Rv0989c encodes a novel (E)-geranyl diphosphate synthase facilitating decaprenyl diphosphate biosynthesis in Mycobacterium tuberculosis

Francis M. Mann
Iowa State University

Jill A. Thomas
Iowa State University

Reuben J. Peters
Iowa State University, rjpeters@iastate.edu

Follow this and additional works at: http://lib.dr.iastate.edu/bbmb_ag_pubs

Part of the Biochemistry, Biophysics, and Structural Biology Commons, Cell and Developmental Biology Commons, Diseases Commons, and the Genetics and Genomics Commons

The complete bibliographic information for this item can be found at http://lib.dr.iastate.edu/bbmb_ag_pubs/103. For information on how to cite this item, please visit http://lib.dr.iastate.edu/howtocite.html.
Rv0989c encodes a novel (E)-geranyl diphosphate synthase
facilitating decaprenyl diphosphate biosynthesis in
Mycobacterium tuberculosis

Francis M. Mann1, Jill A. Thomas1, and Reuben J. Peters2
Department of Biochemistry, Biophysics, and Molecular Biology, Iowa State University, Ames, IA

Abstract

Mycobacterium tuberculosis (Mtb) has a highly complex cell wall, which is required for both
bacterial survival and infection. Cell wall biosynthesis is dependent on decaprenyl diphosphate as
a glyco-carrier, which is hence an essential metabolite in this pathogen. Previous biochemical
studies indicated (E)-geranyl diphosphate (GPP) is required for the synthesis of decaprenyl
diphosphate. Here we demonstrate that Rv0989c encodes the “missing” GPP synthase,
representing the first such enzyme to be characterized from bacteria, and which presumably is
involved in decaprenyl diphosphate biosynthesis in Mtb. Our investigation also has revealed
previously unrecognized substrate plasticity of the farnesyl diphosphate synthases from Mtb,
resolving previous discrepancies between biochemical and genetic studies of cell wall
biosynthesis.

Keywords

Polyisoprenyl phosphates; Terpene Biosynthesis; Tuberculosis; Cell Wall biosynthesis; Genetic
fitness

1. Introduction

Mycobacterium tuberculosis (Mtb) is the main intracellular pathogen responsible for the
human disease Tuberculosis, which is particularly lethal in immune-deficient individuals.
Evolution of drug resistant strains has renewed this pathogen’s threat on the general
population, as these new strains are extremely pathogenic and untreatable by current
antibiotic regimens [1]. Furthermore, decades of research have failed to uncover the
pathogenic mechanism of Mtb, complicating drug design efforts. This elusive mechanism
likely involves a multifactorial attack, employing attack by a variety of virulence factors
over the course of the infection [2]. One set of these virulence factors involve the complex
mycobacterial cell wall and lipids, which have been linked to pathogenesis and survival of
the bacteria during infection [3].
The mycobacterial cell wall is unique amongst bacteria. The cell wall consists of the peptidoglycan, arabinogalactan, and mycolipid layers, as well as a capsule-like layer separate from the core wall components [4]. Important in proper assembly of this complex structure is the glyco-carrier polyprenyl phosphate (Pol-P) [5]. Such Pol-P molecules are synthesized via the addition of isoprene units in the form of isopentenyl diphosphate (IPP) to a variety of allylic isoprenyl diphosphate acceptors, which are found in varying lengths, as well as various double bond isomers. Specifically, Mtb utilizes a 10 isoprene-unit decaprenyl diphosphate to assemble its complex cell wall [6], which is formed by the addition of seven IPP to (Z,E)-farnesyl diphosphate (FPP) by Rv2361c [7]. To synthesize (Z,E)-FPP, one unit of IPP is added to (E)-geranyl diphosphate (GPP) by Rv1086 [7]. Previous work has indicated that this process is dependent on GPP as the allylic acceptor of IPP, yet no GPP synthase has been characterized from Mtb [8] (Figure 1). In addition, while Rv2361c is essential for bacterial survival [9], neither Rv1086 nor the remaining uncharacterized prenyltransferases are similarly essential, complicating identification of the prenyltransferases involved in decaprenyl diphosphate biosynthesis.

While studying the biosynthesis of the immune modulating diterpenoid, isotuberculosinol, we uncovered four previously uncharacterized putative trans-prenyltransferases (Rv0989c, Rv2173, Rv3383c, Rv0562) [10]. Of these prenyltransferases, only Rv0989c failed to synthesize the (E,E,E)-geranylgeranyl diphosphate (GGPP) required for isotuberculosinol or other diterpenoid biosynthesis. These initial assays were not set up to detect shorter chain isoprenoids (i.e. FPP or GPP), leading to a lack of observed activity with Rv0989c. After recognizing that a GPP synthase was expected, we altered our protocol to enable detection of ten and fifteen carbon isoprenoids, and found that Rv0989c acts as a GPP synthase, that seems to assist the production of FPP in Mtb. Furthermore, our analysis indicates that Rv1086 is able to produce small amounts of (Z,E)-FPP from dimethylallyl diphosphate (DMAPP), which may explain the non-essential nature of these prenyltransferases. Accordingly, we describe here the first bacterial GPP synthase, and further clarify the roles of the various isopentyl diphosphate synthases involved in production of the decaprenyl diphosphate required for mycobacterial cell wall biosynthesis, thus resolving the apparent contradictions between biochemical and genetic analysis of these enzymes.

2. Materials and methods

General

IPP, DMAPP, GPP, (E,E)-FPP, and GGPP were purchased from Isoprenoids, LC (Tampa, FL). Corresponding dephosphorylated alcohol standards for gas chromatography (GC) analysis were purchased from Sigma Aldrich (St. Louis, MO). Unless otherwise noted, all other chemicals were purchased from Fisher Scientific (St. Louis, MO) and all molecular biology reagents were purchased from Invitrogen (Carlsbad, CA). Sequence alignments were carried out using the AlignX program from the Vector NTI software package (Invitrogen), using standard parameters.

Cloning, expression, and purification

Rv0989c and Rv3398c were cloned from Mycobacterium tuberculosis CDC1551, while Rv1086 was obtained as a synthetic construct, codon-optimized for expression in Escherichia coli. These were cloned into pENTR/SD/D-TOPO prior to directional recombination into an expression vector containing a 6xHis N-terminal fusion (pDEST17, Gateway, Invitrogen). Proteins were expressed in E. coli strain C41 (Lucigen, Middleton, WI), by growing the recombinant bacteria in NZY liquid media at 37 °C to an OD600 of 0.6, transfer to 16 °C for 1 hr prior to induction with 0.5 mM IPTG, and subsequent 14–16 hr fermentation. Cells were harvested via centrifugation and resuspended in Lysis Buffer (10
mM Tris-Cl, 10% Glycerol, 1 mM Dithiothreitol) for sonication. The resulting lysates were clarified via centrifugation (10 min x 10,000g). The recombinant proteins were then purified over Ni-NTA Superflow resin (Novagen) following manufacturer protocols.

**Prenyltransferase assay**

The purified enzymes were characterized in Assay Buffer (50 mM sodium phosphate (pH 7.0), 10% glycerol, 5 mM MgCl₂, 2 mM Dithiothreitol), with IPP and allylic isoprenyl diphosphate substrate (i.e. DMAPP, GPP, or FPP), in a final volume of 1 mL at 37 °C. Assays were stopped by the addition of 1 ml of Stopping Solution (9:1 Methanol: 3N HCl), which also dephosphorylates the substrates and products, and 1 mL of pentane was added immediately and mixed to allow for full partitioning of the isoprenoid alcohols to the organic solvent layer. This was removed and dried under a stream of N₂ at room temperature prior to resuspension in 50 μL hexanes. Enzymatic products were identified on a gas chromatograph (GC) with mass spectrometry (MS) detection using a Varian 3900 GC with Saturn 2100T ion-trap MS (Varian Inc, Palo Alto, CA) with comparison to authentic standards, and were quantified via an Agilent 6890 GC with flame ionization detection (FID)(Agilent Technologies, Santa Clara, CA), using a previously published method [11].

**Characterization of Rv0989c**

Rv0989c was determined to produce GPP, and then assays with this enzyme optimized for protein concentration and incubation time to obtain initial linear reaction rates. It was necessary to use ≥ 0.5 μM enzyme to achieve linearity, with incubation times of 2 minutes then found to be optimal. Steady-state kinetic constants were determined by varying concentrations of either substrate, i.e. 10–300 μM IPP or 5–200 μM DMAPP, with the other held constant at the highest relevant concentration, which initial analysis had determined was saturating. Concentrations of both the internal standard (cembrene) and product were determined by integration of peaks after separation and detection by GC-FID. The observed data was fit to the Michaelis-Menten equation (KaleidaGraph 4.0, Synergy Software; Reading, PA), resulting in R-values ⩾ 0.97.

**Coupled assays**

Coupled assays were carried out by pre-incubating Rv0989c (5 μM) with DMAPP (100 μM) and IPP (100 μM) for 5 min. prior to the addition of Rv1086 (1 μM) or Rv3398c (0.35 μM), along with an additional 50 μM IPP, and the assay incubated for an additional 5 minutes before stopping. Control assays with Rv1086 or Rv3398c, in the absence of Rv0989c, using 100 μM of GPP or DMAPP as the allylic substrate, were run for 5 minutes. These all were analyzed by GC-MS, as described above, for production of the relevant isomers of FPP using separate methods, allowing for distinct separation of the resulting dephosphorylated farnesol (i.e. (E,E)-farnesol [12] or (Z,E)-farnesol [11]).

**Characterization of Rv3398c and Rv1086**

Rv3398c was kinetically characterized as described above for Rv0989c, using substrate concentrations ranging from 2.5–200 μM. Time course studies with Rv1086 were performed with 100 μM each of IPP and DMAPP. When Rv1086 was assayed alone, Triton X-100 was added at 0.3 % (v/v) to the Assay Buffer to mimic previously reported conditions [8].
3. Results

Characterization of Rv0989c as a GPP synthase

Rv0989c was initially screened for GGPP synthesis based on its homology to other GGPP synthases within the Mtb genome (Rv0562, Rv2173, Rv3383c) [10]. As Rv0989c failed to synthesize GGPP from any allylic precursor (DMAPP, GPP, or FPP), it was then assessed for activity as a GPP or FPP synthase, and found to produce GPP when incubated with DMAPP and IPP (Figure 2). Although production of (E,E)-FPP was occasionally observed in the presence of excess (>1 mM) substrate, this likely indicates non-physiologically relevant reuptake of product for further elongation under these conditions. Steady-state kinetic analysis of Rv0989c revealed unusual behavior. When varying DMAPP concentration in the presence of saturating amounts of IPP, \( k_{cat} \) appeared to be 0.7 min\(^{-1}\), which is over 7.5-fold lower than that measured when varying IPP concentration in the presence of saturating amounts of DMAPP, where \( k_{cat} \) appeared to be 5.1 min\(^{-1}\). Yet the pseudo-substrate binding constant \( K_M \) suggests DMAPP is bound three-fold tighter than IPP (58 versus 163 \( \mu \)M, respectively). This may indicate non-catalytically relevant binding of two molecules of IPP at high concentrations, necessitating displacement by DMAPP that decreases the observed catalytic rate.

Rv0989c facilitates production of (Z,E)-FPP by Rv1086 and (E,E)-FPP by Rv3398c

Decaprenyl diphosphate formation in Mtb proceeds through Rv1086, a (Z,E)-FPP synthase, which has been reported to use exclusively GPP as its allylic isoprenyl diphosphate substrate [8]. Similarly, Rv3398c, the Mtb (E,E)-FPP synthase, also has been reported to utilize GPP exclusively [13]. To demonstrate that Rv0989c synthesizes the required product for further elaboration into these important isoprenyl diphosphates, Rv0989c was assayed with each of these FPP synthases. Coupled assays with Rv0989c and Rv1086 led to production of significant amounts of (Z,E)-FPP (Figure 3), along with smaller amounts (<10%) of (E,E)-FPP (data not shown). Surprisingly, a small amount of (Z,E)-FPP also was produced by Rv1086 from IPP and DMAPP in the absence of Rv0989 (Figure 3). Nevertheless, time course measurements indicate the coupled reaction proceeds at a faster rate and nets higher yields of (Z,E)-FPP (Figure 4), supporting a role for GPP as the preferred substrate for this enzyme. Coupled assays with Rv0989c and Rv3398c produce (E,E)-FPP (Figure 5).

However, again, when assayed only in the presence of DMAPP and IPP, Rv3398c is able to produce (E,E)-FPP at measurable rates. Steady state kinetic analysis of Rv3398c determined that \( k_{cat} \) is 1.5 ± 0.1 min\(^{-1}\), \( K_M \) for IPP is 2.6 ± 0.4 \( \mu \)M, while that for DMAPP is 20 ± 9 \( \mu \)M. Comparing to previous publications [13], this still suggests that GPP (\( K_M = 10 \mu \)M) is the preferred allylic isoprenyl diphosphate substrate for Rv3398c.

4. Discussion

The mycobacterial cell wall has been extensively studied as it is exceptionally reinforced and uniquely immunogenic [3]. Key to the assembly of this complex structure is the glyco-carrier decaprenyl diphosphate [6]. Though clearly required for cell growth, development, and viability [9], the biosynthetic pathways utilized in synthesis of decaprenyl diphosphate have frustrated scientists in recent decades. Genetic and biochemical analysis of the biosynthetic machinery has failed to completely characterize the pathway. While decaprenyl diphosphate production is required for cell wall biosynthesis [14], genetic analysis indicates that only the final step in the production of this important metabolite (that catalyzed by the gene product of Rv2361c) is absolutely required for bacterial viability [9]. While this decaprenyl diphosphate synthase will accept a range of allylic isoprenyl diphosphate substrates [7], it has been reported that the production of its preferred (Z,E)-FPP primer for elongation is completely dependent on the availability of GPP [8]. Previously, neither
genetic nor biochemical analysis was able to identify the corresponding GPP synthase in Mtb, which our results now demonstrate is encoded by Rv0989c.

To the best of our knowledge, Rv0989c is the first characterized bacterial GPP synthase. While this does exhibit homology to other such trans-prenyltransferases, there are a number of surprising substitutions observed in Rv0989c. In particular, while readily aligned with other known GPP synthases (Figure 6), Rv0989c contains remarkable changes within two otherwise well-conserved aspartate-rich motifs (designated as the first and second, FARM and SARM, respectively). In Rv0989c, the second aspartate of the FARM and the first aspartate of the SARM are replaced by arginine, and the final aspartate of the SARM is replaced by an alanine, whereas the only other naturally found substitutions are asparagines. Indeed, site-directed mutations in the first and second aspartate of the SARM of other trans-prenyltransferases to alanine, specifically an (E,E)-FPP synthase from Saccharomyces cerevisiae, have demonstrated a drastic loss of activity [15]. The significance of the observed divergence in the FARM and SARM of Rv0989c is not readily apparent, but their expected deleterious effects may have been a contributing factor to the previous difficulty in identification of a GPP synthase in the Mtb genome.

We have determined Rv0989c is able to provide the GPP precursor required for efficient production of both double bond isomers of FPP in Mtb, as demonstrated by coupled assays with the relevant FPP synthases (encoded by Rv1086 and Rv3398c). However, our enzymatic analysis also revealed that both of these are able to react with the shorter, universal allylic isoprenyl diphosphate substrate DMAPP. This clarifies the contradictory biochemical and genetic evidence regarding the decaprenyl diphosphate biosynthetic pathway in Mtb [8,9,13]. In particular, knockout analysis of these prenyltransferases in Mtb have demonstrated that they are not required for viability, indicating that cell wall biosynthesis was able to proceed in their absence, despite biochemical evidence indicating that decaprenyl diphosphate production depends on their activity/products. Nevertheless, these findings further highlight the need for development and study of functional, non-lethal knockouts in Mtb cell wall biosynthetic machinery, as many questions have yet to be answered regarding bacterial tolerance and flexibility for substitutions in this process. This important component of Mtb viability and pathogenesis needs to be completely understood before the puzzle that is the Mtb pathogenic mechanism will be solved.

Acknowledgments

We thank Dr. Niels Nieuwenhuizen of The New Zealand Institute for Plant and Food Research Limited for the kind gift of a synthetic version of Rv1086, and gratefully acknowledge support for this project from the NIH (GM076324 to R.J.P.).

List of abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mtb</td>
<td>Mycobacterium tuberculosis</td>
</tr>
<tr>
<td>IPP</td>
<td>isopentenyl diphosphate</td>
</tr>
<tr>
<td>DMAPP</td>
<td>dimethylallyl diphosphate</td>
</tr>
<tr>
<td>GPP</td>
<td>geranyl diphosphate</td>
</tr>
<tr>
<td>FPP</td>
<td>farnesyl diphosphate</td>
</tr>
<tr>
<td>GGPP</td>
<td>geranylgeranyl diphosphate</td>
</tr>
<tr>
<td>GC</td>
<td>gas chromatography</td>
</tr>
<tr>
<td>Pol-P</td>
<td>polyprenyl phosphate</td>
</tr>
</tbody>
</table>

FEBS Lett. Author manuscript; available in PMC 2012 February 4.
FID  flame ionization detector
MS  mass spectrometer
FARM  first aspartate-rich motif
SARM  second aspartate-rich motif

References


Figure 1.
Decaprenyl diphosphate biosynthesis in *Mycobacterium tuberculosis*. The roles of Rv1086 and Rv2361c have been demonstrated, yet biochemical analysis indicated they are dependent on a previously uncharacterized GPP synthase, which is demonstrated here to be encoded by Rv0989.
Figure 2. Confirmation of Rv0989c activity
Comparison of dephosphorylated Rv0989c product to GPP-derived GOH. (A) GC-FID chromatograms. (B) GC-MS mass spectra.
Figure 3. Production of (Z,E)-FPP-derived FOH by Rv0989c/Rv1086
GC-MS analysis of product profiles from Rv1086 assayed in the presence of authentic GPP, GPP synthesized by Rv0989c, or DMAPP. (A) Total ion chromatograms. (B) Mass spectra.
Figure 4. Time-dependent production of (Z,E)-FPP by Rv1086
(Z,E)-FPP-derived FOH synthesis by Rv1086 assayed in the presence of 100 μM IPP and
100 μM DMAPP (squares) or 100 μM IPP, 100 μM DMAPP, and Rv0989c (circles). Smooth
fit lines were applied.
Figure 5. Production of (E,E)-FPP-derived FOH by Rv0989c/Rv3398c
GC-MS analysis of product profiles from Rv3398 assayed in the presence of authentic GPP and GPP synthesized by Rv0989c. (A) Total ion chromatograms. (B) Mass spectra.
Figure 6. Alignment of Rv0989c with confirmed GPP synthases
Alignment of the Rv0989c gene product with that of the GPP synthases from *Ips pini* (IpGPS) [16], *Aphis gossypii* (AgGPS) [17], *Arabidopsis thaliana* (AtGPS) [18], and *Citrus sinensis* (CsGPS) [19]. The first- and second aspartate-rich motifs (FARM and SARM, respectively), are underlined, with the non-canonical substitution of arginine or alanine residues for otherwise conserved active site aspartate residues in Rv0989c indicated by * and # above the relevant position, respectively.