Conditional expression is a powerful tool to investigate the role of bacterial genes. Here, we adapt the *Pseudomonas putida*-derived positively regulated XylS/Pm expression system to control inducible gene expression in *Mycobacterium smegmatis* and *Mycobacterium tuberculosis*, the causative agent of human tuberculosis. By making simple changes to a Gram-negative broad-host-range XylS/Pm-regulated gene expression vector, we prove that it is possible to adapt this well-studied expression system to non-Gram-negative species. With the benzoic acid-derived inducer m-toluate, we achieve a robust, time- and dose-dependent reversible induction of Pm-mediated expression in mycobacteria, with low background expression levels. XylS/Pm is thus an important addition to existing mycobacterial expression tools, especially when low basal expression is of particular importance.

Introduction

Tuberculosis (TB), caused by *Mycobacterium tuberculosis* (*Mtb*), continues to be one of the most serious lethal infectious diseases in the world. The World Health Organization reports around 9 million new cases and 1.5 million deaths annually as a result of TB [1]. The spread of drug-resistant mycobacteria adds to the importance of developing new anti-mycobacterial therapies [2]. To improve TB therapeutics, it is crucial to better understand the underlying molecular mechanisms of the disease, and to gain new insights it is important to create a greater selection of new and enhanced molecular tools.

Inducible expression systems have become fundamental engines for genetic research. For instance, over-expression or conditional regulation of genes enables us to map gene function, determine gene essentiality and validate drug targets [3–5]. Regulators of inducible expression systems can be divided into three groups; activators (positive regulation), repressors (negative regulation) and regulators with dual activity (positive and negative regulation) [6]. Upon induction, activators commonly lead to activation of transcription by interacting with the RNA...
polymerase and thereby increase the affinity between the polymerase and the promoter. Conversely, a repressor will typically bind to the operator sequence in the absence of inducer and inhibit transcription initiation, for instance by preventing the binding of RNA polymerase to the promoter. Thus, the steady state level of the repressor in the various bacterial growth phases is important to keep basal expression low. In the presence of inducer the repressor undergoes conformational changes, making it unable to bind the operator and thus allow transcription [6, 7]. Regulators with dual activity can act as repressors or activators of the same promoter, depending on presence or absence of inducer [8]. While the majority of inducible expression systems are negatively regulated, it has been proposed that positively regulated systems, independent of repressor activity, are better candidates when tight control with low basal expression levels is desired [9, 10].

In addition to low basal expression, a good mycobacterial inducible expression system should provide robust and reversible induction by an inexpensive, non-toxic inducer which penetrates cells by diffusion, induce a dose- and time-dependent response and have the potential to work across mycobacterial species as well as in macrophage and animal infection models.

While the lack of a diverse collection of mycobacterial inducible expression tools has hampered research for decades, widely used inducible expression systems were developed for *Escherichia coli* by the early 1980s [11, 12]. The first equivalent system in mycobacteria, the acetamide-inducible system based on the *Mycobacterium smegmatis* (*Msmeg*) endogenous acetamidase promoter [13], was developed as late as in 1997. A few years later, a temperature-inducible expression cassette, based on the *Ptra* promoter and a derivative of the TraR repressor, was demonstrated to be functional but weak in mycobacteria [14]. More recently, several tetracycline (tet) repressor-based inducible systems have been developed [15–18], together with pristinamycin, nitril, arabinose and IPTG-inducible expression tools [19–22]. The most recent contribution to mycobacterial inducible expression is a riboswitch-based theophylline responsive system regulated at a translational level [23]. The aforementioned systems have various strengths and weaknesses as expression tools; all are however dependent on repression to lower expression in the uninduced state. The nitril-inducible NitR/PnitA system adapted from the related Gram-positive *Rhodococcus rhodochrous* is controlled by the AraC-family regulator NitR proposed to work as both a repressor and activator, albeit the dual function of NitR has not yet been experimentally verified [19, 24]. Thus, developing a positively regulated system independent of repression of the inducible promoter could greatly benefit mycobacterial research.

*XylS/Pm* is a well-studied, positively regulated expression system derived from the *Pseudomonas putida* TOL plasmid pWWO [25], and it is known to function in a wide range of Gram-negative bacterial species [26]. The inducible *Pm* promoter originates from the transcriptional regulation of the meta-cleavage pathway for benzoate and alkylbenzoate catabolism controlled by the activator XylS and is to our knowledge fully independent of transcriptional repression in basal expression mode [6]. Upon induction by benzoate-derived effectors, such as *m*-toluate, XylS dimerizes and binds to the *Pm* promoter where it interacts with the transcription initiation machinery and activates transcription [27, 28].

In this study we adapt XylS/Pm for use in *Mtb* and *Msmeg*, introducing the first positively regulated expression system to mycobacteria. We show robust time- and dose-dependent inducible expression from the *Pm* promoter, with a basal expression comparably lower to commonly used mycobacterial expression tools. Importantly, by showing the function of XylS/Pm-controlled expression in non-Gram-negative species, we demonstrate the great potential of expanding the use of this expression system to other Gram-positive bacteria.
## Materials and Methods

### Bacterial strains and culture conditions

*Msmeg* mc²155 [29], *Msmeg* DM22 [30] (kindly provided by Kurt Krause, University of Otago, New Zealand) and *Mtb* H37Rv (ATCC 25618) were cultured in Middlebrook 7H9 (BD Difco) supplemented with 0.2% glycerol, 0.05% Tween80 and 10% ADC (*Msmeg*: 50 g BSA fraction V, 20 g dextrose, 8.5 g NaCl, 0.03 g catalase, dH₂O up to 1 L) or OADC (*Mtb*: BD Difco). Kanamycin was added to 20 μg/ml in mycobacteria and to 50 μg/ml in *E. coli*. Hygromycin was added to 50 μg/ml in mycobacteria. Colonies of *Msmeg* were grown on LB agar plates, whereas colonies of *Mtb* were grown on 7H10 (BD Difco) supplemented with glycerol and 10% OADC (BD Difco). Standard cloning and mycobacterial transformation techniques were used.

### Determination of *Mtb* growth in *m*-toluate

Strain H37Rv was grown to mid log phase in standard 7H9 media. The culture was then diluted to an OD₆₀₀ of 0.01 and aliquoted in a 96-well plate with the final concentrations of *m*-toluate as indicated in the respective figure. Ethanol carrier concentration was equal in all conditions. Plates were incubated at 37°C for 14 days. Cultures were then resuspended well, and OD₆₀₀ was determined using a Spectramax 5 (Molecular Devices).

### Expression vector construction

To create the expression plasmid pMDX, a DNA fragment containing a) *xylS* controlled by *Pmyc1tetO* promoter [16] (called *Ptet* in this study), b) reverse TetR#28 [31] (called reverse TetR in this study) controlled by *Psmyc* promoter [16], c) *Pm* promoter and d) transcriptional terminators strategically placed was synthesized by Genscript USA Inc (NJ, USA). The *xylS* and *Pm* sequences were obtained from pKT1 [32]. The synthetic expression cassette described above (GenBank accession number KT239651) was cloned into the backbone of pMV261 [33] using NotI. pMDX-luc was created by PCR amplification of the firefly luciferase gene from pMH109 (kindly provided by Mark Hickey/David Sherman, Seattle Biomed, Seattle, WA, USA) [34, 35], using primers LucMDX_F and LucMDX_R. pMDX-zeo was created by PCR amplification of the zeocin resistance gene *Sh ble* (GenBank accession number ABW35374.1) from pER10 (unpublished data) using primers ZeoMDX_F and ZeoMDX_R. The amplified genes were separately cloned into pMDX using the EcoRI and NdeI sites, placing the reporter genes under control of the *Pm* promoter. pTET-zeo was created by PCR amplification of the zeocin gene from pMDX-zeo using the primer pair ZeoPacIF and ZeoPstIR. The amplified gene was cloned into the PacI and PstI sites of pUV15TetORm [16]. pMDX-hyg-alr was constructed by amplifying the hygromycin resistance cassette from *Msmeg* ΔfxbA strain [36] using primers pMDXHyG-F-NheI and pMDXHyG-R-SpeI and clone it into NheI and SpeI restriction sites of pMDX, before the complete alanine racemase (*alr*) gene was amplified from *Msmeg* mc²155 [29] with primers pMDXAlr-F-NdeI and pMDXAlr-R- EcoRI and cloned into NdeI and EcoRI restriction sites of pMDX. The pTET-αlr was constructed by amplifying the alanine racemase gene from *Msmeg* mc²155 [29] using primers pTetOA1r-R-PacI and pTetOA1r-R-SbfI and clone it into PacI and SbfI restriction sites of pTET-zeo. pMDXint and pMDXint-luc was created by amplifying integrase and mycobacteriophage L5 attP-site from pMC1s [16] using primers pMC1_F and pMC1_R. The generated PCR product was digested with PciI and EcoRI and cloned into the PciI and EcoRI sites of pMDX and pMDX-luc replacing pAL5000, creating pMDXint and pMDXint-luc, respectively. *pUV15TetORm* and *pMC1s* were kindly provided by Sabine Ehr/Dirk Schnappinger, Weill Cornell Medical College, New York, USA. Details of the plasmids and primer sequences can be found in Table 1 and Table 2, respectively.
**XylS/Pm induction protocol**

*Msmeg* or *Mtb* transformed with either pMDX (negative control), pMDX-luc, pMDX-zeo, pMDXint, pMDXint-luc or pMH109 (positive control for luciferase assays) were grown in 7H9 with supplements and 20 μg/ml kanamycin to stationary phase, then diluted approximately 1:500 and grown to OD₆₀₀ 0.05 – 0.1 at 37°C. At this point, the samples were normalized to identical optical densities (ODs) and induced with the indicated amount of *m*-toluate (Sigma, 1 M stock solution solved in laboratory grade ethanol). For uninduced controls the same

### Table 1. Plasmids used in this study.

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Properties relevant to study</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>pMDX</td>
<td>Kan⁺, XylS controlled by <em>Ptet</em>, reverse TetR controlled by <em>Psmyc, Pm, oriM</em> (pAL5000), oriE (pBR322).</td>
<td>This study</td>
</tr>
<tr>
<td>pMDX-luc</td>
<td>pMDX derivative. Firefly luciferase gene from pMH109 cloned under control of Pm.</td>
<td>This study</td>
</tr>
<tr>
<td>pMDX-zeo</td>
<td>pMDX derivative. <em>Sh ble</em> gene cloned under control of Pm.</td>
<td>This study</td>
</tr>
<tr>
<td>pMDXint</td>
<td>Kan⁺, XylS under control of <em>Ptet</em>, reverse TetR controlled by <em>Psmyc, Pm</em>, integrase, attP-site</td>
<td>This study</td>
</tr>
<tr>
<td>pMDXint-luc</td>
<td>pMDXint derivative, <em>Pm</em> controlling firefly luciferase gene from pMH109, integrase, attP-site.</td>
<td>This study</td>
</tr>
<tr>
<td>pMC1s</td>
<td>Integrate, attP-site</td>
<td>[16]</td>
</tr>
<tr>
<td>pNBV1</td>
<td>Hyg⁺, <em>E. coli</em>—mycobacterial shuttle vector</td>
<td>[37]</td>
</tr>
<tr>
<td>pKT1</td>
<td>Kan⁺, XylS controlled by <em>Ps2</em>, firefly luciferase gene from <em>Photinus pyralis</em> controlled by <em>Pm, oriE</em> (RK2).</td>
<td>[32]</td>
</tr>
<tr>
<td>pMH109</td>
<td>Hyg⁺, attP-site, firefly luciferase gene controlled by MOP.</td>
<td>[34, 35]</td>
</tr>
<tr>
<td>pER10</td>
<td>Zeo⁺ (<em>Sh ble</em> gene: GenBank accession number ABW35374.1)</td>
<td>unpublished data</td>
</tr>
<tr>
<td>pMV261</td>
<td>Kan⁺, <em>oriM</em> (pAL5000), <em>oriE</em> (pBR322).</td>
<td>[33]</td>
</tr>
<tr>
<td>pTET-zeo</td>
<td>Hyg⁺, P₂myc*tetO in control of <em>Sh ble</em>. TetR.</td>
<td>This study</td>
</tr>
<tr>
<td>pUV15TetORm</td>
<td>Hyg⁺, Kan⁺, P₂myc*tetO in control of GFP. Tn10 derived TetR controlled by <em>Psmyc, oriM</em> (pAL5000), oriE (pBR322).</td>
<td>[16]</td>
</tr>
<tr>
<td>pUV15TetORm:: luciferase</td>
<td>Derivative of pUV15TetORm. <em>Pmyc</em>tetO in control of luciferase.</td>
<td>[38]</td>
</tr>
<tr>
<td>pMDX-hyg-air</td>
<td>pMDX derivative. <em>air</em> gene cloned under control of <em>Pm</em>. Hyg⁺ added.</td>
<td>This study</td>
</tr>
<tr>
<td>pTET-air</td>
<td>pTET-zeo derivative. <em>Pmyc</em>tetO in control of <em>air</em>.</td>
<td>This study</td>
</tr>
</tbody>
</table>


doi:10.1371/journal.pone.0134544.t001

### Table 2. Oligonucleotides used in this study.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence, 5'-3'</th>
</tr>
</thead>
<tbody>
<tr>
<td>LucMDX_F</td>
<td>AGTCATGAAACATATGGAGACCAGCCAAAAACAT</td>
</tr>
<tr>
<td>LucMDX_R</td>
<td>ATCGAATTCTCTAGAATTACACGGCGATCTTTCC</td>
</tr>
<tr>
<td>ZeoMDX_F</td>
<td>AGTCATGAAACATATGGCCAAGTGGACCAGTGC</td>
</tr>
<tr>
<td>ZeoMDX_R</td>
<td>ATCGAATTCTCTAGAATCGTCTCGTCTCCGCCAGCA</td>
</tr>
<tr>
<td>ZeoPacIF</td>
<td>CGCATGCTTAATTTGAGGAGGAGATACATATGGCCAGTTGACCAGTGC</td>
</tr>
<tr>
<td>ZeoPstIR</td>
<td>CCTCTAATCTGAGCCGGGGAGGCCATTTTAAATGCTGAGATCTAGTCTCGTCTCCGCCCA</td>
</tr>
<tr>
<td>pMC1_F</td>
<td>TTGTGCTGATACTATGGAAGCCAGCAGCTCTCGT</td>
</tr>
<tr>
<td>pMC1_R</td>
<td>GGATCCAGCTCAGAATTTC</td>
</tr>
<tr>
<td>pMDXHyG-F-Nhel</td>
<td>CTAGCTAGCTGCTACCTTCTACATATAAGGAGAAGAGAGGAGATATACATATGGCCAGTTGACCAGTGC</td>
</tr>
<tr>
<td>pMDXHyG-R-Spel</td>
<td>CACATAGTGGGCTTTAGCTGTAATTAA</td>
</tr>
<tr>
<td>pMDXAir-F- Ndel</td>
<td>CCATAGTGGACCACCCAGCGACCCATG</td>
</tr>
<tr>
<td>pMDXAir-R- EcoRI</td>
<td>GGAATTCTCAATCTTGTTGTCTGCTGC</td>
</tr>
<tr>
<td>pTetOAir-F-Pac I</td>
<td>CCTTAATTAATGACACCCAGGACGCC</td>
</tr>
<tr>
<td>pTetOAir-R-Sbf I</td>
<td>ATTTGTTTCCTGAGGCTCAATCTTGTTGTCTGCC</td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pone.0134544.t002
amount of ethanol was added in place of inducer. The samples were incubated with shaking at 30°C or 37°C for the number of hours specified.

**Mtb tetracycline-inducible luciferase comparison**

*Mtb* H37Rv containing pMDX-luc or pUV15tetORm::luciferase (a gift of Deborah Hung, The Broad Institute [38]) was grown to mid log phase, and then diluted to an OD$_{600}$ of 0.01. For pMDX-luc strains, m-toluate or an equivalent amount of ethanol carrier was added to each roller bottle. For pUV15tetORm::luciferase strains, anhydrous tetracycline or an equivalent amount of water carrier was added to each roller bottle. Cultures were grown with rolling at 37°C. At the indicated time points, aliquots were taken for luciferase assays as described.

**Luciferase reporter gene assays**

After induction, OD$_{600}$ was measured and noted. Cells were spun down at 4000 rpm for 10 minutes and pellets completely dried for any excess media before being resuspended in 100 μl DH$_2$O. 30 μl of the resuspended cells were mixed with 7 μl 5x Passive Lysis Buffer (Promega) in the appropriate 96-well plate and incubated with shaking for 30 minutes at 37°C for sufficient lysis. After incubation, assay substrate (Promega, E1500) was prepared according to manufacturer’s instructions and 75 μl added to each well containing cell lysis mixture. Bioluminescence was subsequently measured according to manufacturer’s recommendations on a Fluoroskan Ascent FL (Labsystems). The relative luminescence units (RLU) were recorded and normalized with regard to OD$_{600}$ or colony forming units (CFU). To measure the amount of luciferase produced, cell pellets from uninduced or induced cultures were resuspended in 1 ml Passive Lysis Buffer and the cells were disrupted using 0.1 mm glass beads and an MP FastPrep-24 (3 x 45 sec, 6000 M/S). The samples were normalized by OD before cell disruption. 50 μl cell lysate was combined with 50 μl luciferase assay substrate and luciferase activity was measured as above. The amount of luciferase produced was determined by comparison to known luciferase concentrations (luciferase obtained from Abcam) and normalized to total protein levels as measured by BioRad Protein Assay.

**Zeo** (Sh ble) gene assay

On solid media: after 5 hours pre-induction with 1.5 mM m-toluate (induced) or ethanol (uninduced), *Msmeg* cultures containing were normalized by OD measurements and serially diluted. The dilutions were spotted on LB agar plates with 20 μg/ml kanamycin and increasing amounts of zeocin (Invitrogen). Induced samples were spotted on plates containing 1.5 mM m-toluate, and uninduced samples were spotted on plates containing ethanol instead of m-toluate. The plates were incubated at 30°C for 2 days.

In liquid media: after 5 hours pre-induction with 1.5 mM m-toluate, induced and uninduced *Msmeg* cultures were normalized to OD$_{600}$ 0.005 and grown in the presence of increasing concentrations of zeocin and 1.5 mM m-toluate (induced) or ethanol (uninduced) in special micro-plate “honeycomb” wells (Oy Growth Curves Ab Ltd). Growth was monitored by Bioscreen (Oy Growth Curves Ab Ltd), registering OD$_{600}$ every 2 hours. For the comparison between pMDX-zeo and pTET-zeo, the cultures were diluted directly to OD$_{600}$ 0.005, without pre-induction. The pMDX-zeo strain was grown in the presence or absence of 1.5 mM m-toluate, while the pTET-zeo strain was grown in the presence or absence of 200 ng/ml anhydro-tetracycline (atc, Sigma).
Functional adaptation of XylS/Pm expression system for regulation of gene expression in mycobacteria

We wanted to investigate whether the Gram-negative, positive regulator system XylS/Pm could be functional in mycobacteria. We determined that the most direct way to evaluate the induction potential of this system would be by simply cloning the entire Gram-negative broad host range expression cassette into a mycobacteria-E. coli shuttle vector, introducing it into mycobacteria and assessing for inducibility. Hence, we took the XylS/Pm cassette from pKT1 [32], cloning XylS with its native constitutive promoter and the reporter gene encoding firefly luciferase expressed from the inducible Pm promoter, into the shuttle vector pNBV1 [37]. However, no induction of luciferase expression was observed in Msmeg, despite optimization of induction time, inducer concentration and growth phase. We could, on the other hand, observe low basal expression from Pm independent of presence or absence of inducer (results not shown). Clearly, several factors could explain the lack of induction; the inducer might be unable to diffuse through the complex mycobacterial cell wall, XylS might not be properly expressed, the RNA polymerase might not interact with XylS or the Pm promoter and/or the chosen reporter gene might not be properly transcribed or translated in mycobacteria. Finally, the reporter gene product could be toxic to the new host cell when over-produced. To unravel potential obstacles, we tested whether the inducer could enter the cell, hypothesizing that increased concentrations of inducer would be toxic if successfully entering the bacteria. Msmeg was grown in the presence of increasing amounts of m-toluate, and indeed, we saw increasing toxicity proportional to increasing inducer concentration (S1 Fig indicating that the inducer enters the cells.

We then changed the native promoter driving expression of xylS to a mycobacterial promoter, ensuring proper constitutive expression of the activator. We chose the Pmyc,tetO promoter (called Ptet in our study) from Ehrt et al.’s adapted tet-inducible system, based on its constitutive function in the absence of the TetR repressor [16]. Excess of XylS has previously been shown to stimulate expression from Pm in the absence of inducer [28, 39, 40]. To overcome the potential increase in background expression due to over-production of XylS, we introduced a mechanism for turning off xylS transcription by adding a mutant version of TetR to our construct. In contrast to standard TetR, the mutant acts as a reverse repressor, binding rather than releasing the operators in Ptet upon tetracycline induction (TetR#28 from [31]). In the event of high basal expression due to excessive XylS, we could tune down XylS production by adding anhydro-tetracycline (atc) to the media, facilitating tighter regulatory control and lower levels of basal expression.

To address the possibility of luciferase reporter toxicity to our cells, the firefly luciferase reporter gene was changed from wild type to a modified version of the firefly luciferase previously shown to be functional in mycobacteria [34, 35]. As presented in Fig 1, the adapted XylS/Pm expression system is in an OFF mode at baseline (no induced transcription from Pm) in the absence of the inducer m-toluate (Fig 1, middle panel). However, to potentially further minimize background expression, atc can be added to repress xylS transcription, leaving the system in a more fully OFF mode (Fig 1, lower panel). Transcription from Pm is fully ON in the presence of m-toluate and in the absence of tetracycline (Fig 1, upper panel).

Induction of Pm with m-toluate is robust, time- and dose-dependent in mycobacteria

Following modification of the XylS/Pm expression cassette, we examined whether gene expression could, in fact, be induced in mycobacteria by addition of m-toluate. We introduced our
adapted expression system with luciferase as a reporter gene (pMDX-luc) to \textit{Msmeg}, and by optimizing inducer concentration and induction time we were able to achieve robust induction. When adding 1.5 mM \textit{m}-toluate, we reached up to 90-fold induction (ratio of induced to uninduced expression levels) during late log phase (Fig 2A and 2B). Furthermore, we demonstrated that induction was time-dependent, peaking after 24 hours, and dose-dependent, rising nearly proportionally with increased inducer concentrations. In contrast, uninduced background levels of luciferase did not increase over time (Fig 2A and 2C). Normalizing luciferase expression to CFU instead of OD$_{600}$ gave an enhanced induction pattern after 24 hours; we reached 240-fold induction by CFU normalization (S2 Fig).

The amount of luciferase expressed was determined by measuring relative luminescence units, RLUs, which are relative. To get a sense of expression from \textit{Pm} relative to other standard mycobacterial promoters, we compared our induced levels of luciferase with the stably expressed levels from the strong constitutive mycobacterial optimal promoter, MOP [35]. We transformed \textit{Msmeg} with pMH109, an integrative plasmid containing our luciferase gene under control of MOP [35]. As seen in Fig 2D, when \textit{Pm} was induced with 2 mM \textit{m}-toluate, luciferase expression exceeded the level expressed from MOP, indicating that a high level of expression can be achieved using XylS/\textit{Pm}. Furthermore, to approximate the amount of luciferase produced, sample luciferase activity was compared to the activity of known luciferase concentrations. Fig 2E and 2F show the level of luciferase produced for uninduced and induced samples at mid log and stationary phase, respectively. When normalized to the total protein...
concentration of the lysed cell samples, the amount of luciferase constituted approximately 0.022% (0.1μg/ml luciferase) and 0.003% (0.012μg/ml luciferase) of total protein in uninduced samples in mid log and stationary phase, respectively. Induced samples contained 0.11% (mid log, 0.64μg/ml) and 0.096% (stationary, 0.38μg/ml) luciferase.

Our pMDX-luc plasmid is episomal (pAL5000 ori) with an estimated copy number of 3–10 in mycobacteria [41]. To determine whether we could also induce expression from a chromosomally integrated single copy of the XylS/Pm expression cassette, we constructed pMDXint-luc, a vector integrating at the mycobacteriophage L5 attachment site. Again using luciferase as the reporter gene, we induced pMDXint-luc-transformed Msmeg alongside our Msmeg pMDX-luc strain, demonstrating inducible expression (~10-fold induction) even when the expression system was chromosomally integrated (S3 Fig).

Fig 2. Induction of Pm with m-toluate is robust, time- and dose-dependent. Msmeg transformed with the expression vectors pMDX-luc or the empty vector pMDX (no reporter gene) treated with increasing concentrations of m-toluate (induced) or ethanol carrier (uninduced). Cells were incubated at 30°C, and luciferase expression was determined at 2.5, 5.5, 11, 23, 31 and 49 hours after addition of m-toluate. (A) Fold induction of RLU in induced samples compared to uninduced samples. (B) Growth of uninduced and induced samples of pMDX-luc corresponding to samples in (A). (C) Time course of luciferase induction from pMDX-luc with 1.5 mM m-toluate or ethanol carrier. (D) Maximal induction of pH109 and pMDX-luc-transformed Msmeg induced with 2 mM m-toluate. (E) Amount of luciferase produced as determined by the activity of known luciferase concentrations (0, 0.1, 0.2, 0.4, 0.6, 0.8 and 1 μg/ml luciferase) in mid log phase or (F) stationary phase. Luciferase fraction of total bacterial protein shown in brackets. RLUs were normalized to the OD600 of the samples before luciferase assay. All results are representative of two or more independent experiments.

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was induced in a similar manner as described for Msmeg. As shown in Fig 3A, the Pm promoter was clearly inducible in Mtb, and showed a time-dependent response peaking in late log phase. Maximum induced expression was overall lower in Mtb than in Msmeg (Fig 3A and 3B). Mtb showed a similar tolerance pattern towards m-toluate as Msmeg (S1 Fig). Taken together,

Fig 3. Induction of Pm in Mtb. Mtb transformed with the expression vector pMDX-luc or the empty vector pMDX were induced with indicated concentrations of m-toluate (induced), or ethanol carrier (uninduced). Samples were maintained rolling at 37°C and analyzed for luciferase expression levels at 2, 3, 4 and 6 days after induction. (A) Fold induction of RLU of induced to uninduced samples normalized for OD600. (B) Growth of uninduced and induced samples containing pMDX-luc corresponding to samples in (A). The results are representative for two independent experiments.

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our results show that we are able to achieve a robust, time- and dose-dependent induction of the Pm-promoter in mycobacteria.

**Addition of anhydro-tetracycline restricts Pm-mediated induction through inhibition of xylS expression**

Positively regulated expression systems are proposed to have lower basal transcription rates than negatively regulated systems [9, 10]. We wanted to investigate whether repressing XylS production would decrease the basal expression from Pm even further. First, to confirm that adding anhydro-tetracycline (atc) to the media would activate the reverse TetR to bind the operator and reduce transcription of the xylS gene, we incubated our *Msmeg* pMDX-luc strain with or without 100 ng/ml atc over night, thus giving TetR time to bind and inhibit *Ptet*-controlled *xylS*-transcription. Aside from addition of atc, the samples were treated identically, and induced to maximum expression of luciferase (Fig 4A). We could clearly see that the addition of atc markedly reduced expression of luciferase from the Pm promoter, most likely due to the lack of activator in the cells. However, inhibiting xylS expression by atc did not seem to further lower the basal expression from Pm (Fig 4A), suggesting that background expression is likely to be mediated by *Msmeg* endogenous transcription machinery rather than excessive XylS production.

**Induced expression from Pm can be reversed by inducer washout**

Mapping gene function and essentiality are important aspects of mycobacterial research. To investigate phenotypes arising from very low expression levels of the genes of interest, being able to reverse induced expression is crucial. To investigate whether removing the inducer by washout could reduce expression from Pm in mycobacteria, we induced our *Msmeg* pMDX-luc strain to maximum luciferase expression (about 90-fold, results not shown) before washing the cells in inducer-free media upon subcultivation or directly subcultivation without washing. When the subcultures reached late log phase, we measured luciferase activity (Fig 4B). After...
subcultivation, only the unwashed sample showed induction (about 40 fold), demonstrating that inducer washout before re-cultivation is enough to completely return the cells to an uninduced state.

We furthermore investigated whether a combination of inducer washout and addition of atc to turn off xylS expression would lower the basal Pm expression. We added 100 ng/ml atc to washed cells upon subcultivation, and allowed them grow to late log phase in the presence of atc before measuring luciferase. Compared with wash alone, no further decrease in background expression levels was observed in the presence of atc (Fig 4B). Together, our results demonstrate that we could successfully revert to uninduced levels of expression by inducer washout prior to subcultivation.

Regulation of the zeocin resistance gene (Sh ble) by XylS/Pm

After demonstrating that we could induce gene expression via Pm in a precisely controlled fashion, we wanted to investigate whether Pm could be used for conditional regulation of biologically active genes in mycobacteria. Zeocin is an antibiotic of the bleomycin/phleomycin family that causes cell death by cleaving DNA. Resistance to zeocin is conferred by the Sh ble gene encoding a small acidic protein, which inhibits the antibiotic by binding to it in a stoichiometric manner [44]. By taking advantage of these properties we can mimic the regulation of a conditional essential gene in mycobacteria, making the Sh ble gene essential for survival in the presence of zeocin. Hence, the Sh ble gene was cloned under control of XylS/Pm, creating pMDX-zeo. Msmeg transformed with pMDX-zeo was induced for 5 hours to give the Sh ble gene a chance to be expressed at low levels. The culture was then serially diluted and spotted on agar plates containing both m-toluate and increasing concentrations of zeocin (Fig 5A); plates were then incubated for 48 hours. Uninduced controls were included in the experiment by simultaneous spotting on plates lacking m-toluate. As shown in Fig 5A, the initial number of bacteria seen on the uninduced and induced plates were identical (compare uninduced and induced samples, 0 μg/ml zeocin). However, induced cells could survive concentrations of zeocin up to 100–200 μg/ml due to the elevated production of the conditionally essential Sh ble gene product; a few small colonies were still visible in induced sample at the 200 μg/ml concentration. Moreover, the smaller colony sizes seen in the uninduced sample in the presence of zeocin concentrations as low as 1 μg/ml suggest that the uninduced cells have limited ability to grow under even low levels of zeocin stress (Fig 5A).

To get a more detailed picture of the zeocin-mediated growth inhibition, we diluted pre-induced cultures of Msmeg pMDX-zeo in increasing concentrations of zeocin with or without m-toluate. OD₆₀₀ was measured every two hours over a 48 hour period, and the absorbance measured for cells grown in increasing zeocin concentrations was determined at the time point the sample, induced or uninduced, grown in absence of zeocin had reached mid log phase (Fig 5B). Uninduced sample showed a significantly reduced growth rate in zeocin concentrations as low as 1 μg/ml, and growth was nearly completely inhibited at 2.5 μg/ml. Induced samples grew close to normally with up to 15 μg/ml zeocin before the growth rate was gradually reduced. Controls with empty vector lacking Sh ble grew only in media without zeocin (results not shown). These results show that the XylS/Pm expression system could serve as a tool to regulate conditionally essential genes in mycobacteria.

Pm compares favorably to Ptet in basal expression levels in Msmeg

So far, we have established XylS/Pm as a functional expression tool in Msmeg and Mtb, and demonstrated that it is a potential tool for conditional control of essential genes in mycobacteria. We wanted to investigate next how XylS/Pm compared to a commonly used inducible
mycobacterial expression tool, with emphasis on basal expression levels. We cloned the \textit{Sh} ble gene conferring resistance to zeocin under control of \textit{Ptet} in pUV15TetORm (Tn10-derived TetR controlled by the intermediate strong promoter \textit{Pimyc} and pAL5000 mycobacterial ori) [16] for direct comparison to XylS/\textit{Pm}, creating pTET-zeo. The \textit{Pm} and \textit{Ptet}-regulated systems were introduced to \textit{Msmeg} pMDX-zeo in increasing concentrations of zeocin, when the respective sample grown in the absence of zeocin reached mid log phase. Error bars represent standard deviations and the results represent three independent experiments.

\textbf{Fig 5. Regulation of the zeocin resistance gene in the presence of zeocin.} (A) \textit{Msmeg} transformed with pMDX-zeo was grown to OD\textsubscript{600} 0.05–0.1 before addition of 1.5 mM \textit{m}-toluate (induced) or ethanol carrier (uninduced) was added. Cells were then incubated for 5 hours at 30°C. The cells were normalized by OD\textsubscript{600}, serially diluted, and spotted on plates containing increasing amounts of zeocin and 1.5 mM \textit{m}-toluate (induced) or ethanol (uninduced) and incubated at 30°C for 2 days. (B) \textit{Msmeg} transformed with pMDX-zeo was pre-induced for 5 hours as described above, then diluted to OD\textsubscript{600} 0.005 and grown in triplicates in micro-plate wells in the presence of increasing concentrations of zeocin (0, 0.5, 2.5, 5, 7.5, 10, 15, 25, 50, 100, 150, 200 and 250 μg/ml) and 1.5 mM \textit{m}-toluate (induced) or ethanol carrier (uninduced) shaking at 37°C. Growth was monitored by Bioscreen, registering OD\textsubscript{600} every other hour. The samples are presented by the OD\textsubscript{600} of uninduced or induced \textit{Msmeg} pMDX-zeo in increasing concentrations of zeocin, when the respective sample grown in the absence of zeocin reached mid log phase. Error bars represent standard deviations and the results represent three independent experiments.

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\textit{Pm} and \textit{Ptet}-mediated basal expression was compared through determination of the amount of zeocin tolerated by the uninduced expression strains in log phase or stationary phase. While we observed comparable sensitivity to increasing concentrations of zeocin in uninduced \textit{Pm} and \textit{Ptet}-regulated strains in log phase (Fig 6A), the \textit{Ptet}-regulated strain could grow in the presence of higher concentrations of zeocin than the \textit{Pm}-regulated strain in stationary phase in the absence of inducer (Fig 6B). The pMDX-zeo strain was severely growth restricted in the presence of as little as 5 μg/ml zeocin, whereas the pTET-zeo strain could tolerate up to 250 μg/ml zeocin before reaching the same level of growth inhibition in stationary phase (Fig 6B). These results suggest that basal expression from \textit{Ptet} generates significantly higher levels of \textit{Sh} ble transcripts than \textit{Pm} during prolonged growth.

Comparison of induced samples of pMDX-zeo and pTET-zeo revealed higher levels of \textit{Sh} ble expression in \textit{Ptet} than \textit{Pm}-regulated cells. pTET-zeo cultures grew nearly unaffected by the
antibiotic up to 50 μg/ml zeocin in mid log phase (under optimized conditions for tet-inducible expression [16]) (Fig 6D), while growth of pMDX-zeo cultures was markedly inhibited by 10 μg/ml zeocin under optimized conditions (Fig 6C). Overall, Ptet-regulated cells showed around 10-fold higher induced levels of the zeocin resistance protein compared to Pm.

Although Pm drives lower levels of basal expression than commonly used inducible promoter systems during extended growth, Pm-mediated background expression might still be of biological relevance. In S6 Fig we cloned the alanine racemase gene (alr), which gene product catalyzes the conversion of L-alanine to D-alanine, under the control of XylS/Pm. For this particular gene, the background levels of alr expression were sufficient for Msmeg to produce enough D-alanine for normal growth. The same was seen for Ptet-regulated alr expression, and similar results were obtained for both Ptet and Pm-regulated alr expression on solid (S6 Fig) and liquid media (results not shown). These results underline that successful regulation of conditionally essential genes using inducible expression systems depends largely on the gene in question.

Fig 6. Pm–mediated basal expression is low and compares favorably to Ptet-mediated basal expression in Msmeg. (A-B) Msmeg transformed with pMDX-zeo or pTET-zeo was diluted to OD_{600} 0.005 in the presence or absence of m-toluate (1.5 mM) or atc (200 ng/ml), respectively, and increasing amounts of zeocin (0, 0.5, 2.5, 5, 7.5, 10, 15, 25, 50, 100, 150, 200, 250 or 500 μg/ml). Samples were grown in triplicates and monitored for 120 hours by a Bioscreen, registering OD_{600} every other hour. Basal expression from Pm and Ptet is presented by growth of Msmeg pMDX-zeo and Msmeg pTET-zeo in increasing concentrations of zeocin, when the respective sample grown in the absence of zeocin reached mid log phase (A) or stationary phase (B). (C) Induced and uninduced samples of pMDX-zeo strain in late log phase. (D) Induced and uninduced samples of pTET-zeo strain in mid log phase. The results represent two independent experiments.

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**Pm** compares favorably to **Ptet** in basal expression levels in **Mtb**

The behavior of **Pm** in **Mtb** was investigated by transformation of pMDX-luc or pUV15tetORm::luciferase, and comparing **Pm** and **Ptet**-mediated basal expression through determination of the levels of luciferase produced by the two strains under induced and uninduced conditions. As seen for **Sh ble**, pUV15tetORm::luciferase revealed higher levels of luciferase expression compared to **Pm**-regulated cells during exponential growth (Fig 7A). We measured luciferase activity in the uninduced samples and we observed a background expression from **Pm** that was low throughout the experiment. For pUV15tetORm::luciferase we observed an increase in background expression over time (Fig 7B).

Taken together, our results indicate that **Mtb** basal expression mediated by **Pm** is significantly lower than **Ptet** basal expression, under the given conditions. While **Ptet**-regulated constructs result in higher induction, XylS/**Pm** may provide an advantage for experiments where consistent low background expression is of particular importance.

**Discussion**

In this study we have successfully introduced a benzoic acid-inducible expression system to mycobacteria, and expanded the repertoire of mycobacterial tools for studying gene function and essentiality. XylS/**Pm** has previously been thoroughly studied and applied in Gram-negatives (reviewed in [6]). Importantly, by adapting **Pm**-regulated expression to mycobacteria, we demonstrate for the first time the great potential of expanding the use of this expression system to non-Gram-negative species.

Upon induction with **m**-toluate, we show a typical 90-fold increase for **Pm**-mediated reporter gene expression in **Msmeg** (Fig 2A), an induction ratio comparable to or higher than the reported ratios of all other published inducible mycobacterial expression systems based on reporter gene assays [15–19, 21, 23]. The exception is the pristamycin-inducible system with a reported 400-fold induction in **M. tuberculosis** [21]. We also report a dose-dependent response, with higher concentrations of inducer leading to increased induction ratios (Fig 2A). Although we might reach even greater expression levels by further increasing inducer concentrations,
due to a decrease in growth rate in the presence of more than ~1.5 mM \( m \)-toluate (S1 Fig) we chose to largely avoid concentrations above 1.5 mM. Furthermore, in this study we focused on one of many XylS/Pm-inducers known to work in Gram-negative bacteria [45]. Other inducers might compare favorably to \( m \)-toluate when it comes to both inducibility and toxicity in mycobacteria.

While Pm-mediated expression clearly functions in \( Mtb \), we are not able to reach the same level of induced expression as in \( Msmeg \), and we do not see the same dose-dependent response (Fig 3A). Nevertheless, as in \( Msmeg \), we see a time-dependent response, with a peak in expression in late log phase (Fig 3A and 3B). The ~10-fold increase in induction observed using 0.15 mM \( m \)-toluate is comparable to the 0.15 mM-induction of \( Msmeg \), but increasing inducer concentration further does not boost expression in \( Mtb \). Mycobacterial drug efflux pumps have been identified in \( Mtb \) for inducers like tetracycline [46, 47], and we cannot exclude similar active pumping of \( m \)-toluate out of the cell, resulting in a decreased maximal response to induction. Alternatively, \( m \)-toluate could be metabolized by mycobacteria, decreasing the concentration of inducer in the cell, as oxidation of benzoate and related substances was reported in early mycobacterial research [48]. Another inducer might thus be superior to \( m \)-toluate for XylS/Pm-induction in \( Mtb \).

XylS/Pm can be suitable to regulate conditionally essential genes in mycobacteria. We cloned the \( Sh \) ble gene, which confers zeocin resistance, under the control of Pm (pMDX-zeo) in \( Msmeg \). Our results demonstrated that induced cells containing pMDX-zeo survived significantly better in the presence of zeocin than uninduced cells, with growth of uninduced cells inhibited by as little as 1 \( \mu \)g/ml zeocin (Fig 5). Although the growth rate of uninduced cells that carry the resistance gene showed inhibition at 1 \( \mu \)g/ml zeocin, cultures did grow, albeit slowly, in the presence of up to 5–10 \( \mu \)g/ml of the antibiotic. In contrast, cells transformed with a vector lacking \( Sh \) ble, did not grow at 0.5 \( \mu \)g/ml zeocin (results not shown). This observation suggests that the induced basal expression of the resistance gene is appreciably low, but might be biologically relevant depending on the gene in question. When we controlled expression of the alanine racemase gene with the Pm and Ptet promoters, we observed that background expression of the conditionally essential gene was sufficient to maintain a wild type phenotype.

Comparing Pm-mediated background expression with the background expression of the widely used tet-inducible Ptet-promoter [16], Pm in fact drives lower levels of basal expression than Ptet both in \( Msmeg \) and \( Mtb \) (Figs 6B and 7B). Here, Ptet basal expression level (unlike Pm) is dependent on active repression of transcription. A codon usage adaption of the Tn10-derived TetR for optimal gene regulation in \( Mtb \) resulted in 3-fold improved repression in \( Msmeg \) when the repressor was expressed episomally, due to increased TetR steady state levels [31]. It is plausible that the increase in Ptet-mediated background expression we observed in stationary phase is due to a decrease in TetR steady state levels during prolonged growth, making the repressor-independent nature of Pm-mediated basal levels a likely explanation for the lower background expression seen in XylS/Pm. Reverse gene regulation tools, where the addition of the effector molecule turns transcription off instead of on, have been developed to silence genes in mycobacteria [31, 49, 50]. This approach has proved useful to study conditional mutants [50, 51], however, these systems are still dependent on sufficient expression of the repressor to keep the gene expression low. In our system, the consistent low basal expression coupled to titratable, strong induction makes Pm an appealing option for regulated gene expression in mycobacteria, and XylS/Pm may be a particularly good alternative when a consistent, low background expression is important.

Basal rates of Pm-mediated expression were not reduced by addition of atc, which suggests that the XylS production is unlikely to be high enough to drive transcription from the uninduced construct. Instead, the low background expression observed in our experiments is most
probably due to leaky transcription from the \textit{Pm} promoter itself initiated at a low rate by the endogenous transcription machinery. Consequently, future work toward even tighter control of \textit{Pm} basal transcription in mycobacteria would include random mutagenesis of the promoter or the 5’ untranslated region, which previously has been shown to improve XylS/\textit{Pm}-mediated regulation in \textit{E. coli} [43, 52].

An important aspect of mycobacterial research is the modulation of gene expression during infection. Tet-inducible expression systems are thus far the only tools for induction in macrophage or animal models of mycobacterial infection [15, 16, 18]. The XylS/\textit{Pm}-inducers are weak acids passively diffusing through plasma membranes. Medina \textit{et al.} successfully induced expression of dTomato reporter protein under the control of the \textit{Pm} promoter in \textit{Salmonella} during the infection of HeLa cells, using salicylate as inducer [53]. These results support the potential function of our adapted XylS/\textit{Pm} expression system during mycobacterial infections of macrophages. Benzoic acid is commonly used as a food preservative [54, 55], which together with the fact that benzoic acids can readily diffuse across plasma membranes suggests that inducers of \textit{Pm}-mediated expression could be excellent candidates for oral administration in animal models. Thus, XylS/\textit{Pm} has potential to become the second inducible regulator of mycobacterial expression applicable in animal models of tuberculosis.

### Supporting Information

**S1 Fig. Growth curves in the presence of \textit{m}-toluate.** (A) \textit{Msmeg} and (B) \textit{Mtb} grown in 7H9 medium with increasing concentrations of \textit{m}-toluate. OD\textsubscript{600} was determined after 2.5, 5, 8, 10 and 24 hours (\textit{Msmeg}), or 4, 10 and 14 days (\textit{Mtb}).

(TIF)

**S2 Fig. Induced luciferase expression normalized to OD\textsubscript{600} and RLU.** \textit{Msmeg} transformed with pMDX or pMDX-luc was grown to stationary phase, subcultured and grown to OD\textsubscript{600} 0.05–0.1 before induction by 1.5 mM \textit{m}-toluate (induced) or ethanol carrier (uninduced). Cultures were incubated for 24 hours with shaking at 37°C before luciferase expression was measured. RLU was normalized to (A) OD\textsubscript{600} or (B) CFU, reaching 110 and 240-fold induction, respectively.

(TIF)

**S3 Fig. XylS/\textit{Pm} is inducible when integrated in the chromosome.** \textit{Msmeg} transformed with pMDXint, pMDXint-luc or pMDX-luc were grown to stationary phase, subcultured and grown to OD\textsubscript{600} 0.05–0.1 before addition of 1.5 mM \textit{m}-toluate (induced) or ethanol carrier (uninduced). Cultures were incubated for 24 hours with shaking at 30°C before luciferase expression was measured. Results are presented as RLU normalized to OD\textsubscript{600}. Results are representative for three individual experiments.

(TIF)

**S4 Fig. Induction of \textit{Pm} is robust at both 30 and 37°C.** \textit{Msmeg} transformed with pMDX-luc was grown to stationary phase, subcultured and grown to OD\textsubscript{600} 0.05–0.1 before addition of 0.05, 0.25, 1 or 1.5 mM \textit{m}-toluate (induced) or ethanol carrier (uninduced). Cultures were incubated for 24 hours with shaking at 30 or 37°C before luciferase expression was measured. Results are presented as fold induction of RLU normalized to OD\textsubscript{600} of induced compared to uninduced samples. The fold inductions of RLUs comparing 30°C and 37°C were obtained in separate luciferase assays. Results are representative for two or more independent experiments.

(TIF)
S5 Fig. Comparison of Ptet and Pm-mediated expression. Msmeg containing pTET-zeo or pMDX-zeo was grown to stationary phase, then diluted to OD$_{600}$ 0.005 in the presence or absence of inducer. pMDX-zeo was induced with 1.5 mM m-toluate, and pTET-zeo was induced with 200 ng/ml anhydro-tetracycline (atc). The samples were grown in triplicates in micro-plate wells in increasing amounts of zeocin (0, 0.5, 2.5, 5, 7.5, 10, 15, 25, 50, 100, 150, 200, 250 or 500 μg/ml). Growth was monitored for 120 hours using a Bioscreen, shaking at 37°C, registering OD$_{600}$ every other hour. All growth curves represent the average of 3 replicates.

(TIF)

S6 Fig. Alanine racemase gene controlled by Pm and Ptet. Msmeg wt, DM22 mutant and DM22 containing pTET-alr or pMDX-hyg-alr were grown to stationary phase then serial diluted and spotted onto agar plates with or without D-alanine.

(TIF)

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Author Contributions

Conceived and designed the experiments: MD AB NK MM THF SV EJR MS. Performed the experiments: MD AB NK MM MS. Analyzed the data: MD AB NK MM THF SV EJR MS. Contributed reagents/materials/analysis tools: AB SV EJR MS. Wrote the paper: MD AB NK MM THF EJR MS.

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