The role of GlnD in ammonia assimilation in *Mycobacterium tuberculosis*

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**Summary**

The control of ammonia assimilation in *Mycobacterium tuberculosis* is poorly understood. We have been investigating a regulatory cascade predicted to control the activity of glutamine synthetase (GS). We previously demonstrated that the GS-modifying protein, GlnE (an adenylyl transferase), is essential for *M. tuberculosis* growth. GlnD, a uridylyl transferase, is involved in the control of GlnE activity in other bacteria. In *M. tuberculosis*, *glnD* is arranged in an apparent operon with *amt* and *glnB*; all three genes are up-regulated in a low-ammonia medium. We constructed an in-frame deletion of *glnD* by homologous recombination. The mutant had no growth defect in media containing different nitrogen sources. Total GS activity in culture filtrates was markedly reduced in the mutant, although activity in cell-free extracts remained normal. Virulence was unaffected in both in vitro and in vivo model systems of infection, indicating that the presence of extra-cellular GS is not critical for virulence and that the residual intra-cellular GS activity is sufficient. Thus although GlnD does play a role in the control of ammonia assimilation, it is not required for virulence.

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**Introduction**

*Mycobacterium tuberculosis* is responsible for the largest number of human deaths from a single bacterial agent. Nearly two million people die from tuberculosis each year and more than eight million are newly infected.\(^1\) A better understanding of the basic metabolism of this pathogen could lead to new strategies for eradication. Although progress has been made in understanding some of the nutritional requirements of this organism both in vitro and in vivo, in particular its carbon source acquisition,\(^2,3\) little is known about nitrogen metabolism.

*M. tuberculosis* has four enzymes with glutamine synthetase (GS) activity (GlnA1-4).\(^4,5\) Of these, GlnA1, GlnA3 and GlnA4 synthesise L-glutamine, whereas GlnA2 synthesises D-glutamine and D-isoglutamine required for cell wall biosynthesis.\(^5\) The major GS, GlnA1, is expressed to a high
level and is exported. The role of the secreted enzyme is not clear, but it has been suggested that it may play a role in pH modulation, although the biosynthetic reaction does require a source of ATP.

GS catalyses the production of glutamine from glutamate and ammonia. Since this reaction requires ATP, it needs to be strictly regulated in the presence of excess ammonia to conserve both energy and glutamate pools in the cell. GS activity can be controlled by several mechanisms, including feedback inhibition, transcriptional control of gene expression and by post-translational modification. In *Escherichia coli* a regulatory cascade of three proteins, GlnD, PII and GlnE, is involved in the latter mechanism (Fig. 1). GlnD is a uridylyl transferase which modifies the PII protein. PII in turn controls the activity of GlnE. GlnE is an adenylyl transferase which controls the interconversion of GS and GS-AMP. Transfer of the AMP moiety to the GS enzyme reduces its activity. GS-AMP is hydrolysed enzymatically to GS with the predominant reaction being determined by interaction with the PII protein; PII promotes the adenylylation reaction, whereas PII-UMP promotes the deadylylation reaction. In this way the cells can rapidly control GS biosynthetic activity in response to ammonia availability. An *M. tuberculosis* GlnA1 mutant is auxotrophic for glutamine, and attenuated in macrophages and guinea pigs, suggesting that the assimilation of ammonia via this pathway is required in vivo.

We have previously shown that, in contrast to other bacteria including closely related organisms such as *Streptomycetes coelicolor*, GlnE is an essential gene in *M. tuberculosis*. Thus the control of GS activity seems critical to normal growth. Here we show that *glnD* is not essential and that although it has an effect on GS activity in the cells, it is not required for virulence.

**Materials and methods**

**Culture**

*M. tuberculosis* H37Rv (ATCC25618) was grown in Middlebrook 7H9 plus 10% v/v OADC supplement (Becton Dickinson) and 0.05% w/v Tween 80, Middlebrook 7H10 agar with 10% v/v OADC supplement or TSM media (1.5 g/L K2HPO4, 0.5 g/L KH2PO4, 0.5 mg/L MgSO4, 0.5 mg/L CaCl2, 0.1 mg/L ZnSO4, 0.1 mg/L CuSO4 and 50 mg/L ferric chloride) supplemented with 10% v/v OADC and 0.05% w/v Tween 80. Working pH 7.2. For TSM-high ammonia, 30 mM (NH4)2SO4 was added; for TSM-low ammonia, 0.1 mM (NH4)2SO4 was added, L-amino acids (alanine asparagine, glutamine and glutamate) were added to 3 mM. Growth curves were obtained in 12 mm diameter borosilicate tubes with 4–5 ml media and stirring at 250 rpm with an 8 mm flea. Hygromycin was used at 100 μg/ml and kanamycin at 20 μg/ml.

**Quantitative RT-PCR**

Probes and primers were designed for quantitative PCR for *sigA* (endogenous control), *amt*, *glnB* and *glnD* using the software Primer Express (Table 1). cDNA was synthesised from RNA using RT and random hexamer primers using AMV reverse transcriptase. PCR was carried out in a Taqman 7900 using a standard PCR master mix. For *sigA*, *amt*, *glnB* and *glnD*, the primer pairs were *SigA*-R and *SigA*-F, *Amt*-R and *Amt*-F, *GlnB*-R and *GlnB*-F, and *GlnD*-R, respectively, and the probes used were *SigA*-T, *Amt*-T, *GlnB*-T and *GlnD*-T. The primer and probe concentrations were first optimised. The optimal primer concentration was 300 nM for all four genes, the probe concentration was 100 nM for *sigA* and *glnB*, 125 nm for *glnD* and 200 nm for *amt*. In order to measure relative gene expression levels, standard curves for each primer-probe set were generated using genomic DNA. CT values were converted into the equivalent of ng using the standard curve. Control reactions without RT were used to confirm that there was no significant contaminating genomic DNA present. CT values for genomic DNA were converted to ng and subtracted from the plus RT values. In order to standardise the samples to ensure that equal amounts of cDNA were used, each value was standardised to *sigA* to generate unit-less values. At least three independent RNA samples were assayed in triplicate for each gene.

**Construction of glnD mutant**

We used our previous method for generating delivery vectors with a marker cassette. The delivery vector was constructed by amplifying two regions flanking *glnD* such that an...
in-frame deletion was engineered and cloning them into p1NIL. Primer pairs gap7 CACAAGGATACACACAAC and gap8 CGTCAATGCTGTTGCTGC, and gap5 CAAGACCTGGGGAGACGC and gap6 CAGTTTGTCGGTGCCCTC were used to amplify the upstream and downstream regions and the PCR products were cloned into pGEM EasyT (Promega). The two regions were then excised as KpnI–EcoRI and EcoRI–HindIII fragments respectively and cloned into the KpnI–HindIII sites of p1NIL thereby deleting 1.7 kbp of the glnD gene. The marker cassette from pGOAL19 (hyg, lacZ, sacB) was introduced as a PacI fragment to generate the final delivery vector pKOD3.

A deletion mutant was constructed according to our previous method. Briefly, we treated the vector DNA with UV and electroporated M. tuberculosis to generate a single cross-over strain. This strain was streaked out without antibiotics and double cross-overs selected and screened on 2% w/v sucrose and 50μg/ml X-gal. White colonies were patch tested for hygromycin and kanamycin sensitivity and then screened by PCR for the deletion gene. Potential mutants were confirmed by Southern blotting using XhoI digested genomic DNA and hybridising to a probe derived from the upstream flanking region.

GS assays

Cell-free extracts were generated using the MiniBeadBeater. Culture filtrates were prepared using 0.2 μM filters and the filtrates were concentrated using Centricon 20 units (Amicon). Total GS activity was assayed using the transferase assay, GS activity is given in nmol per minute per mg of total protein.

Virulence assays

THP-1 cells were maintained in culture, treated with PMA to induce differentiation, washed and then infected as described. A total of 5 × 10³ macrophages were infected at MOIs of 1:50 and 1:5 bacteria to macrophage. Extracellular bacteria were removed by washing several times.

**Table 1 Primsers**

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequence</th>
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<tbody>
<tr>
<td>gap5</td>
<td>CAAGACCTGGGGAGACGC</td>
</tr>
<tr>
<td>gap6</td>
<td>CAGTTTGTCGGTGCTGC</td>
</tr>
<tr>
<td>gap7</td>
<td>CACAAGGATACACACAAC</td>
</tr>
<tr>
<td>gap8</td>
<td>CGTCAATGCTGTTGCTGC</td>
</tr>
<tr>
<td>SigA-R</td>
<td>TCTCCTGCAAGGATACACACAAC</td>
</tr>
<tr>
<td>SigA-F</td>
<td>CGTCAGGATACACACAAC</td>
</tr>
<tr>
<td>Amt-R</td>
<td>GAGACCTTGAGACACCCGATTTG</td>
</tr>
<tr>
<td>Amt-F</td>
<td>CGTCTACTCGATTG</td>
</tr>
<tr>
<td>GlnB-R</td>
<td>AATGCTGCCAGCACCTTGGTC</td>
</tr>
<tr>
<td>GlnB-F</td>
<td>GGCAGCCGAGGTTTAC</td>
</tr>
<tr>
<td>GlnD-F</td>
<td>CAGTGCTGCGAAGGGTAGGA</td>
</tr>
<tr>
<td>GlnD-R</td>
<td>GGCAGGCTTCCAGCAT</td>
</tr>
<tr>
<td>SigA-T</td>
<td>AGCCACACTGGCGCTTGGTT</td>
</tr>
<tr>
<td>Amt-T</td>
<td>CATGCGGCAGCACCAGGAC</td>
</tr>
<tr>
<td>GlnB-T</td>
<td>CAATGGAATCGTCAACACACGCTCGAT</td>
</tr>
<tr>
<td>GlnD-T</td>
<td>TTGCATCGCAAATTCCGATCTGA</td>
</tr>
</tbody>
</table>

Determination of the initial inoculum was assessed by plating serial dilutions and the number of intra-cellular bacteria was monitored over 7 days. Mice were infected with approximately 10⁶ viable mycobacteria in 200μl of pyrogen-free saline via a lateral tail vein. Where appropriate, infected mice were killed by cervical dislocation in accordance with humane endpoint protocols under the Animals Scientific Procedures Act, 1986 (UK).

**Results**

Expression of glnD and nitrogen regulation

We are interested in the regulatory cascade that controls the activity of GS by post-translational modification. We previously demonstrated that GlnE, an adenyllyl transferase which modifies GS, is essential in M. tuberculosis. The activity of GlnE is modified by the PII protein (encoded by GlnK or GlnB) and PII is controlled by the GlnD protein (Fig. 1). In order to gain a better understanding of this pathway, we extended our work to look at the other members of the cascade.

The glnD gene is arranged in an apparent operon with two other genes, amt and glnB (Fig. 3A). The start and stop codons of amt and glnB overlap, but there is a 60bp gap between glnB and glnD which could theoretically contain a promoter. We wanted to determine if the genes in this

**Figure 2** Expression in response to ammonia availability. Levels of mRNA were measured using RT-qPCR. The amount of mRNA is given as an arbitrary value standardised to sigA expression values. The mean±standard deviation of at least three independent samples assayed in triplicate is given. Asterisks indicate significant induction in low ammonia (Student’s t-test p<0.05). Key: black bars—low ammonia; white bars—high ammonia.
region are controlled by ammonia availability. We used RT-quantitative PCR to look at the expression levels of the three genes relative to sigA (Fig. 2). Amt and glnB were up-regulated three-fold and glnD was up-regulated two-fold in low-ammonia medium as compared to high-ammonia medium. The lesser induction of glnD is not unexpected as it is often seen with genes that are at the 3’ end of the operon. Alternatively, glnD could be independently expressed from a promoter located in the intergenic region.

**Construction of glnD mutant**

In order to characterise the role of GlnD in the regulatory cascade which controls GS activity, we constructed a deletion mutant. An in-frame deletion of the gene was made in the vector p1NIL and the gene cassette from pGOAL19 containing the hyg, lacZ, sacB genes was inserted. A two-step homologous recombination process was used to generate the mutant. The in-frame deletion and expected genotype was confirmed by Southern blotting (Fig. 3). Out of 32 double cross-overs screened, 12 were mutants. One strain (Tame 69) was selected for further study.

We analysed the ability of the glnDΔ strain to grow in various nitrogen sources, since it has been shown that mutations in the control of GS activity can lead to deleterious effects on growth. We measured growth of the deletion mutant Tame 69 with various nitrogen sources. The mutant was able to use all the nitrogen sources and there was no difference in the growth rates from the wild-type strain (data not shown). Thus the mutant is not compromised in its ability to utilise any of these substrates.

**Control of total GS activity**

In other bacteria, GlnD is involved in the post-translational modification cascade which ultimately controls GS activity by modulating its adenylylation state. It has also been shown to play a role in regulation of nitrogen-controlled genes in Corynebacteria. Therefore, we investigated total GS activity of the mutant grown with ammonia or glutamate.

![Figure 3](image_url) **Figure 3** Construction of glnD mutant. An in-frame deletion covering 1.7 kb of the gene was constructed in the delivery vector pKOD3 and used in a two-step process to make a deletion mutant. (A) The in-frame deletion showing the two PCR products amplified and cloned into the delivery vector. (B) PCR screening for the deletion strains. Primers were used to amplify the glnD deletion region—the size of the amplified product in the deletion strains was 0.5 kbp. Lanes 1–13—double cross-over strains; Lane 14—plasmid pKOD3; Lane 15—negative control; Lanes 3, 4, 6 and 9 have deletion alleles. (C) Southern blotting confirmed the expected genotype. Wild-type (Wt) (5.8 and 3.7 kb) and deletion (Del) (4.1 and 3.7 kb) bands are indicated.
as nitrogen sources (Fig. 4) to determine if deletion of \textit{glnD} affected the activity of GS. The majority of GS is found in the culture filtrate in \textit{M. tuberculosis} and our data confirmed that we had higher activity in this fraction than in the cell-free extracts for the wild-type strain. Interestingly, we found that total GS activity was higher in ammonia-rich conditions (30 mM ammonium sulphate) as compared to either low ammonia (0.1 mM) or 7H9 (glutamate) in the wild-type strain. This was in agreement with RT-qPCR data for \textit{glnA1} which showed increased expression in high-ammonia medium (Pashley and Parish, unpublished data), although \textit{glnA2} expression was unchanged.9 There was no significant difference in the total GS activity between the mutant and wild-type in the cell-free extracts. However, in culture filtrates the situation was very different. We found a large reduction in total GS activity in the mutant strain grown in all media except low ammonia (where the level of activity was at its lowest). This was particularly pronounced in ammonia-rich conditions, where the wild-type activity was at its highest.

**Virulence in macrophages and SCID mice**

\textit{M. tuberculosis} \textit{glnA1} mutants are glutamine auxotrophs and showed a reduced ability to multiply in macrophages and guinea pigs.\textsuperscript{7} Since extra-cellular GS activity was reduced in the \textit{glnD} mutant, we determined whether GlnD played any role in intra-cellular survival. We used two measures of this: (1) the ability of the mutant to grow in macrophages and (2) its ability to cause disease in mice. For the macrophage infection assay we used the human macrophage-like THP-1 cell line. Infections were carried out at two different MOI (Fig. 5). The results showed that there was no difference between the wild-type and mutant strain, indicating that disruption of GlnD function had no effect on intra-cellular survival. We also tested virulence in the SCID mouse model.
to detect any strong attenuation profile (Fig. 5). Again, the mutant strain behaved just as the wild-type and was not attenuated.

Discussion

We have shown that expression of glnB and amt is controlled in response to ammonia levels. The up-regulation of these genes in low-ammonia conditions is in agreement with previous findings in other related bacteria. For example, in Corynebacterium glutamicum, these genes are transcribed as an operon under the control of nitrogen availability; the operon is switched off in nitrogen-rich media and is turned on during nitrogen starvation. 17,18

Our previous results indicate that at least one member of the GS regulatory cascade is essential, as we have been unable to construct GlnE mutants in M. tuberculosis. 6,19 Our current hypothesis is that GlnE is required to inactivate GlnA1 because the latter is expressed at such a high level. If it were all enzymatically active, the intra-cellular levels of glutamate and/or ATP would quickly be depleted. In agreement with this we have shown that the amidolylation function is essential, but the deamidolylation function is not (Pashley et al., unpublished). Deletion of GlnD would result in a lack of amidolylation of the PII protein. Since unmodified PII promotes the amidolylation activity of GlnE, inactivation of GS should still occur in the mutant and therefore deletion of GlnD is possible. However, our hypothesis predicts that PII is likely to be essential since it directly modulates GlnE activity. This concurs with our preliminary evidence that we are unable to make a glnB deletion mutant, in contrast to the relative ease of constructing a glnD mutant (Pashley and Parish, unpublished data).

Although the glnD mutant was viable, there were substantial changes in the level of GS activity. Normally, a large amount of GS is exported from the cell during growth. 6 Export of GS has been linked to high levels of expression, rather than any specific export mechanism. 19 If glnD deletion leads to reduced expression of glnA1, this would explain why only extra-cellular levels are depleted. The observation that total GS activity in the culture filtrates was markedly reduced in the glnD mutant is an intriguing one. It is possible that overall expression of GlnA1 is markedly reduced and therefore little of it will be exported to the outside. Alternatively, GlnA1 production could be completely abolished. The enzyme assay we used does not distinguish between the four M. tuberculosis GS enzymes, so GS activity measured could arise from GlnA2-4. However, this seems unlikely, since it has already been demonstrated that deletion of the glnA1 gene leads to glutamine auxotrophy, implying that all the biosynthetic activity in the cell is GlnA1. 7

It is likely that the GlnD regulatory cascade only controls activity of GlnA1, although we cannot rule out the possibility that GlnA3 activity would also be modulated. However, GlnA2 and GlnA4 will not be controlled by this cascade (since they cannot be modified by GlnE). Thus GS activity arising from enzymes other than GlnA1 cannot be ruled out. Further work to determine the levels of expression of each GS would help to further elucidate the mechanism of control of GS activity in the whole cell. However, the fact that GlnA1 is the major GS in M. tuberculosis confirms the importance of this post-transcriptional regulatory control system.

Although extra-cellular GS was markedly reduced in the glnD mutant, the strain was still fully virulent in macrophages and SCID mice. This was a surprising observation, since it has already been demonstrated that GlnA1 is required for virulence. 7 Our data indicate that GS is not required at high levels for virulence in the SCID mouse and that the presence of the lower level of intra-cellular GS is sufficient for bacterial survival. It is possible that a defect may be seen in the glnD mutant under conditions in which intra-cellular multiplication is seen or in activated, rather than resting macrophages. However, since we have not measured extra-cellular GS directly during infection, we cannot exclude the possibility that extra-cellular GlnA1 is found in the glnD mutant in vivo. Therefore, the attenuation of GlnA1 mutants most likely results from their auxotrophic nature and the lack of availability of glutamine in the host environment.

In conclusion, we have shown that glnD is not an essential gene, but that it is required for the normal expression and activity of GS in culture filtrates of M. tuberculosis. Future work to address the question of how deletion of glnD results in altered expression of GS by determining the mechanism of regulation for ammonia-regulated genes is underway.

Acknowledgements

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References


