# IpsA, a novel Lacl-type regulator, is required for inositol-derived lipid formation in Corynebacteria and Mycobacteria 

Meike Baumgart ${ }^{1}$, Kerstin Luder ${ }^{1}$, Shipra Grover ${ }^{2}$, Cornelia Gätgens ${ }^{1}$, Gurdyal S Besra² ${ }^{2}$ and Julia Frunzke ${ }^{1 *}$


#### Abstract

Background: The development of new drugs against tuberculosis and diphtheria is focused on disrupting the biogenesis of the cell wall, the unique architecture of which confers resistance against current therapies. The enzymatic pathways involved in the synthesis of the cell wall by these pathogens are well understood, but the underlying regulatory mechanisms are largely unknown. Results: Here, we characterize IpsA, a Lacl-type transcriptional regulator conserved among Mycobacteria and Corynebacteria that plays a role in the regulation of cell wall biogenesis. IpsA triggers myo-inositol formation by activating ino1, which encodes inositol phosphate synthase. An ipsA deletion mutant of Corynebacterium glutamicum cultured on glucose displayed significantly impaired growth and presented an elongated cell morphology. Further studies revealed the absence of inositol-derived lipids in the cell wall and a complete loss of mycothiol biosynthesis. The phenotype of the C. glutamicum ipsA deletion mutant was complemented to different extend by homologs from Corynebacterium diphtheriae (dip1969) and Mycobacterium tuberculosis (rv3575), indicating the conserved function of IpsA in the pathogenic species. Additional targets of IpsA with putative functions in cell wall biogenesis were identified and IPsA was shown to bind to a conserved palindromic motif within the corresponding promoter regions. Myo-inositol was identified as an effector of IpsA, causing the dissociation of the IpsA-DNA complex in vitro. Conclusions: This characterization of IpsA function and of its regulon sheds light on the complex transcriptional control of cell wall biogenesis in the mycolata taxon and generates novel targets for drug development.


Keywords: Transcriptional regulator, Corynebacterium glutamicum, Corynebacterium diphtheriae, Mycobacterium tuberculosis, Cell wall synthesis, Mycolic acids, Mycothiol, Inositol, LM, LAM

## Background

The human pathogens Mycobacterium tuberculosis and Corynebacterium diphtheriae, as well as many related species of the order Corynebacteriales, share a distinctive cell wall architecture that plays an important role in virulence and imparts effective protection against harsh environmental conditions and resistance to therapies [1]. One unique feature of these cell walls is the presence of long-chain $\alpha$-alkyl, $\beta$-hydroxy fatty acids, called mycolic acids, which form a second bilayer close to the cell surface that is similar to the outer membrane of Gram-negative bacteria [2].

[^0]The enzymes and regulatory mechanisms involved in biosynthesis of the cell walls are primary targets for the development of new drugs.
A great deal is known about these bacterial cell walls (Figure 1A). The plasma membranes are rich in phospholipids, such as phosphatidylglycerol, diphosphatidylglycerol and phosphatidylinositol (PI). While inositol is an essential polyol in eukaryotes, in eubacteria it is present only in the order Corynebacteriales where it serves both as a cell wall component and as a building block of mycothiol (MSH), the major antioxidant in Mycobacteria, similar to glutathione [3]. One function of PI is to anchor cell wall lipoglycans, specifically PI mannosides (PIMs), lipomannan (LM) and liporarabinomannan (LAM), to the plasma membrane


Figure 1 IpsA plays an essential role in the integrity of the cell wall of Corynebacteriales. (A) Current model of the structure of the cell wall of Corynebacteria and Mycobacteria. (B) Comparison of the organization of the IpsA genome locus in C. glutamicum and related species. IpsA and homologous proteins are shown in black. Data were taken from [5,6]. The percentage identity of the amino acid sequences to IpsA from $C$. glutamicum was taken from NCBI Blast. hp, hypothetical protein. (C, D) Growth curves (C) and phenotype (D) of C. glutamicum wild-type and $\Delta i p s A$. Results in (C) show the growth (backscatter) from sequential cultures in minimal medium with glucose or myo-inositol as carbon source. After reaching the stationary phase, the cells were diluted into fresh medium at an initial $\mathrm{OD}_{600}$ of 1 . Shown are three glucose cultures and the third myo-inositol culture ( $n=3$ ). In ( $D$ ), DNA was stained with 4',6-diamidino-2-phenylindole (DAPI) (cyan) and lipophilic regions with nile red (red). The scale bar is $5 \mu \mathrm{~m}$.
$[1,4]$. The biogenesis of LAM is performed in consecutive steps in the following order: $\mathrm{PI} \rightarrow \mathrm{PIM} \rightarrow \mathrm{LM} \rightarrow \mathrm{LAM}$.
In Mycobacteria, there is a three-step process for synthesis of PI. Step 1 is cyclization of glucose-6-phosphate to 1D-myo-inositol-3-phosphate, a reaction that is catalyzed by a myo-inositol phosphate synthase, Ino1 (Rv0046c) [7]. In step 2, the 1D-myo-inositol-3-phosphate is dephosphorylated by inositol monophosphatase (IMP) [8] to yield myo-inositol. Step 3 is the transfer of diacylglycerol (DAG) from cytidine diphosphate-diacylglycerol (CDP-DAG) to myo-inositol by a PI synthase, PgsA (Rv2612c) [9]. The enzymes Ino1 and PgsA are essential for the viability of Mycobacteria [10,11]. An ino1 mutant of M. tuberculosis grows in vitro only in the presence of high inositol concentrations and was no longer able to kill SCID mice (see Movahedzadeh et al., [10]). In Mycobacterium smegmatis, ino1 has been shown to be essential for growth in the
absence of exogenous inositol [12]. Ino1 has therefore been proposed as a target for the development of new antibiotics [10,13].
In contrast to the wealth of knowledge about the biogenesis of cell walls, little is known about the regulatory circuits that control these processes [4]. IpsA was chosen as target for further studies from a set of mutants strains as a $\Delta i p s A$ mutant displayed a drastically altered cell morphology hinting towards a function in the control of cell wall biosynthesis. The catabolism of myoinositol has been examined in C. glutamicum $[14,15]$ but regulation of myo-inositol formation has not been deeply investigated up until now. Here, a regulator of inositol synthesis in the model organism C. glutamicum is described. Named IpsA, it is a novel LacI-type regulator that acts as an inositol-dependent transcriptional activator of the myo-inositol phosphate synthase gene. IpsA
function is conserved throughout Corynebacteria and Mycobacteria, including the prominent pathogenic species C. diphtheriae and M. tuberculosis.

## Results

IpSA is a novel Lacl-type regulator conserved in the Corynebacteriales
IpsA was assigned by homology to the Lacl family of transcriptional regulators, which often coordinate the available nutrients with the expression of catabolic genes [16]. The corresponding genomic loci display a remarkable level of conservation across the Corynebacteriales (Figure 1B). In previous studies, the structure of IpsA (Cg2910) was analyzed (PDB:3H5T); its physiological function is the subject of this study.

## IpsA deletion affects cell shape and growth

To gain insight into the function and possible target genes of IpsA, an in-frame deletion mutant was constructed. Growth rates and final backscatter of $\Delta i p s A$ cells were significantly below control levels using minimal medium with glucose as carbon source, and these shortfalls increased in magnitude with successive subcultivation (Figure 1C). The morphology of the mutant strain was altered, as revealed by fluorescence microscopy with staining of DNA ( 4,6 -diamidino-2-phenylindole (DAPI)) and membranes (nile red). $\triangle i p s A$ cells formed chain-like structures and failed to divide properly. DNA in the cells was unevenly distributed and was frequently found in the extracellular space, indicative of severe damage to the cell membrane and/or the cell wall (Figure 1D, for a larger sample see Additional file 1: Figure S1). Membrane staining revealed several intact cell septa in the unshaped mutant cells.
Both the growth defect and altered morphology could be partially overcome by plasmid-encoded ipsA under control of the native or an inducible promoter (data not shown). These efforts revealed that increased levels of IpsA also affect growth of C. glutamicum. Full restoration of the wild-type phenotype was only achieved when ipsA was reintegrated into the mutant strain (intergenic region cg1121 to cg1122) and expressed from its native promoter (Figure 2A, B). When the cells were cultivated with myo-inositol instead of glucose as carbon source, no differences were observed between the wildtype and $\triangle i p s A$ strains in terms of growth and morphology (Figure 1C, D).

## The impact of ipsA mutation on transcription

To elucidate the transcriptional changes caused by the deletion of ipsA, DNA microarray experiments were performed. More than 450 genes showed alterations in mRNA level greater than a factor of 2 , using cells in the exponential growth phase $\left(\mathrm{OD}_{600}=5\right)$ cultivated in minimal


Figure 2 Complementation of the $\triangle i p s A$ phenotype with ipsA and the target gene ino1 (cg3323). For chromosomal complementation, the gene ipsA was integrated into the intergenic region of cg 1121 to cg 1122 under control of the native promoter. (A) Growth of the strains in CGXII minimal medium with $2 \%(\mathrm{w} / \mathrm{v})$ glucose as carbon source. The average and standard deviation of three biological replicates is shown. (B) Fluorescence microscopy images of stationary phase cells of the cultures presented in (A). (C) Plasmid-based complementation with the target gene inol (cg3323). The strains were cultivated in CGXII minimal medium with $2 \%(\mathrm{w} / \mathrm{v})$ glucose and $50 \mu \mathrm{M}$ isopropyl $\beta$-D-1-thiogalactopyranoside (IPTG) for induction of inol expression.
medium with glucose as carbon source (Additional file 1: Table S3, full data deposited in GEO database [17] under accession number GSE50210).
The expression of the gene cg3323, which encodes a myo-inositol phosphate synthase (ino1), and of the operon (cg0044 to cg0046), which encodes an uncharacterized $A B C$ transporter, was reduced by more than a factor of ten in the mutant strain (Table 1). Other downregulated genes included a flavin-containing monooxygenase (cg3195), a hypothetical endoglucanase (cg2896), and two further transporters (cg0621 to cg0623 and cg2181 to cg2184) (Additional file 1: Table S3). Among the strongly upregulated genes was a cluster encoding enzymes for inositol catabolism (cg3389 to cg3392), an operon encoding both menE and a putative integral membrane protein (cg0533 to cg0534), and a putative dinucleotide-binding enzyme (cg1421). Many of the affected genes are components of the DtxR stimulon, which plays a central role in iron homeostasis in $C$. glutamicum $[18,19]$. We speculate that these changes are the result of an impaired iron uptake of the mutant strain due to the defects in the cell wall. In combination with the observed growth phenotype of the mutant strain, the alterations in gene expression indicate that IpsA is a major regulator of inositol-derived cell wall components.

## Identification of direct IpsA target promoters

The microarray experiments revealed several putative target genes of IpsA. To determine whether these genes are directly regulated by IpsA, the promoter regions of the respective genes were tested in electrophoretic

Table 1 Transcriptome analysis of an IpsA deletion mutant

| Locus tag | Gene | Annotated function | Ratio |
| :---: | :---: | :---: | :---: |
| cg3323 | ino 1 | Myo-inositol phosphate synthase | 0.047 |
| cg0044 | rbsB | ABC transporter/periplasmic D-ribose-binding protein | 0.082 |
| cg0045 | - | Probable ABC transport protein, membrane component | 0.138 |
| cg2910 | - | Transcriptional regulator, Lacl family | 0.173 |
| cg3195 | - | Flavin-containing monooxygenase | 0.178 |
| cg0046 | - | Probable ABC transport protein, ATP-binding component | 0.323 |
| cg3210 | - | Cell envelope-related transcriptional regulator | 0.411 |
| cg1421 | - | Putative dinucleotide-binding enzyme | 5.214 |
| cg0533 | menE | O-succinylbenzoic acid-CoA ligase | 5.467 |
| cg0534 | - | Putative integral membrane protein | 5.597 |

mRNA ratios ( $\Delta i p s A /$ wild-type) of IpsA target genes in microarray experiments are shown. Presented are all genes where binding of IpsA could be shown in electrophoretic mobility shift assays (EMSAs) as well as IpsA itself, confirming the deletion.
mobility shift assays (EMSAs) for the formation of complexes with purified IpsA protein. We analyzed the promoters of all genes that were more than fourfold regulated but excluded ribosomal proteins and genes belonging to the DtxR/RipA regulon. DNA fragments (approximately 500 bp ) covering the promoter regions of putative target genes, were incubated with increasing IpsA-His concentrations. IpsA-His bound with high affinity to the promoter regions of cg0044 and inol (cg3323) and with lower affinity to the promoter regions of cg0534 (a putative integral membrane protein), cg1421 (a putative dinucleotide binding enzyme), cg1476 (thiC, thiamine biosynthesis protein ThiC), cg1918 (a putative secreted protein), and cg3195 (flavin containing monooxygenase) (Figure 3; all tested fragments are shown in Additional file 1: Figure S3).

## Determination of the IpsA DNA binding site

Using EMSAs, two separate binding sites were identified in the promoter of ino1 and one in the promoter of cg0044 (Figure 3A and Additional file 1: Figure S4). Analysis of these sites using the motif-based sequence analysis tool MEME [20] revealed a conserved palindromic motif, which was subsequently verified by mutational analysis (Figure 4). In particular, the central six base pairs appear to be crucial for IpsA binding. The preliminary motif was used to search for further binding sites in putative target genes and across the genome of C. glutamicum. Binding sites were identified by EMSAs using oligonucleotides in the promoters of cg1421, cg3195, cg3210 and cg0534 (Table 1, Figure 3A and Additional file 1: Figure S5). The identified motifs were used to define the IpsA consensus motif shown in Figure 3C.

## Myo-inositol is an effector molecule of IpsA

The enzyme encoded by the main IpsA target gene is Ino1, a myo-inositol phosphate synthase that catalyzes the formation of 1D-myo-inositol-3-phosphate from glucose-6-phosphate [7]. We tested several metabolites involved in the relevant pathway for their influence on the formation of the IpsA-DNA complex. A clear backshift was observed for myo-inositol but not for 1D-myo-inositol-1-phosphate, 1D-myo-inositol-3-phosphate or glucose-6phosphate (Figure 3B). The effect of myo-inositol could be observed at concentrations down to 0.2 mM (Additional file 1: Figure S6). This effect is specific for IpsA DNA binding, as the addition of up to 50 mM myo-inositol did not have any effect on the formation of an unrelated proteinDNA complex (for example, AcnR and the acn promoter, Additional file 1: Figure S7).

## IpsA is required for inositol-dependent regulation of ino1

 The major function of IpsA revealed thus far is the activation of ino1 in the absence of external inositol.

Figure 3 Target genes, effector and binding sites of IpsA. (A) Binding of IpsA to 30 bp oligonucleotides covering the binding sites in the putative target promoters. As negative control the oligonucleotide cg3323-d (sequence AGGTCTGATTTCTGCTGGGAATCCCCACAT) was used, which is located immediately downstream of the IpsA binding sites in the cg3323 promoter. (B) The influence of intermediates in inositol metabolism ( 5 mM each) on the binding of IpsA to the ino 1 promoter. myo-I, myo-inositol; I1P, 1D-myo-inositol-1-phosphate; I3P, 1D-myo-inositol-3-phosphate; G6P, glucose-6-phosphate. (C) The IpsA consensus motif predicted by MEME [20] and an overview of the sites in the corresponding promoter regions. (D) Promoter-fusion studies using the promoter of ino1 fused to eyfp (pJC1-Pcg3323-eyfp). The specific fluorescence of ATCC 13032 containing this plasmid on glucose was set to 1 and the other values were calculated accordingly.

Consistent with this, the growth and cell morphology of the $\triangle i p s A$ strain is complemented by the constitutive expression of ino1 under standard conditions (Figure 2C). To study the influence of IpsA and myo-inositol on the expression of ino1 in vivo, we fused the inol promoter to eyfp (enhanced yellow fluorescent protein) and monitored the fluorescence output of the wild-type and the $\Delta i p s A$ strain during growth on glucose and myo-inositol (Figure 3D). Strong promoter activity was observed in the wild-type during growth on glucose. Expression of eyfp was significantly decreased in the ipsA deletion strain and was lower still when either strain was cultivated with myo-inositol as the carbon source.

## IpsA function is conserved in C. diphtheriae and $M$. tuberculosis

The conservation of IpsA in the Corynebacteriales suggests that it may play a similar role in pathogenic relatives. To assess this possibility, we tested in vivo complementation of C. glutamicum $\triangle i p s A$ with the IpsA homologs of M. tuberculosis (Rv3575) and C. diphtheriae (DIP1969). Partial complementation of growth and morphology was achieved (Figure 5A-C). Growth was still slightly retarded but the final OD was significantly increased in comparison with the $\Delta i p s A$ strain. Likewise, the morphological phenotype of the complemented strains fell in between the wildtype and the $\Delta i p s A$ strain, with a slightly better complementation by DIP1969 compared to Rv3575. Most cells
displayed the wild-type morphology and, while some chain-like cell clusters were observed (Figure 5C), these were shorter than clusters of the $\Delta i p s A$ strain (Figure 1D).

We also tested the binding of IpsA to the promoter regions of C. diphtheriae and M. tuberculosis homologs of cg3323 and cg0044 using EMSAs (Figure 5D). C. glutamicum IpsA bound strongly to the C. diphtheriae promoters DIP0115 and DIP0021 and the binding sites were determined (Figure 5E and Additional file 1: Figure S8). In $M$. tuberculosis, Rv0046c is organized in an operon with Rv0047c. A binding site was identified within the open reading frame (ORF) of Rv0047c, suggesting the occurrence of a second promoter upstream of inol (Figure 5E).

## Mycothiol synthesis is abolished in C. glutamicum $\triangle i p s A$

Besides being a precursor of PI and derived lipids, myoinositol is one of the building blocks of mycothiol, the major antioxidant of Corynebacteria and Mycobacteria. Therefore, we measured mycothiol in a set of our strains (Figure 6) using a method previously published for $C$. glutamicum [21]. The mycothiol peak was identified by comparison of wild-type chromatograms with those of a mutant depleted in mycothiol synthesis ( $\Delta m s h C$ ). To proof that the corresponding peak, lacking in $\Delta m s h C$, represents mycothiol, thiols were blocked with NMM ( N -methyl-maleimide) prior to derivatization with bromobimane. In this sample, the largest peak of the wildtype at 5.6 minutes was indeed absent the two control


Figure 4 Mutational analysis of one of the IpsA binding sites in the ino1 promoter. The importance of the predicted DNA sequence motif for IpsA binding was tested in electrophoretic mobility shift assays (EMSAs) with DNA fragments in which three nucleotides of the proposed motif were exchanged, as indicated. A + indicates that the mutated fragment was bound with the same affinity as the unaltered wild-type fragment (positive control); (+) indicates that the mutated fragment was shifted, but with lower affinity; - indicates that the mutated fragment was not shifted. In the case of M3 and M4, binding was completely abolished by the mutation, indicating that the six central base pairs are crucial for IpsA binding. For M1, M2 and M6 a slight decrease of binding was observed. The binding of M5 was unchanged.
samples. In fact, this mycothiol peak was absent in the $\triangle i p s A$ strain but could fully be restored by plasmidbased expression of ipsA (Figure 6). Interestingly, the mycothiol measurements showed a partial complementation with the ipsA homologs from C. diphtheriae and M. tuberculosis, which is in line with our previous findings regarding growth and morphology of crosscomplemented strains (Figure 5).

## PI-based glycolipid synthesis is lost in C. glutamicum $\Delta i p s A$

To study the effect of IpsA on the formation of PI-based phosphoglycolipids, $\left[{ }^{14} \mathrm{C}\right]$-labeled polar lipids were extracted from wild-type and $\Delta i p s A$ C. glutamicum. The
glycolipid profiles were examined using two-dimensional thin-layer chromatography (2D-TLC) (Figure 7). The wild-type strain showed presence of $\mathrm{AcPIM}_{2}$, PI, Gl-A (GlcAGroAc 2 ) and Gl-X (ManGlcAGroAc ${ }_{2}$ ) in lipid extracts as compared to known standards and previous studies [22], isolated from the cells grown with either myo-inositol or glucose. Under normal conditions, C. glutamicum produces both PI and Gl-X, which is an alternative glycolipid anchor to PI for synthesis of Cg-LM. The glycolipid Gl-X is derived from Gl-A, its precursor product. Although both PI and Gl-X can be visualized on TLC analysis, the formation of Gl-X is distinguished only in the case of mutants lacking PI, as both Gl-X and PI do not separate out as distinct spots [23,24]. However,



E

| cg3323-B | ACCTGATC $G A T C A A G C$ |
| :--- | :--- |
| cg3323-A | AATTGATC $G$ ATCAAGC |
| DIP0115 | ACTTAACC $G$ ATTAACC |
| DIP0021 | TCTTTACC $A T T C A G T ~$ |
| Rv0047 | GGTGACTCGATCAACA |
| cg0044 | ACTGAATCAATAAAGA |



Figure 5 IpsA in Corynebacterium diphtheriae and Mycobacterium tuberculosis. Corynebacterium glutamicum $\Delta i p s A$ carrying the plasmids pAN6, pAN6-DIP1969 or pAN6-rv3573 and C. glutamicum wild-type with pAN6 as control were cultivated in CGXII with glucose without isopropyl $\beta$-D-1-thiogalactopyranoside (IPTG) (A) or with $50 \mu \mathrm{M}$ IPTG (B). (C) Microscopic phenotypes of the complemented strains. DNA was stained with 4',6-diamidino-2-phenylindole (DAPI) (cyan) and lipophilic regions with nile red (red), scale bar $5 \mu \mathrm{~m}$. (D) Oligonucleotides ( $30 \mathrm{bp}, 1 \mu \mathrm{M}$ for DIP0021 and DIP0115, $0.5 \mu \mathrm{M}$ for Rv0047 and the negative control) covering the predicted binding sites in the promoter regions of the respective genes were incubated with IpsA at the given concentrations and analyzed on $15 \%$ native polyacrylamide gels. In M. tuberculosis, Rv0046 is organized in an operon with Rv0047. A binding site was identified within the open reading frame (ORF) of Rv0047, suggesting the occurrence of a second promoter upstream of ino1. (E) lpsA binding motif derived from the high affinity C. glutamicum targets and the C. diphtheriae and $M$. tuberculosis binding sites.

Gl-A appears as a distinct spot on the TLC results in both wild-type and mutant strains. For convenience, the TLC results for polar lipid extracts from the ipsA
mutant defective in PI synthesis have been marked to depict $\mathrm{Gl}-\mathrm{X}$ at the same position at which PI is depicted in the wild-type and complemented strains. The labeling


Figure 6 Determination of mycothiol production in Corynebacterium glutamicum wild-type and different mutants and complemented strains. The mycothiol was derivatized using bromobimane, separated by high-performance liquid chromatography (HPLC) and monitored using a fluorescence detector (390 nm excitation and 475 nm emission). As control, the mycothiol deficient strain $\Delta m s h C$ was used as well as a wild-type sample that had been treated with N -methyl-maleimide (NMM) to block thiols prior to derivatization. The peak assumed to be mycothiol is marked with a dotted line. Presented is a representative chromatogram of three biological replicates each.
for Gl-A remained unchanged. Interestingly, the complete absence of $\mathrm{AcPIM}_{2}$ was observed in lipid extracts isolated from C. glutamicum $\Delta i p s A$ when cultured in minimal media supplemented with glucose indicating the loss of PI-based glycolipid synthesis in the mutant (Figure 7B). The lipid profiles of complemented strains C. glutamicum
$\triangle i p s A:: \mathrm{pK} 18-\mathrm{ipsA}$ and C. glutamicum $\operatorname{\Delta ips} A$ harboring cg3323 and DIP1969 in plasmid pAN6, were identical to the wild-type when grown on glucose, demonstrating successful complementation of the loss of gene function in these strains (Figure 7C-E). However, the lipid extracts from C. glutamicum $\Delta i p s A$ harboring Rv3575 in plasmid pAN6 cultured under the same conditions displayed no complementation as none of the components was stained with phosphate stain (results not shown). The lipid extracts isolated from the mutant and complemented strains cultured in minimal media supplemented with myo-inositol were similar to wild-type, with distinct $\mathrm{AcPIM}_{2}$ and PI production (Figure 7G, H and Additional file 1: Figure S9).

## Analysis of lipoglycan extracts

Since the lipoglycans LM and LAM are built on PIMs which are mannosylated products of PI, the effect of IpsA was also studied on these lipoglycans extracted from the wild-type, C. glutamicum $\Delta i p s A$ and the complemented strains (Additional file 1: Figure S10A). LAM and Cg-LMA/B were clearly present in extracts from the wild-type cultured in glucose or myo-inositol. However, the lipoglycan extracts from the mutant C. glutamicum $\Delta i p s A$ grown on glucose revealed no LAM while a band corresponding to Cg-LM-B could be observed. On glucose the complemented strains C. glutamicum $\Delta i p s A:: \mathrm{pK} 18-\mathrm{ipsA}$ and $C$. glutamicum $\Delta i p s A$ containing plasmid encoded Cg3323 and DIP1969 exhibited lipoglycan profiles identical to the wild-type as distinct bands corresponding to both LAM and $\mathrm{Cg}-\mathrm{LM}-\mathrm{A} / \mathrm{B}$. However, no band corresponding to LAM was observed in lipoglycan extracts of C. glutamicum $\triangle i p s A$ complemented with Rv3575c in pAN6 plasmid grown in minimal media supplemented with glucose. Instead, a band corresponding to Cg-LM-B was evident. The lipoglycan extracts isolated from the mutant and complemented strains cultured in minimal media supplemented with myo-inositol were similar to wild-type with distinct LM and LAM (Additional file 1: Figure S10B).

## Discussion

The metabolic pathways for biogenesis of the cell wall in Mycobacteria and Corynebacteria are well characterized, with few gaps remaining to be filled. In contrast, the regulation of these processes remains largely undiscovered. Here, we describe the novel transcriptional regulator IpsA, which functions as an activator of ino1 and other target genes most likely involved in the synthesis of inositol-derived cell wall components and mycothiol (MSH) in Corynebacteria and Mycobacteria (Figure 8). Myo-inositol was shown to act as effector of IpsA and our studies are consistent with a conserved function for this regulator throughout the Corynebacteriales.
The polyol myo-inositol has an important function in Mycobacteria and Corynebacteria: it serves as the key


Figure 7 Two-dimensional thin-layer chromatography (2D-TLC) analysis of $\left[{ }^{14} \mathrm{C}\right]$-labeled polar lipids from Corynebacterium glutamicum cells grown in glucose (A-F) or myo-inositol (G-H). (A) and (G) C. glutamicum ATCC 13032, (B) and (H) ATCC 13032 $\Delta i p s A$, (C) ATCC $13032 \Delta i p s A$ pAN6-cg3323, (D) ATCC 13032 $\Delta i p s A::$ pK18int-ipsA, (E) ATCC 13032 $\Delta i p s A$ pAN6-DIP1969 and (F) ATCC 13032 IipsA pAN6-Rv3575. The cells were cultured in CGXII with either glucose (A-F) or myo-inositol (G, H). PI, phosphatidylinositol; TMCM, trehalose monocorynomycolate; GMCM, glucose monocorynomycolate; $\mathrm{Ac}_{1} \mathrm{PIM}_{2}$, monoacylated phosphatidyl myo-inositol dimannoside; $\mathrm{GlcAGroAc}_{2}$, 1,2-di-O-C $\mathrm{C}_{16} / \mathrm{C}_{18,1}$-(a-D-glucopyranosyluronic acid)-( $1 \rightarrow 3$ )-glycerol (GL-A); ManGlcAGroAc2 1,2-di-O-C $\mathrm{C}_{16} / \mathrm{C}_{18: 1}$ -(a-D-mannopyranosyl)-(1 $\rightarrow 4$ )-(a-D-glucopyranosyluronic acid)$(1 \rightarrow 3)$-glycerol (GL-X).
building block for a large class of cell wall phospholipids, as well as for the antioxidant MSH [1,3]. Several studies have illustrated the potential of the underlying biosynthetic pathways as putative drug targets [13]. For example, an ino1 mutant of $M$. tuberculosis can only grow in the presence of high concentrations of myo-inositol [10] and a M. tuberculosis mutant defective in MSH biosynthesis requires catalase for growth on plates [25]. Our studies have shown that the deletion of ipsA leads to a complete loss of mycothiol biosynthesis, demonstrating the importance of IpsA also for this metabolic pathway.
Myo-inositol and its phosphate derivatives are common substances in eukaryotes and are therefore present in significant quantities in the natural habitats of $M$. tuberculosis, C. diphtheriae (the human body) and C. glutamicum (soil) [26]. It has been shown that myoinositol can be used by C. glutamicum as a carbon and energy source, and several transporters, enzymes and regulators of the catabolic pathway have been identified $[14,15]$. In the present study, microarray analysis revealed that the two main target genes of IpsA, ino1 and cg0044, are strongly downregulated during growth on inositol compared to growth on glucose, which is in line with IpsA being an inositol-dependent activator of both genes/operons [14].
Bioinformatic analysis of $\operatorname{ips} A$ and its homologs in other Corynebacteria and Mycobacteria revealed conservation of the gene among these species (Figure 1B). In many genomes of the Corynebacteriales, ips $A$ is located next to $z n u A C B$, which encodes a putative zinc transport system regulated by Zur [27]. Furthermore, genes involved in the biosynthesis of trehalose (otsAB), which is also an important component of the cell wall of the Corynebacteriales, are frequently located downstream of ips $A$, but their orientations are divergent [28,29]. The product of the main target gene of IpsA, Ino1, contains a zinc ion in its catalytic center and needs $\mathrm{NAD}^{+}$for its reaction [30].
The transcriptional organization of ino1 differs between Corynebacteria and Mycobacteria. While ino1 is the only gene in the transcriptional unit in Corynebacteria, it is cotranscribed with a preceded, uncharacterized PadR-like transcriptional regulator in Mycobacteria [31] (Additional file 1: Figure S11). Given the genomic organization, we suspect that this regulator is an additional player in the control of ino1 expression.

That ino1 is the main target gene of IpsA is supported by the observation that the ipsA phenotype was complemented by overexpression of ino1. One could assume that this is the only real target gene of IpsA, but it is not unlikely, that other target genes only appear to be important under different conditions, for example, another carbon source or certain stress conditions. Based on DNA


Figure 8 Model of IpsA function. In Corynebacteria and Mycobacteria, myo-inositol (mlno) is an important building block for cell wall components and mycothiol. mino can be taken up from the culture medium or synthesized from glucose-6-phosphate (G6P) via 1D-myo-inositol-3-phosphate (I3P). When mino concentration is low, IpsA activates ino1, encoding myo-inositol phosphate synthase, which catalyzes the formation of I3P from G6P. Besides ino1, IpsA activates or represses several other targets of unknown function.

Microarrays and EMSAs we identified several other direct target genes of IpsA, all of which were previously uncharacterized. The operon cg0044 to cg0046 encodes an ABC transporter of unknown function. It may be involved in the uptake of inositol(-derived) carbohydrates required for cell wall biogenesis. The product of $\operatorname{cg} 3210$ is annotated as a cell envelope-related transcriptional regulator of the LytR type. It has a transmembrane domain and is conserved in Corynebacteria and Mycobacteria. Cg1421 is a putative dinucleotide-binding enzyme that has some similarity with coenzyme F420-dependent NADP oxidoreductases. Cg3195 is annotated as a flavin-containing monooxygenase and has, like Cg1421, a $\operatorname{NAD}(\mathrm{P})$ binding domain. As many MSH-dependent detoxification reactions require NAD or NADP as cofactors [3], Cg1421 and Cg3195 may be involved in cofactor supply or regeneration for these processes. The operon cg0534 to cg0533 encodes a putative integral membrane protein of the YphA family and men $E$, an $O$-succinylbenzoic acid-CoA ligase. Although the functions of the target genes besides inol are largely unknown, there are some clear links to cell wall synthesis and regulation, and likely also to MSHassociated reactions. Thus, our analysis of the IpsA regulon has revealed several novel and interesting targets for further studies.
Among the target genes of IpsA are not only activated but also repressed genes. The localization of the IpsA binding site in the particular promoter regions matches very well with the type of regulation (Additional file 1: Figure S5). For the activated genes (cg3323, cg0044, cg3210, and cg3195) IpsA binding sites are located upstream of the -35 region. In the promoters of the repressed genes (cg1421 and cg0534) the IpsA binding sites overlap with the -35 and/or the -10 region. This is a further
indication that the altered mRNA level we see in the DNA microarrays is a direct effect of the IpsA deletion.

The structure-function relationship of IpsA is comparable to other members of the LacI family of transcriptional regulators. They all contain a DNA-binding domain at the N-terminus of the protein, composed of three $\alpha$-helices that form a helix-turn-helix motif that binds to the major groove of the operator. The tetramerization of LacI takes place at the C-terminus, at residues 340 to 357, (Additional file 1: Figure S12) [32]. These residues are missing in IpsA, which is in agreement with our gel filtration analysis that indicated a dimeric conformation for IpsA. A computational analysis of many LacI regulators and binding sites revealed a highly conserved hydrophobic amino acid at position 54 (mostly leucine, L64 in IpsA), which inserts into a conserved, central CG group in the operator. This group is present in both binding motifs of the ino1 promoter in C. glutamicum [33].
The formation of 1D-myo-inositol-3-phosphate from glucose-6-phosphate by Ino1 represents the first step of PI synthesis; it provides the scaffold on which cell wall components such as PIMs, LM and LAM are built. Our chromatographic analysis of radiolabelled polar lipid extracts of the mutant C. glutamicum $\Delta i p s A$ grown in minimal media with glucose generated a characteristic phenotype, namely that the mutant strain was deficient in $\mathrm{AcPIM}_{2}$ and PI while Gl-A and Gl-X were unaffected. In addition, the mutant strain grown on minimal media supplemented with glucose failed to produce PI-based LAM. Instead, the GL-X-based LM, Cg-LM-B was apparent on sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) analysis of lipoglycans, which could explain why C. glutamicum $\Delta i p s A$ can grow over several cultivations without supplementation with
myo-inositol [1]. This LM-B was not found in Mycobacteria. The polar lipid and lipoglycan extracts from the complemented strain C. glutamicum $\Delta i p s A::$ pK18-ipsA grown on minimal media supplemented with glucose displayed normal levels of $\mathrm{AcPIM}_{2}$, PI, LM and LAM biosynthesis, with profiles identical to wild-type. These data are consistent with the view that IpsA affects an early stage of PI biosynthesis, namely the expression of ino1. In fact, the polar lipid and lipoglycan extracts obtained from $\Delta i p s A$ complemented with plasmid-encoded ino1 also displayed lipid profiles identical to that of the wild-type, indicating involvement of IpsA in transcriptional activation of ino1 expression. The results imply that in absence of IpsA, PI-based PIMs, LM and LAM are not synthesized as the mutant strain no longer produces the inositol monophosphate synthase required for synthesis of myo-inositol, a substrate for PI production. The polar lipid extracts of C. glutamicum $\Delta i p s A$ complemented with the homolog from C. diphtheriae exhibited a profile akin to wild-type. Thus DIP1969 is a functional equivalent of IpsA and can be designated as transcriptional activator of ino1 (DIP0115) in this pathogenic species. For the homolog of Rv3575 the situation is not as clear as for DIP1969. Although we see a positive effect on growth and a partial complementation in terms of mycothiol biosynthesis and morphological phenotype, no PI derived lipids could be identified on the TLC results. One possible explanation for this is that most of the inositol formed is channeled towards mycothiol. Consequently, a low amount of PI derived lipids might be formed, sufficient for partial complementation of the phenotype but still too low for detection by TLC. Furthermore, the differences in M. tuberculosis IpsA structure, binding affinity and specificity can also be a reason for the only partial complementation. The fact that we did observe partial complementation with respect to most phenotypes tested hints at least to a very similar function in M. tuberculosis.

## Conclusions

In this study, we have characterized the LacI regulator IpsA as a first regulator of inositol anabolism in the Corynebacteriales. The findings provide insights into the regulatory mechanisms involved in processes that have been identified as promising targets for drug development. IpsA itself could be such a target, if a myo-inositol analog could be developed that blocks activation of ino1; it is noteworthy that a trehalose analog with a bacteriostatic effect on Mycobacterium aurum has been reported [34]. Finally, the IpsA regulon delivers a promising starting point for the analysis of further, so far unstudied players in cell wall synthesis in this important class of bacteria.

## Methods

## Bacterial strains, plasmids and growth media

The bacterial strains and plasmids used in this study are listed in Additional file 1: Table S1. The C. glutamicum type strain ATCC 13032 was used as wild-type. Growth experiments were performed at $30^{\circ} \mathrm{C}$ and $1,200 \mathrm{rpm}$ in a Biolector system (m2p-labs, Baesweiler, Germany) in 48well FlowerPlates containing $750 \mu \mathrm{~L}$ CGXII minimal medium [35] supplemented with 3,4-dihydroxybenzoate ( $30 \mathrm{mg} / \mathrm{L}$ ) and carbon source as indicated. If appropriate, $25 \mu \mathrm{~g} / \mathrm{mL}$ kanamycin was added. All cloning was performed in Escherichia coli $\mathrm{DH} 5 \alpha$ cultivated at $37^{\circ} \mathrm{C}$ in lysogeny broth (LB, [36]) with $50 ~ \mu \mathrm{~g} / \mathrm{L}$ kanamycin.

## Recombinant DNA work

Routine methods such as polymerase chain reaction (PCR), DNA restriction and ligation were performed using standard protocols [36-38]. The oligonucleotides used in this study were obtained from Eurofins MWG Operon (Ebersberg, Germany) and are listed in Additional file 1: Table S2. DNA sequencing was performed by Eurofins MWG Operon (Ebersberg, Germany). The $\operatorname{ipss} A$ mutant of C. glutamicum was constructed via a two-step homologous recombination protocol as described previously [39]. For further details regarding plasmid and mutant construction, see Additional file 1.

## DNA microarrays

For transcriptome analysis, C. glutamicum wild-type and $\Delta i p s A$ cells were grown in 5 mL BHI (brain-heart infusion, Becton, Dickinson and Company, Le Pont de Claix, France) for about 6 h at $30^{\circ} \mathrm{C}$. A second precultivation was performed in CGXII minimal medium containing $2 \%(\mathrm{w} / \mathrm{v})$ glucose as carbon source. The main cultures were inoculated to an OD of 0.5 in CGXII minimal medium with $2 \%$ (w/v) glucose. At an OD of 5 the cells were harvested by centrifugation ( $4,120 \mathrm{~g}, 10$ minutes and $4^{\circ} \mathrm{C}$ ). The cell pellet was subsequently frozen in liquid nitrogen and stored at $-70^{\circ} \mathrm{C}$. The preparation of total RNA was performed as described previously with the RNeasy Kit from Qiagen (Hilden, Germany) [40]. Synthesis of fluorescently-labeled cDNA was carried out as described in [40,41]. Purified cDNA samples to be compared were pooled and the prepared two-color samples were hybridized at $65^{\circ} \mathrm{C}$ while rotating for 17 h using Agilent's Gene Expression Hybridization Kit (Agilent, Böbligen, Germany), hybridization oven and hybridization chamber. After hybridization, the arrays were washed using Agilent's Wash Buffer Kit according to the manufacturer's instructions. Fluorescence of hybridized DNA microarrays was determined at 532 nm (Cy3) and $635 \mathrm{~nm}(\mathrm{Cy} 5)$ at $5 \mu \mathrm{~m}$ resolution with a GenePix 4000B laser scanner and GenePix Pro 6.0 software (Molecular Devices, Sunnyvale, CA, USA). Fluorescence images were saved to raw data files in TIFF format (Gene

Pix Pro 6.0). Quantitative TIFF image analysis was carried out using GenePix image analysis software and results were saved as GPR-file (GenePix Pro 6.0). For background correction of spot intensities, ratio calculation and ratio normalization, GPR-files were processed using the Bio Conductor R-packages limma and marray [42]. Array data were deposited in the GEO database [43] under accession number GSE50210.

## Overproduction and purification of IpsA

E. coli BL21(DE3) carrying the expression plasmids pET-TEV-ipsA or pAN6-ipsA-STREP were grown in LB medium at $37^{\circ} \mathrm{C}$ and 120 rpm . IpsA overproduction was induced by addition of $50 \mu \mathrm{M}$ isopropyl $\beta$-D-1-thiogalactopyranoside (IPTG) followed by cultivation at $16^{\circ} \mathrm{C}$ for 20 h before the cells were harvested by centrifugation. Nickel chelate and StrepTactin affinity chromatography were performed as described previously [44,45]. For further purification and determination of the molecular weight, gel filtration was performed using a Superdex 200 10/300 GL column (GE Healthcare, Munich, Germany) (buffer: 50 mM Tris- $\mathrm{HCl} \mathrm{pH} 8,250 \mathrm{mM} \mathrm{NaCl}, 1 \mathrm{mM}$ dithiothreitol (DTT)). Note that the IpsA-variant with the C-terminal STREP-tag formed less aggregates than the variant with the His $_{10}$-tag and eluted at a volume corresponding to 79 kDa , suggesting dimer formation (Additional file 1: Figure S2). The protein was concentrated, flash frozen in liquid nitrogen and stored in the gel filtration buffer at $-70^{\circ} \mathrm{C}$.

## Electrophoretic mobility shift assays (EMSAs)

EMSAs were performed as described previously with the following modifications [46]. Unless otherwise stated, the STREP-tag variant was used for the EMSAs. Purified IpsA was incubated with DNA fragments (30 to 500 bp , final concentration 0.028 to $1 \mu \mathrm{M}$ ) in binding buffer ( 50 mM Tris- $\mathrm{HCl} \mathrm{pH} 7.5,40 \mathrm{mM} \mathrm{KCl}, 5 \mathrm{mM} \mathrm{MgCl} 2$ ) in a total volume of $10 \mu \mathrm{~L}$. Electrophoresis was performed with $10 \%$ to $15 \%$ native polyacrylamide gels at room temperature and 150 or 180 V for 45 to 60 minutes (depending on the size of the DNA fragments). For the testing of effector molecules, IpsA was first incubated with the putative effector for 20 minutes at room temperature, followed by addition of the DNA, further 20 minutes incubation and electrophoresis as described above. As control AcnR protein and the acn promoter fragment was used as described in [47] (Additional file 1: Figure S7).

## Fluorescence microscopy

For recording fluorescence microscopy images, cells were centrifuged and resuspended in phosphate-buffered saline (PBS, $137 \mathrm{mM} \mathrm{NaCl}, 2.7 \mathrm{mM} \mathrm{KCl}, 4.3 \mathrm{mM} \mathrm{Na} 2 \mathrm{HPO}_{4}$, $\left.1.4 \mathrm{mM} \mathrm{KH} \mathrm{KO}_{4}, \mathrm{pH} 7.3\right)$ containing $200 \mathrm{ng} / \mathrm{mL}$ DAPI and $300 \mathrm{ng} / \mathrm{mL}$ nile red. After 5 to 10 minutes incubation at room temperature, the cells were analyzed on agar pads
using a Zeiss Axioplan 2 imaging microscope that was equipped with an AxioCam MRm camera and a PlanApochromat $100 \times$, 1.40 Oil DIC oil-immersion objective. Digital images were acquired and analyzed with AxioVision 4.6 software (Zeiss, Göttingen, Germany).

## Promoter fusion studies

To analyze the regulation of the ino1 promoter by IpsA in vivo, a DNA fragment covering the cg3323 promoter region was fused to the eyfp-coding sequence ( $\mathrm{pJC1}$ -Pcg3323-eYFP). Wild-type and $\operatorname{\Delta ips} A$ cells were transformed with the resulting plasmid and the fluorescence output was analyzed on minimal medium with $2 \%$ (w/v) glucose or myo-inositol in the Biolector system.

## Extraction and determination of mycothiol

The determination of mycothiol was carried out as described [21]. Briefly, cells were first cultivated in BHI medium for 6 h and then twice in CGXII minimal medium with $4 \%(\mathrm{w} / \mathrm{v})$ as carbon source at $30^{\circ} \mathrm{C}$ and 120 rpm for 16 h and 24 h , respectively. 200 mg of wet cells were resuspended in 1 mL of warm $\left(60^{\circ} \mathrm{C}\right)$ acetonitrile containing 20 mM Tris-HCl, pH 8.0 and 2 mM bromobimane (Sigma-Aldrich, Taufkirchen near Munich, Germany), sonificated for 20 s and further incubated in a $60^{\circ} \mathrm{C}$ water bath for 15 minutes in the dark. The samples were acidified with $5 \mu \mathrm{~L}$ of 5 M methanesulfonic acid and the cellular debris was removed by centrifugation at $16,000 g$ for 10 minutes. The supernatants were filtered and diluted fivefold in 10 mM methanesulfonic acid. A wild-type control sample was incubated with 5 mM NMM to block the thiol groups prior to derivatization with bromobimane. The sample were analyzed using a $\mathrm{C}_{18}$ column (LiChrospher RP 18, $125 \times 4 \mathrm{~mm})$ and the solvents $\mathrm{A}(100 \mathrm{mM}$ sodium acetate buffer pH 7.2 ) and B ( $100 \%$ methanol) a flow rate of $0.7 \mathrm{~mL} /$ minute and the following gradient: 0 minutes, $15 \%$ B; 1 minute, $15 \%$ B; 1.5 minutes, $35 \%$ B; 11 minutes, $80 \%$ B; 12 minutes, $95 \%$ B; 13 minutes $100 \%$ B. The peak assumed to be mycothiol ( 5.61 minutes) was identified by comparison of the wild-type chromatogram with the one from the $\Delta m s h C$ strain and the NMM treated sample. Furthermore this peak appears shortly after glutathione (5.14) as published for a similar method [48].

## Growth of bacteria for lipid and lipoglycan analysis

Overnight cultures of C. glutamicum ATCC 13032, C. glutamicum $\Delta i p s A$, C. glutamicum $\Delta i p s A::$ :pK18int-ipsA, and C. glutamicum $\Delta i p s A$ harboring ino1 (cg3323), Rv3575 and DIP1969 in plasmid pAN6 were pregrown in 20 mL of $3.7 \%$ brain-heart-glucose bouillon (SigmaAldrich, Gillingham, UK) containing Kanamycin ( $25 \mu \mathrm{~g} /$ mL ). The cells were washed once with PBS, used to inoculate a second preculture to an $\mathrm{OD}_{600}$ of 1 in 50 mL CGXII minimal medium ( 50 mL ) supplemented with $2 \%$ glucose
and incubated with gentle shaking overnight at $30^{\circ} \mathrm{C}$. The cells of the second preculture were washed once with PBS and used to inoculate the main cultures to an $\mathrm{OD}_{600}$ of 0.1 in 10 mL CGXII minimal medium ( 10 mL ), supplemented with either $2 \%$ glucose or $2 \%$ myo-inositol and grown until early $\log$ phase. At early log phase, the cells were labeled with $1 \mu \mathrm{Ci} / \mathrm{mL}\left[1,2-{ }^{14} \mathrm{C}\right]$-acetate ( 1.66 to $2.22 \mathrm{GBq} / \mathrm{mmol}$, PerkinElmer Inc., Waltham, Massachusetts, USA) and incubated overnight at $30^{\circ} \mathrm{C}$ with gentle shaking. The cells were recovered following centrifugation at $3,500 g$ and dried.

## Analysis of polar lipids and lipoglycans

The cell cultivation procedure for the following experiments is described in the Additional file 1. Polar and apolar lipid extracts were prepared from the dried cell pellets using established procedures [49]. The dried polar lipid extracts were resuspended in $\mathrm{CHCl}_{3}: \mathrm{CH}_{3} \mathrm{OH}(100 \mu \mathrm{~L}, 2: 1$, $\mathrm{v} / \mathrm{v}$ ) and the incorporation of $\left[1,2-{ }^{14} \mathrm{C}\right]$ acetate in lipid extracts determined by counting an aliquot (5\%) using scintillation fluid ( 5 mL ). To analyze the lipid profiles, equal counts of polar lipid extracts ( $20,000 \mathrm{cpm}$ ) were applied to silica thin layer chromatography (silica gel $60 \mathrm{~F}_{254}$ Merck 5554, TLC plates) and developed using the solvent system: $\mathrm{CHCl}_{3}: \mathrm{CH}_{3} \mathrm{OH}: \mathrm{H}_{2} \mathrm{O}(60: 30: 6, \mathrm{v} / \mathrm{v} / \mathrm{v})$ in the first dimension and $\mathrm{CHCl}_{3}: \mathrm{CH}_{3} \mathrm{CO}_{2} \mathrm{H}: \mathrm{CH}_{3} \mathrm{OH}: \mathrm{H}_{2} \mathrm{O}(40: 25: 3: 6, \mathrm{v} / \mathrm{v} / \mathrm{v} / \mathrm{v})$ in the second dimension. The autoradiograms were obtained by exposing the TLC results to X-ray films (Kodak-Biomax MR Kodak, Rochester, New York, USA) for 48 h .
Lipoglycans were extracted using previously described protocols [50]. Briefly, dried cell pellets from a 10 mL volume culture were resuspended in water and refluxed five times with equal volume of $50 \%$ aqueous $\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{OH}$ at $85^{\circ} \mathrm{C}$, for 6 h intervals. The suspension was centrifuged and the supernatant recovered between each extraction following centrifugation. The combined supernatant fractions were dried and subjected to hot phenol- $\mathrm{H}_{2} \mathrm{O}$ treatment at $65^{\circ} \mathrm{C}$. The aqueous phase containing the crude lipoglycan fraction was recovered and dialyzed (molecular weight cutoff 3.5 KDa ) against water. The dialyzed fraction was dried and $20 \mu \mathrm{~g}$ of lipoglycans loaded on a $15 \%$ SDS-PAGE gel. The lipoglycans were visualized using Pro-Q emerald glycoprotein stain (Life Technologies, Darmstadt, Germany).

## Additional file

## Additional file 1: Additional methods, figures and tables.

## Abbreviations

BHI: brain-heart infusion; CDP-DAG: cytidine diphosphate-diacylglycerol; DAG: diacyl-glycerol; eyfp: enhanced yellow fluorescent protein; GL-A: GlcAGroAc $2 ; 1,2$-di-O-C ${ }_{16} / \mathrm{C}_{18: 1}$-(a-D-glucopyranosyluronic acid)-( $1 \rightarrow 3$ )glycerol; GL-X: ManGlcAGroAc $2 ; 1,2$-di-O-C $16 /$ C $_{18: 1}$-(a-D-mannopyranosyl)$(1 \rightarrow 4)$-(a-D-glucopyranosyluronic acid)-(1 $\rightarrow 3$ )-glycerol; IMP: inositol monophosphatase; LAM: liporarabinomannan; LM: lipomannan; MSH: mycothiol; PBS: phosphate-buffered saline; PI: phosphatidylinositol; PIMs: PI mannosides.

## Competing interests

The authors declare that they have no competing interests.

## Authors' contributions

MB designed and coordinated the study, constructed the mutant, performed the microarray experiments, the mycothiol measurements and fluorescence microscopy and prepared the figures. KL and CG carried out the recombinant DNA work, the electrophoretic mobility shift assays (EMSAs), the protein purification, the promoter fusion studies and the growth experiments. SG and GSB performed and analyzed the cell wall lipid studies and prepared the respective parts of the manuscript. MB and JF conceived of the study and wrote the manuscript. All authors read and approved the final manuscript.

## Acknowledgements

This work was supported by the German Ministry of Education and Research (BMBF, grant 0316017B) and by the Helmholtz Association (Young Investigator grant VH-NG-716). We thank Tino Polen for help with the microarray analyses, Jan van Ooyen for help with high-performance liquid chromatography (HPLC) measurements, the students Jakob Schlegel and Sebastian Schnorrenberg for their contribution to this project, Antonia Heyer for providing plasmid pJC1-venus-term, Jörn Kalinowski for information about transcriptional start sites as well as Lothar Eggeling and Simon Klaffl for discussion.

## Author details

Institut für Bio- und Geowissenschaften, IBG-1: Biotechnologie, Forschungszentrum Jülich, 52425 Jülich, Germany. ${ }^{2}$ School of Biosciences, University of Birmingham, Edgbaston, Birmingham, B15 2TT, UK.

Received: 13 September 2013 Accepted: 17 December 2013
Published: 30 December 2013

## References

1. Mishra AK, Driessen NN, Appelmelk BJ, Besra GS: Lipoarabinomannan and related glycoconjugates: structure, biogenesis and role in Mycobacterium tuberculosis physiology and host-pathogen interaction. FEMS Microbiol Rev 2011, 35:1126-1157.
2. Eggeling L, Besra GS, Alderwick L: Structure and synthesis of the cell wall. In Corynebacteria: genomics and molecular biology. Edited by Burkovski A. Wymondham, UK: Caister Academic Press; 2008:267-294.
3. Fahey RC: Glutathione analogs in prokaryotes. Biochim Biophys Acta 1830, 2013:3182-3198.
4. Morita YS, Fukuda T, Sena CBC, Yamaryo-Botte Y, McConville MJ, Kinoshita T: Inositol lipid metabolism in mycobacteria: biosynthesis and regulatory mechanisms. Biochim Biophys Acta 1810, 2011:630-641.
5. http://www.microbesonline.org
6. Alm EJ, Huang KH, Price MN, Koche RP, Keller K, Dubchak IL, Arkin AP: The MicrobesOnline web site for comparative genomics. Genome Res 2005, 15:1015-1022.
7. Bachhawat N, Mande SC: Identification of the INO1 gene of Mycobacterium tuberculosis H37Rv reveals a novel class of inositol-1phosphate synthase enzyme. J Mol Biol 1999, 291:531-536.
8. Nigou J, Besra GS: Characterization and regulation of inositol monophosphatase activity in Mycobacterium smegmatis. Biochem J 2002, 361:385-390.
9. Movahedzadeh F, Wheeler PR, Dinadayala P, Av-Gay Y, Parish T, Daffe M, Stoker NG: Inositol monophosphate phosphatase genes of Mycobacterium tuberculosis. BMC Microbiol 2010, 10:50.
10. Movahedzadeh F, Smith DA, Norman RA, Dinadayala P, Murray-Rust J, Russell DG, Kendall SL, Rison SC, McAlister MS, Bancroft GJ, McDonald NQ, Daffe M, Av-Gay Y, Stoker NG: The Mycobacterium tuberculosis ino1 gene is essential for growth and virulence. Mol Microbiol 2004, 51:1003-1014.
11. Jackson M, Crick DC, Brennan PJ: Phosphatidylinositol is an essential phospholipid of mycobacteria. J Biol Chem 2000, 275:30092-30099.
12. Haites RE, Morita YS, McConville MJ, Jacobe HB: Function of phosphatidylinositol in mycobacteria. J Biol Chem 2005, 280:10981-10987.
13. Hernick M: Mycothiol: a target for potentiation of rifampin and other antibiotics against Mycobacterium tuberculosis. Expert Rev Anti Infect Ther 2013, 11:49-67.
14. Krings E, Krumbach K, Bathe B, Kelle R, Wendisch VF, Sahm H, Eggeling L: Characterization of myo-inositol utilization by Corynebacterium glutamicum: the stimulon, identification of transporters, and influence on L-lysine formation. J Bacteriol 2006, 188:8054-8061.
15. Klaffl S, Brocker M, Kalinowski J, Eikmanns B, Bott M: Complex regulation of the PEP carboxykinase gene pck and characterization of its GntR-type regulator IIIR as a repressor of myo-inositol utilization genes in Corynebacterium glutamicum. J Bacteriol 2013, 195:4283-4296.
16. Swint-Kruse L, Matthews KS: Allostery in the Lacl/GaIR family: variations on a theme. Curr Opin Microbiol 2009, 12:129-137
17. www.ncbi.nIm.nih.gov/geo
18. Wennerhold J, Bott M: The DtxR regulon of Corynebacterium glutamicum. J Bacteriol 2006, 188:2907-2918.
19. Wennerhold J, Krug A, Bott M: The AraC-type regulator RipA represses aconitase and other iron proteins from Corynebacterium under iron limitation and is itself repressed by DtxR. J Biol Chem 2005, 280:40500-40508.
20. Bailey TL, Elkan C: "Fitting a mixture model by expectation maximization to discover motifs in biopolymers", Proceedings of the Second International Conference on Intelligent Systems for Molecular Biology. Menlo Park, Ca: AAAI Press; 1994:28-36.
21. Feng J, Che YS, Milse J, Yin YJ, Liu L, Rückert C, Shen XH, Qi SW, Kalinowski J, Liu SJ: The gene ncgl2918 encodes a novel maleylpyruvate isomerase that needs mycothiol as cofactor and links mycothiol biosynthesis and gentisate assimilation in Corynebacterium glutamicum. J Biol Chem 2006, 281:10778-10785.
22. Tatituri RV, Illarionov PA, Dover LG, Nigou J, Gilleron M, Hitchen P Krumbach K, Morris HR, Spencer N, Dell A, Eggeling L, Besra GS: Inactivation of Corynebacterium glutamicum NCgl0452 and the role of MgtA in the biosynthesis of a novel mannosylated glycolipid involved in lipomannan biosynthesis. J Biol Chem 2007, 282:4561-4572.
23. Mishra AK, Klein C, Gurcha SS, Alderwick LJ, Babu P, Hitchen PG, Morris HR, Dell A, Besra GS, Eggeling L: Structural characterization and functional properties of a novel lipomannan variant isolated from a Corynebacterium glutamicum pim ${ }^{\prime}$ mutant. Antonie Van Leeuwenhoek 2008, 94:277-287.
24. Mishra AK, Batt S, Krumbach K, Eggeling L, Besra GS: Characterization of the Corynebacterium glutamicum $\Delta$ pimB $^{\prime} \Delta m g t A$ double deletion mutant and the role of Mycobacterium tuberculosis orthologues Rv2188c and Rv0557 in glycolipid biosynthesis. J Bacteriol 2009, 191:4465-4472.
25. Vilchèze C, Av-Gay Y, Attarian R, Liu Z, Hazbón MH, Colangeli R, Chen B, Liu W, Alland D, Sacchettini JC, Jacobs WR Jr: Mycothiol biosynthesis is essential for ethionamide susceptibility in Mycobacterium tuberculosis. Mol Microbiol 2008, 69:1316-1329.
26. Turner BL, Paphazy MJ, Haygarth PM, McKelvie ID: Inositol phosphates in the environment. Philos T Roy Soc B 2002, 357:449-469.
27. Schröder J, Jochmann N, Rodionov DA, Tauch A: The Zur regulon of Corynebacterium glutamicum ATCC 13032. BMC Genomics 2010, 11:12.
28. Tropis M, Meniche X, Wolf A, Gebhardt H, Strelkov S, Chami M, Schomburg D, Kramer R, Morbach S, Daffe M: The crucial role of trehalose and structurally related oligosaccharides in the biosynthesis and transfer of mycolic acids in corynebacterineae. J Biol Chem 2005, 280:26573-26585.
29. Murphy HN, Stewart GR, Mischenko W, Apt AS, Harris R, McAlister MSB, Driscoll PC, Young DB, Robertson BD: The OtsAB pathway is essential for trehalose biosynthesis in Mycobacterium tuberculosis. J Biol Chem 2005, 280:14524-14529.
30. Norman RA, McAlister MSB, Murray-Rust J, Movahedzadeh F, Stoker NG, McDonald NQ: Crystal structure of inositol 1-phosphate synthase from Mycobacterium tuberculosis, a key enzyme in phosphatidylinositol synthesis. Structure 2002, 10:393-402.
31. Roback P, Beard J, Baumann D, Gille C, Henry K, Krohn S, Wiste H, Voskuil MI, Rainville C, Rutherford R: A predicted operon map for Mycobacterium tuberculosis. Nucleic Acids Res 2007, 35:5085-5095.
32. Bell CE, Lewis M: The Lac repressor: a second generation of structural and functional studies. Curr Opin Struct Biol 2001, 11:19-25.
33. Camas FM, Alm EJ, Poyatos JF: Local gene regulation details a recognition code within the Lacl transcriptional factor family. PLoS Comput Biol 2010, 6:11.
34. Belisle JT, Vissa VD, Sievert T, Takayama K, Brennan PJ, Besra GS: Role of the major antigen of Mycobacterium tuberculosis in cell wall biogenesis. Science 1997, 276:1420-1422
35. Keilhauer C, Eggeling L, Sahm H: Isoleucine synthesis in Corynebacterium glutamicum - Molecular analysis of the Ilvb-Ilvn-Ilvc Operon. J Bacteriol 1993, 175:5595-5603.
36. Sambrook J, Russell DW: Molecular cloning: A Laboratory Manual. 3rd edition. Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press; 2001.
37. Hanahan D: Studies on transformation of Escherichia coli with plasmids. J Mol Biol 1983, 166:557-580.
38. van der Rest ME, Lange C, Molenaar D: A heat shock following electroporation induces highly efficient transformation of Corynebacterium glutamicum with xenogeneic plasmid DNA. Appl Microbiol Biotechnol 1999, 52:541-545.
39. Niebisch A, Bott M: Molecular analysis of the cytochrome $b c_{1}-a a_{3}$ branch of the Corynebacterium glutamicum respiratory chain containing an unusual diheme cytochrome $c_{1}$. Arch Microbiol 2001, 175:282-294.
40. Möker N, Brocker M, Schaffer S, Krämer R, Morbach S, Bott M: Deletion of the genes encoding the MtrA-MtrB two-component system of Corynebacterium glutamicum has a strong influence on cell morphology, antibiotics susceptibility and expression of genes involved in osmoprotection. Mol Microbiol 2004, 54:420-438.
41. Lange C, Rittmann D, Wendisch VF, Bott M, Sahm H: Global expression profiling and physiological characterization of Corynebacterium glutamicum grown in the presence of L-valine. Appl Environ Microbiol 2003, 69:2521-2532.
42. Smyth GK: Limma: linear models for microarray data. In Bioinformatics and Computational Biology Solutions Using R and Bioconductor. Edited by Gentleman R, Carey VJ, Huber W, Irizarry RA, Dudoit S. New York, NY: Springer; 2005:397-420.
43. Edgar R, Domrachev M, Lash AE: Gene Expression Omnibus: NCBI gene expression and hybridization array data repository. Nucleic Acids Res 2002, 30(1):207-210.
44. Garcia-Nafria J, Baumgart M, Bott M, Wilkinson AJ, Wilson KS: The Corynebacterium glutamicum aconitase repressor: scratching around for crystals. Acta Crystallogr Sect F Struct Biol Cryst Commun 2010, 66:1074-1077.
45. Niebisch A, Kabus A, Schultz C, Weil B, Bott M: Corynebacterial protein kinase $G$ controls 2-oxoglutarate dehydrogenase activity via the phosphorylation status of the Odhl protein. J Biol Chem 2006, 281:12300-12307.
46. Heyer A, Gätgens C, Hentschel E, Kalinowski J, Bott M, Frunzke J: The two-component system ChrSA is crucial for haem tolerance and interferes with HrrSA in haem-dependent gene regulation in Corynebacterium glutamicum. Microbiology 2012, 158:3020-3031.
47. Garcia-Nafria J, Baumgart M, Turkenburg JP, Wilkinson AJ, Bott M, Wilson KS: Crystal and solution studies reveal that the transcriptional regulator AcnR of Corynebacterium glutamicum is regulated by Citrate- $\mathrm{Mg}^{2+}$ binding to a non-canonical pocket. J Biol Chem 2013, 288:15800-15812.
48. Anderberg SJ, Newton GL, Fahey RC: Mycothiol biosynthesis and metabolism - Cellular levels of potential intermediates in the biosynthesis and degradation of mycothiol in Mycobacterium smegmatis. J Biol Chem 1998, 273:30391-30397.
49. Dobson G, Minnikin DE, Minnikin SM, Parlett JH, Goodfellow M, Ridell M, Magnusson M: Systematic analysis of complex mycobacterial lipids. In Chemical Methods in Bacterial Systematics, Volume 1. Edited by M. G, E. MD. London, UK: Academic Press; 1985:237-265.
50. Ludwiczak P, Brando T, Monsarrat B, Puzo G: Structural characterization of Mycobacterium tuberculosis lipoarabinomannans by the combination of capillary electrophoresis and matrix-assisted laser desorption/ionization time of-flight mass spectrometry. Anal Chem 2001, 73:2323-2330.
doi:10.1186/1741-7007-11-122
Cite this article as: Baumgart et al.: IpsA, a novel Lacl-type regulator, is required for inositol-derived lipid formation in Corynebacteria and Mycobacteria. BMC Biology 2013 11:122.

[^0]:    * Correspondence: j.frunzke@fz-juelich.de
    ${ }^{1}$ Institut für Bio- und Geowissenschaften, IBG-1: Biotechnologie, Forschungszentrum Jülich, 52425 Jülich, Germany
    Full list of author information is available at the end of the article

