-dependent utilization of propionate derived from cholesterol catabolism, a possibility that requires further investigation.

Although designed to detect a putative vitamin B₁₂ transporter, our screen also established the essentiality of the PduO-type adenosyltransferase for the assimilation of exogenous corrinoids. The M. tuberculosis genome contains both pduO and cobO adenosyltransferases; therefore, the inferred inactivity of the alternative enzyme in vitro might indicate functional adaptation of CobO to de novo B₁₂ biosynthesis [3], or to specific environmental conditions, including anaerobiosis [41]. Intracellular trafficking of B₁₂ in humans requires the sequential activity of multiple proteins which fulfil dual roles as molecular chaperones and in the enzymatic modification of the cofactor [58]. For example, MMACHC catalyses the reductive decyanation of CNCbl [59] while mediating LMBD1-dependent [60] transfer from the lysosome into the cytoplasm. Recent evidence further suggests that this process is facilitated by the interaction of LMBD1 with ABCD4 [36]—an ABC transporter and homologue of Rv1819c (see the electronic supplementary material,

figure S8). In a subsequent step, the ATP:corrinoid adenosyltransferase attaches the axial ligand and ensures delivery of the resulting AdoCbl cofactor across the mitochondrial membrane to its target enzyme, methylmalonyl-CoA mutase [61]. Given that Tn-mediated disruption of pduO alleviated the B₁₂ sensitivity of the *metH* mutant in the primary screen, it is tempting to speculate that PduO might function not merely in enzymatic conversion of exogenous corrinoids, but also in delivery of the active cofactor into the cytoplasm. Our current model for the translocation of B₁₂ across the mycobacterial cell wall into the cytoplasm therefore proposes the sequential activity of the ABC transporter, Rv1819c and the PduO-type adenosyltransferase (figure 6a).

Our Tn screen also identified four low-frequency insertions associated with compromised B_{12} uptake (figure 2b). Targeted sequencing of Rv1819c and the metE riboswitch in these strains excluded spontaneous mutations as the underlying cause of the observed B₁₂ phenotypes. A single mutant carried an insertion in rpfB, which encodes a resuscitation-promoting factor. To explore this result further, we retested $\Delta metH$ rpfB::Tn in parallel with an rpfB deletion mutant of H37Rv, constructed previously [62]. Although the Tn mutant was not able to use exogenous CNCbl for growth in propionate-containing medium, the $\Delta rpfB$ knockout strain phenocopied wild-type H37Rv in this assay (data not shown), thereby excluding a role for RpfB in B₁₂ uptake. It is possible that rpfB::Tn possesses an additional, unidentified polymorphism, affecting B₁₂ assimilation; alternatively, polar effects on the downstream gene, ksgA, encoding dimethyladenosine transferase, might contribute to the observed phenotype [63], a possibility under investigation. Two additional Tn insertions mapped to mymA, encoding a putative flavin-dependent monooxygenase. The predicted role of MymA in the maintenance of cell wall ultrastructure [64] suggests that compromised B₁₂ uptake in these

mutants might be non-specific; however, this requires further investigation, and is complicated by the fact that *mymA* is the first gene in a seven-gene operon [65]. We also isolated a mutA::Tn mutant, whose inability to use propionate for growth in B₁₂-containing medium is consistent with impaired methylmalonyl-CoA mutase function. The basis for the B₁₂ resistance of this mutant in the primary screen is unclear, however, and probably also the result of an additional spontaneous mutation. The final Tn insertion mapped to Rv2927c, a gene which previous saturation mutagenesis studies have predicted as essential for growth of M. tuberculosis in vitro [66,67]. Although the function of Rv2927c is unknown, it has been proposed to operate as part of the cell division machinery [68]. It seems probable that, like the mymA::Tn mutants, the failure of Rv2927c::Tn to assimilate B₁₂ is non-specific. However, given the inferred requirement for PduO-dependent adenosylation in the assimilation of exogenous B₁₂, the prediction that Rv2927c might function in de novo adenosine nucleotide biosynthesis [69] is intriguing, and the subject of current investigation.

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