β-Lactamase can function as a reporter of bacterial protein export during *Mycobacterium tuberculosis* infection of host cells

Jessica R. McCann,1 Justin A. McDonough,1 Martin S. Pavelka2 and Miriam Braunstein1

1Department of Microbiology and Immunology, University of North Carolina School of Medicine, Chapel Hill, NC 27599-7290, USA
2Department of Microbiology and Immunology, University of Rochester School of Medicine and Dentistry, Rochester, NY 14642, USA

*Mycobacterium tuberculosis* is an intracellular pathogen that is able to avoid destruction by host immune defences. Exported proteins of *M. tuberculosis*, which include proteins localized to the bacterial surface or secreted into the extracellular environment, are ideally situated to interact with host factors. As a result, these proteins are attractive candidates for virulence factors, drug targets and vaccine components. Here we describe a β-lactamase reporter system capable of identifying exported proteins of *M. tuberculosis* during growth in host cells. Because β-lactams target bacterial cell-wall synthesis, β-lactamases must be exported beyond the cytoplasm to protect against these drugs. When used in protein fusions, β-lactamase can report on the subcellular location of another protein as measured by protection from β-lactam antibiotics. Here we demonstrate that a truncated TEM-1 β-lactamase lacking a signal sequence for export (‘BlaTEM-1) can be used in this manner directly in a mutant strain of *M. tuberculosis* lacking the major β-lactamase, BlaC. The ‘BlaTEM-1 reporter conferred β-lactam resistance when fused to both Sec and Tat export signal sequences. We further demonstrate that β-lactamase fusion proteins report on protein export while *M. tuberculosis* is growing in THP-1 macrophage-like cells. This genetic system should facilitate the study of proteins exclusively exported in the host environment by intracellular *M. tuberculosis*.

INTRODUCTION

Tuberculosis is responsible for nearly two million deaths each year (WHO, 2007). *Mycobacterium tuberculosis*, the causative agent of this disease, is an intracellular pathogen and the ability of this bacterium to survive and grow in macrophages is essential to its virulence. Multiple processes are likely employed by *M. tuberculosis* to avoid destruction in macrophages. These include residing in a phagosome that fails to mature into an acidified phagolysosome and in macrophages. These include residing in a phagosome or secreted into the extracellular environment. Consequently, exported and secreted proteins make good candidates for virulence factors, drug targets for disease intervention, and vaccine antigens.

*Mycobacteria* possess two conserved pathways for exporting proteins: the general secretion (Sec) pathway and the twin-arginine translocation (Tat) pathway (Braunstein et al., 2001; Kurtz & Braunstein, 2005; McDonough et al., 2005; Owens et al., 2002; Posey et al., 2006; Saint-Joanis et al., 2006). These systems recognize precursor proteins synthesized with amino-terminal signal sequences and transport them across the cytoplasmic membrane (DeLisa et al., 2003; Mori & Ito, 2001). The proteins exported by these pathways can remain associated with the cell envelope or be further secreted by the bacterium. The signal sequences of Sec and Tat substrates share a similar domain structure; however, Tat substrates are distinguished by the presence of the twin-arginine motif, R-R-x-Φ-Φ (Φ = uncharged residue). The two pathways also differ in their mode of transport. Sec substrates are translocated across the cytoplasmic membrane in an unfolded state, whereas Tat substrates are translocated in a folded conformation. *M. tuberculosis* also has at least two specialized protein export pathways: the SecA2-dependent system and the ESX-1 (ESAT-6) system (Braunstein et al., 2003; Guinn et al., 2004; Hsu et al., 2003; Pym et al., 2003; Stanley et al., 2003). Interestingly, both pathways appear capable of secreting specific subsets of proteins that lack conventional Sec or Tat signal sequences.
In *M. tuberculosis*, proteomic and genetic methods have been used to experimentally identify proteins exported beyond the cytoplasm (reviewed by Kurtz & Braunstein, 2005). The genetic methods rely on reporter enzymes that are fused to *M. tuberculosis* protein sequences and report on the subcellular location of the fusion proteins (Braunstein et al., 2000; Chubb et al., 1998; Downing et al., 1999; Lim et al., 1995; Wiker et al., 2000). Surrogate hosts such as non-pathogenic *Mycobacterium smegmatis* or *Escherichia coli* have been used in most of these studies, often because endogenous enzyme activities in *M. tuberculosis* precluded their use directly in the pathogen. The use of surrogate hosts is a problem for identifying proteins that are only exported by pathogenic *M. tuberculosis*.

β-Lactamase is an export reporter that was not initially employed directly in *M. tuberculosis* because of endogenous β-lactam resistance. β-Lactamase catalyses the hydrolysis of β-lactams, a class of antibiotic that targets cell-wall biosynthetic enzymes located outside of the cytoplasmic membrane. Therefore, β-lactamase must be exported beyond the cytoplasm to protect the bacterium from the drug. For this reason, when fused to another protein, it can be used as an export reporter with β-lactam resistance as a powerful indicator of export. We recently reported that a ΔblaC mutant of *M. tuberculosis*, lacking the chromosomally encoded β-lactamase BlaC, is β-lactam sensitive (Flores et al., 2005). Further, we showed that BlaC is a native Tat substrate and that a truncated 'BlaC lacking a signal sequence can function as a reporter of Tat-dependent export directly in a ΔblaC mutant of *M. tuberculosis* (McDonough et al., 2005). This was shown by fusing a Tat signal sequence to 'BlaC and demonstrating that the resulting hybrid protein confers resistance to the β-lactam antibiotic carbenicillin in the ΔblaC background. Interestingly, the 'BlaC reporter works with Tat- but not Sec-expressed proteins. Here we expanded the β-lactamase tools that can be used directly in *M. tuberculosis* by demonstrating that the TEM-1 β-lactamase (BlaTEM-1), originally identified in a clinical isolate of *E. coli* (Datta & Kontomichalou, 1965), functions as an export reporter in the ΔblaC mutant of *M. tuberculosis*. The 'BlaTEM-1 reporter has the significant advantage of being compatible with both Sec and Tat signal sequences.

The proteomic and genetic approaches used in previous work for identifying exported proteins of *M. tuberculosis* are limited by their reliance on *in vitro*-grown bacteria. Consequently, a potentially interesting collection of proteins only exported or secreted while *M. tuberculosis* are inside host cells are missed. In this paper, we demonstrate that β-lactamase reporters have the novel capability of identifying *M. tuberculosis* proteins that are exported during intracellular growth in β-lactam-treated THP-1 macrophage-like cells. The system we describe will be of significant value for identifying the most interesting category of exported *M. tuberculosis* proteins – those exported during growth in the host environment.

**METHODS**

**Bacterial strains, media and growth conditions.** *Escherichia coli* DH5α was grown in Luria–Bertani medium (Fisher) supplemented with the following concentrations of antibiotics as required: carbenicillin, 100 μg ml⁻¹; kanamycin, 40 μg ml⁻¹. *M. tuberculosis* strains H37Rv (wild-type), PM638 (ΔblaC, H37Rv) (Flores et al., 2005) and all derivative strains were cultured in Middlebrook 7H9 medium or on Middlebrook 7H10 agar medium (Difco; BD Biosciences) supplemented with 10 % ADS [0.5 % BSA, fraction V (Roche); 0.2 % glucose (dextrose); and 0.85 % NaCl], 0.5 % glycerol and 0.05 % Tween 80 (Fisher). Antibiotics for mycobacteria were used at the following concentrations: carbenicillin, 50 μg ml⁻¹; kanamycin, 20 μg ml⁻¹. 7H10 plates supplemented with carbenicillin lacked Tween, as the combination of Tween and carbenicillin appeared detrimental to growth of fusion-expressing strains.

**Construction of ‘blaTEM-1’ fusion plasmids**

Plasmids used in this study are listed in Table 1. All subcloned PCR products were sequenced and determined to be error free. Sequence encoding the mature domain (lacking the N-terminal signal sequence) of *E. coli* BlaTEM-1 was amplified from pUC19 plasmid DNA (Invitrogen) using the following primers: TEMbla1 (5’- AGATCTCACCAGAAGCTGTGAAAAG-3’) and TEMbla2 (5’- GTTACCAATGCTTAATCAGTGAGGCACC-3’). The resulting PCR product was cloned into the pCC1 vector (Epitope) to generate pJM114. The ‘blaTEM-1’ reporter was subcloned as a BglII–BanHI fragment into each of the multi-copy vectors described below.

(i) Δss, ‘blaTEM-1’. ‘blaTEM-1’ was digested from pJM114, end-filled with Klenow and cloned into MscI-cut pMV261. The resulting plasmid, pJES102, contains the ‘blaTEM-1’ reporter without a fused signal sequence cloned downstream of the hsp60 promoter.

(ii) ssplcB–‘blaTEM-1’. The ‘blaTEM-1’ fragment was subcloned into BamHI-cut pMB222. The resulting plasmid, pJES101, contains an in-frame fusion of DNA encoding the signal sequence of PglB/Rv2350c (ssplcB) to ‘blaTEM-1’ under the control of the hsp60 promoter.

(iii) ssmp63–‘blaTEM-1’. The ‘blaTEM-1’ fragment was subcloned into BamHI-cut pMB227. The resulting plasmid, pJES103, contains an in-frame fusion of ssmp63 (Rv1926c) to ‘blaTEM-1’ under the control of the hsp60 promoter.

(iv) ssmp83–‘blaTEM-1’. DNA encoding the signal sequence and the first 31 amino acids of the mature *M. tuberculosis* Mpt83 (Rv2873) protein along with the native mpt83 promoter (Juarrez et al., 2001) was amplified from *M. tuberculosis* genomic DNA using the following primers: mpt83HindIIIF (5’-CAAGCTTTCGTTCCATGGATGAGGAAGG-3’) and mpt83HindIII (5’-CAAGCTTTCGTTCCATGGATGAGGAAGG-3’) and cloned into the pCR2.1 vector (Invitrogen) to generate pJES125. A HindIII fragment from pJES125, carrying ssmp83 and upstream genomic sequence was cloned into HindIII-cut pJES128 (Table 1). The resulting plasmid, pJES129, contains an in-frame fusion of ssmp83 to ‘blaTEM-1’ under the control of the native mpt83 promoter (P_mpt83).

**Protein quantification by immunoblotting.** Whole-cell lysates of *M. tuberculosis* strains were prepared as described previously (Braunstein et al., 2001) with the following modifications. *M. tuberculosis* cultures were grown in 5 ml volumes to mid-exponential phase. The cultures were washed twice and resuspended in PBS containing 0.02 % Tween 80. An equal volume of 10 % formalin was added to the washed cultures, which were then incubated at room temperature for 1 h with frequent mixing by inversion. The formalin-fixation step was necessary to kill *M. tuberculosis* before further processing. Bacteria were then harvested by centrifugation at...
Macrophage infections. THP-1 cells were maintained in RPMI (Gibco)/10% heat-inactivated fetal calf serum (FCS) at 37 °C and 5% CO₂. To prepare THP-1 monolayers for infection, cells were spun down at 300 g, washed once in RPMI, then resuspended in RPMI/10% FCS at a concentration of 1 x 10⁶ cells ml⁻¹. Cells were seeded into eight-well tissue culture slides at 2 x 10⁵ cells well⁻¹ and treated with phorbol myristate acetate (PMA) at a final concentration of 50 ng ml⁻¹ for 48 h. M. tuberculosis was grown to mid-exponential phase (OD₆₀₀ 0.5–1.0). Immediately prior to infection, the bacterial culture was pelleted, washed once in PBS containing 0.05% Tween 80 (PBS-Tw), and resuspended in an equal volume of PBS-Tw. The culture was then briefly sonicated to break up clumps of bacteria, diluted in RPMI/10% FCS medium and added to the THP-1 monolayer at m.o.i. = 0.1.

THP-1 monolayers were infected with M. tuberculosis strains for 4 h at 37 °C and 5% CO₂. Overlying medium was then removed, the monolayers were washed three times with RPMI to remove non-cell-associated bacteria, and triplicate wells were lysed and plated to determine uptake (day 0 time point). The infected monolayers were then overlaid with RPMI/10% FCS, or RPMI/10% FCS supplemented with carbenicillin, and maintained at 37 °C and 5% CO₂. At 3 days post-infection, the overlying medium was replenished with RPMI/10% FCS medium or medium supplemented with carbenicillin, as appropriate. On days 1, 3 and 5 post-infection, triplicate wells for each infection were washed to remove antibiotic and lysed with 0.05% SDS. The resulting lysates were diluted and plated on 7H10 agar to enumerate intracellular bacteria during the course of infection. On day 0 and day 5 of the infection, cell lysates were also plated on 7H10 agar supplemented with 50 μg carbenicillin ml⁻¹. This demonstrated that selection of spontaneous β-lactam-resistant mutants did not occur during the course of infection. To determine the appropriate carbenicillin concentration necessary to kill intracellular bacteria, THP-1 infection experiments were performed with a range of antibiotic concentrations (see Fig. 4b). Carbenicillin at 1 mg ml⁻¹ was determined to be the lowest concentration of M. tuberculosis and E. coli used for each experiment. pCC1 was transformed into E. coli DH5α and pJM109 was transformed into E. coli DH5α. pMB192 and pMB227 were transformed into E. coli DH5α to confer resistance to ampicillin and carbenicillin, respectively. Plasmid pCC1 was transformed into E. coli DH5α and pJM109 was transformed into E. coli DH5α. pMB192 and pMB227 were transformed into E. coli DH5α to confer resistance to ampicillin and carbenicillin, respectively. Plasmid pCC1 was transformed into E. coli DH5α and pJM109 was transformed into E. coli DH5α. pMB192 and pMB227 were transformed into E. coli DH5α to confer resistance to ampicillin and carbenicillin, respectively.

### Table 1. Plasmids used in this study

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Genotype</th>
<th>Description</th>
<th>Source</th>
</tr>
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<tbody>
<tr>
<td>pCC1</td>
<td>cat oriV ori2</td>
<td>CopyControl (single copy) blunt cloning vector</td>
<td>Epicentre</td>
</tr>
<tr>
<td>pCR2.1</td>
<td>blapho CoIE1</td>
<td>TA cloning vector</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pJM101</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM227</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM110</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM111</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM113</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM114</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM125</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM128</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
<tr>
<td>pJM129</td>
<td>P_rPM261 ssmp63 hsp60 Cat M. tuberculosis oriM CoIE1</td>
<td>M. tuberculosis cpdB signal sequence in pMM261 under control of hsp60promoter</td>
<td>McDonough et al. (2005)</td>
</tr>
</tbody>
</table>

3000 r.p.m., washed once in PBS 0.02% Tween to remove residual formalin, and bead-beaten lysates were then obtained from each sample. Protein concentration for each lysate was measured using a bicinchoninic acid protein quantification kit (Pierce). Lysates were boiled for 10 min, subjected to SDS-PAGE and immunoblots were performed using standard conditions. Primary antibodies specific for BlaTEM-1 were used at a concentration of 1:20000. Bands were visualized using Western Lightning Chemiluminescent Reagent Plus (Perkin Elmer) and quantified using ImageJ Image Processing and Analysis software (http://rsb.info.nih.gov/ij/). Whole-cell lysates with the highest level of expression were diluted to enable direct comparison of all hybridization signals on a single blot. The comparative quantification was determined by measuring pixel density of an equal area for each blotted lysate in duplicate. Signal intensity per μg of whole-cell lysate loaded was determined and is reported as the amount relative to protein detected in the `BlaTEM-1`-expressing strain.

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Results

'BlaTEM-1 is exported by Sec and Tat signal sequences in M. tuberculosis

β-Lactamase is an ideal reporter for protein export because it must be localized beyond the bacterial cytoplasmic membrane to effectively protect the bacterium from β-lactam antibiotics. Therefore, it can be used in protein fusions to identify proteins that are extracytoplasmic. An attractive feature of a β-lactamase reporter is that a selection for β-lactam-resistant colonies can be performed, as opposed to a more labour-intensive screen. In the past, we showed that the endogenous β-lactamase of M. tuberculosis BlaC can function as a reporter of export exclusively by the Tat pathway when expressed in the β-lactam-sensitive ΔblaC mutant of M. tuberculosis or ΔblaS mutant of M. smegmatis (McDonough et al., 2005). Since the Sec pathway is likely responsible for the majority of protein export in M. tuberculosis, we were interested in utilizing a β-lactamase reporter that additionally works with Sec-exported proteins. For this reason, we tested the E. coli TEM-1 β-lactamase (BlaTEM-1), which has been used in other bacteria to report on proteins exported by Sec, Tat, type II and type III secretion systems (Broome-Smith et al., 1990; Charpentier & Oswald, 2004; Sauvonnet & Pugsley, 1996; Stanley et al., 2002). A series of multi-copy kanamycin-marked 'blaTEM-1' plasmids were constructed and electroporated into the ΔblaC mutant of M. tuberculosis (Fig. 1). The resulting kanamycin-resistant strains were tested for the ability to grow in the presence of 50 μg ml⁻¹ of the β-lactam antibiotic that caused optimal killing of sensitive intracellular M. tuberculosis and was used in subsequent experiments.

Finaly, we tested the signal sequence of PlcB, a proven cell-wall-associated lipoprotein, Mpt83 (Hewinson et al., 1996), was fused to 'BlaTEM-1. This construct also conferred β-lactam resistance to ΔblaC M. tuberculosis (Fig. 1). Of note, the ssMpt83-'BlaTEM-1 fusion protein included the first 31 amino acids of the mature Mpt83 protein as well as the native mpt83 promoter, which is reported to be active at very low levels in vitro (Hewinson et al., 1996; Said-Salim et al., 2006).

To determine whether the ssPlcB-'BlaTEM-1 fusion was expressed by the Tat pathway, it was tested in ΔblaS M. smegmatis and in a ΔtatA ΔblaS M. smegmatis double mutant (McDonough et al., 2005) in two independent experiments. When the ssPlcB-'BlaTEM-1 fusion protein was expressed in ΔblaS M. smegmatis, 92% of colonies...
were carbenicillin resistant. However, when the same construct was expressed in the ΔtatA ΔblaS mutant, only an average 7% of colonies were carbenicillin resistant, indicating that the Tat pathway functions in the export of this fusion protein. To show that a functional Tat pathway was not required for export of the Sec signal sequence 'BlaTEM-1 fusion, we similarly evaluated export of ssMpt63-'BlaTEM-1. When expressed in ΔblaS and the ΔtatA ΔblaS mutants, ssMpt63-'BlaTEM-1 conferred carbenicillin resistance to 90% and 95% of colonies, respectively. This indicated, as expected, no role for the Tat pathway in exporting a Sec signal sequence 'BlaTEM-1 fusion.

In each example where a M. tuberculosis signal sequence (Sec or Tat) was fused to 'BlaTEM-1, ΔblaC M. tuberculosis was protected from β-lactam attack. To demonstrate that the inability of the 'BlaTEM-1 reporter lacking a signal sequence to protect against carbenicillin was due to lack of export, as opposed to lack of expression, whole-cell extracts of 'BlaTEM-1 expression strains were prepared and assayed for cell-associated β-lactamase. To test for enzyme activity, we used the chromogenic β-lactam nitrocefin, which turns red following cleavage by β-lactamase (O’Callaghan et al., 1972). During a 15 min incubation the nitrocefin was hydrolysed by all strains expressing 'BlaTEM-1 constructs, while ΔblaC M. tuberculosis demonstrated no activity, similar to PBS alone (data not shown). Importantly, β-lactamase activity was detected with the truncated 'BlaTEM-1 reporter lacking a signal sequence. In fact, the lysate from the 'BlaTEM-1 strain converted nitrocefin to the red product almost instantaneously and faster than any other strain tested. We similarly detected β-lactamase activity in whole-cell lysates of ΔblaC M. tuberculosis expressing the 'BlaC reporter lacking its native signal sequence.

We also compared the level of each 'BlaTEM-1 fusion protein present in whole-cell lysates from the respective M. tuberculosis strains by immunoblots with antibodies specific for BlaTEM-1. This revealed a wide variation in the amount of 'BlaTEM-1 protein produced by the different strains (Fig. 3). The non-exported 'BlaTEM-1 expressed off the hsp60 promoter (P_hsp60) was the most abundant protein detected. P_hsp60 is considered a relatively strong promoter and is, therefore, present on many mycobacterial shuttle vectors (Stover et al., 1991). In comparison, the P_hypo-driven ssPibC-'BlaTEM-1 and ssMpt63-'BlaTEM-1 were expressed at lower levels (59% and 0.9% of the level of the non-exported 'BlaTEM-1 construct, respectively). Since mpt83 is expressed at relatively low levels in vitro, we expected the ssMpt83-'BlaTEM-1 fusion to be weakly expressed (Hewinson et al., 1996; Said-Salim et al., 2006; Schnappinger et al., 2003). In fact, it was nearly undetectable by immunoblotting, present at only 0.4% of the amount of non-exported 'BlaTEM-1 construct. The bands detected on the immunoblot are in general agreement with the predicted molecular mass of the expressed proteins. 'BlaTEM-1, lacking a signal sequence, has a predicted size of 28 kDa. Since whole-cell lysates were analysed in these experiments it is possible to see processed

**Fig. 2.** 'BlaTEM-1 does not provide β-lactam resistance to ΔblaC M. tuberculosis. Plasmids encoding the indicated ‘blaTEM-1 fusions were electroporated into M. tuberculosis ΔblaC. The resulting strains were then plated on 7H10 plates supplemented with either kanamycin or 0.05% Tween or kanamycin and carbenicillin without Tween. Plates were inspected for growth following 21–25 days incubation. Not shown are colonies expressing ssMpt63-'BlaTEM-1 and ssPibC-'BlaTEM-1; growth on plates containing carbenicillin for these strains was similar to that conferred by ssMpt63-'BlaTEM-1.

**Fig. 3.** 'BlaTEM-1 fusion proteins are detected at different amounts in M. tuberculosis whole-cell lysates. Proteins present in whole-cell lysates (WCL) from each of the indicated ΔblaC strains were separated by SDS-PAGE and immunoblotted using primary antibody specific for BlaTEM-1. Comparative signal was quantified by measuring the pixel density of an equal area for each blotted lysate in duplicate. Average signal intensity per mg WCL is reported as the amount relative to protein detected in the 'BlaTEM-1 expressing strain. Due to the different amounts of protein in each strain, it was necessary to load dilutions of the 'BlaTEM-1- and ssPibC-'BlaTEM-1-expressing lysates so that signal from less abundant protein fusions could be simultaneously detected. There was no detectable signal with the WCL from the ΔblaC mutant carrying empty pMV261 plasmid.
The ΔblaC mutant of M. tuberculosis is sensitive to β-lactams during intracellular growth in human THP-1 cells

β-Lactam antibiotics can be used for clinical treatment of intracellular pathogens such as Listeria monocytogenes (Safdar & Armstrong, 2003), and have been shown to reduce the population of phagocytosed Staphylococcus aureus (Barcia-Macay et al., 2006). This indicates that β-lactams can enter macrophages and inhibit intracellular growth of some bacteria. The ΔblaC mutant of M. tuberculosis is sensitive to β-lactams in vitro, and we set out to test if this mutation also makes M. tuberculosis susceptible to β-lactams during growth in host cells.

Intracellular growth of the ΔblaC mutant was not previously evaluated; therefore, we first tested the ability of this mutant to grow within human monocytic THP-1 cells. THP-1 cells were infected at a m.o.i. of 0.1 with either the ΔblaC mutant or the virulent parental H37Rv strain. After a 4 h period of infection, the THP-1 monolayer was washed to remove non-cell-associated bacilli and fresh medium was added back. Growth over 5 days was assessed by plating of infected host-cell lysates for viable bacilli. The ΔblaC mutant showed no difference in intracellular growth when compared to H37Rv (Fig. 4a). Of note, we confirmed that M. tuberculosis does not grow in the THP-1 culture medium as previously reported (Zhang et al., 1998).

To determine if the ΔblaC mutant was sensitive to β-lactams during intracellular growth, THP-1 cells were infected with ΔblaC M. tuberculosis and, following the washes to remove extracellular bacilli, medium containing different concentrations of carbenicillin was added to the infected monolayers. After 5 days incubation, the infected monolayers were washed to remove carbenicillin and lysed to plate for viable bacilli. In the absence of carbenicillin, the ΔblaC mutant grew in THP-1 cells as previously seen. However, as the concentration of carbenicillin during the intracellular growth period increased, growth of the mutant diminished. At carbenicillin concentrations of ≥0.8 mg ml⁻¹, substantial killing of the mutant was observed (Fig. 4b). These results indicated that the ΔblaC mutant is sensitive to β-lactam antibiotics during intracellular growth, and it suggested that the β-lactamase reporters could be used to study protein export during intracellular growth. Additional experiments showed that a carbenicillin concentration of 1 mg ml⁻¹ was sufficient to achieve significant killing of the ΔblaC mutant of M. tuberculosis in THP-1 cells, and this concentration was used in all subsequent experiments.

Export of β-lactamase protects intracellular ΔblaC M. tuberculosis from β-lactam antibiotics

A reporter system that works with intracellularly growing M. tuberculosis would be of great value for identifying exported proteins that are expressed and exported only during infection. Having shown that the ΔblaC mutant was

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The ΔblaC mutant of M. tuberculosis does not have a growth defect and is sensitive to β-lactam antibiotic in human THP-1 macrophage-like cells. (a) THP-1 cells were seeded into eight-well chamber slides, and triplicate wells were infected with either H37Rv (wild-type) or ΔblaC M. tuberculosis at a m.o.i. of 0.1 bacilli per macrophage. At 4 h (day 0), and 1, 3 and 5 days post-infection, infected wells were washed, lysed and plated for intracellular bacteria. Error bars represent standard deviation of the mean of c.f.u. in triplicate wells. (b) THP-1 cells were infected with ΔblaC M. tuberculosis as in (a). Following a 4 h uptake period, wells were washed, and the indicated concentrations of carbenicillin were added to infected wells. Infected cells were lysed and plated at 4 h (Day 0) and 5 days post-infection to enumerate intracellular bacteria. The dashed line represents average intracellular c.f.u. at 4 h post-infection. Error bars represent standard error of the mean of quadruplicate wells combined from two replicates.
sensitive to β-lactams during intracellular growth, we tested if β-lactamase could be used to report on protein export by M. tuberculosis growing in host cells. We tested fusion proteins expressing the ΔblaC and ΔblaTEM-1 reporters for the ability to protect the ΔblaC mutant in β-lactam-treated THP-1 cells. In each experiment we compared an exported fusion protein to the truncated reporter alone. To test the ΔblaC reporter, which works with Tat-exported proteins only, THP-1 cells were infected with the M. tuberculosis ΔblaC mutant expressing ssPlcB-ΔblaC or ΔblaC only. Medium with or without 1 mg carbenicillin ml⁻¹ was added and the course of infection was monitored over 5 days. In the absence of carbenicillin, both strains grew in THP-1 cells during the course of the experiment. However, in the presence of carbenicillin, the strain expressing the truncated reporter alone did not grow and was reduced by 10-fold over 5 days while the strain expressing the exported ssPlcB-ΔblaC fusion protein was protected from carbenicillin and grew normally (Fig. 5a).

The ΔblaTEM-1 fusions were similarly tested. When THP-1 cells were infected with ΔblaC M. tuberculosis expressing either the exported ssMpt63-ΔblaTEM-1 or the ΔblaTEM-1 reporter alone, only the strain expressing ssMpt63-ΔblaTEM-1 fusion grew in THP-1 cells in the presence of carbenicillin. The non-exported ΔblaTEM-1 strain was sensitive to the β-lactam and was reduced in number by 10-fold (Fig. 5b). Similarly, ΔblaC M. tuberculosis exporting ssMpt83-ΔblaTEM-1 fusion was able to grow in carbenicillin-treated THP-1 cells, while the non-exported ΔblaTEM-1 construct did not confer resistance to the ΔblaC mutant (Fig. 5c).

These experiments demonstrated that both the Tat-specific ΔblaC reporter and the more permissive ΔblaTEM-1 reporter can report on protein export while M. tuberculosis is growing in β-lactam-treated host cells. The use of β-lactamase reporters with intracellular M. tuberculosis represents a powerful tool for the study and identification of proteins exported during growth in host cells.

**DISCUSSION**

The exported proteins of M. tuberculosis have been the subject of research attention for some time. This stems from the well-established fact that the majority of bacterial virulence factors and antigens are proteins exported out of the cytoplasm to the bacterial cell envelope or secreted out from the bacterium (Finlay & Falkow, 1997). In fact, there is a growing list of M. tuberculosis exported and secreted proteins shown to contribute to virulence or to development of a host immune response (Kurtz & Braunstein, 2005). Genetic reporters have proven to be powerful tools for identifying these extracytoplasmic proteins. The construction of a β-lactam-sensitive ΔblaC mutant of M. tuberculosis opened the door for using β-lactamases as reporters of protein export directly in M. tuberculosis. The ΔblaC reporter can be used as a Tat-specific reporter, while the ΔblaTEM-1 reporter, as shown here, can work with Sec or Tat signal sequences. An advantage of β-lactamase reporters is that they can be used to select for exported fusion proteins, as opposed to more labour-intensive
screening. In addition, we showed here for the first time that resistance to \( \beta \)-lactam antibiotics can be used to report on protein export during intracellular growth of bacteria. Even in more genetically tractable bacterial pathogens, the identification of proteins exported or secreted from within host cells is a challenge.

Because \( \beta \)-lactams target cell-wall-modifying enzymes, \( \beta \)-lactamases must be exported in order to protect against these drugs. This export requirement was previously exploited with fusion proteins expressed in \( E. \ coli \) and other bacteria grown in vitro (Broome-Smith et al., 1990; Lee & Hughes, 2006). Here we showed that BlaTEM-1 can also report on protein export directly in \( \Delta \)blaC \( M. \) tuberculosis. The three \( M. \) tuberculosis signal sequences tested in our study are from well-established secreted or cell-wall-associated proteins. Mpt63 (Rv1926c, 16 kDa protein) has a predicted Sec signal sequence and is one of the four most abundant \( M. \) tuberculosis proteins secreted into culture media during in vitro growth (Horwitz et al., 1995). Mpt83 (Rv2873) is a glycosylated lipoprotein (Hewinson et al., 1996; Sutcliffe & Harrington, 2004) that is exported to the cell wall of \( M. \) tuberculosis. Mpt83 has a predicted Sec signal sequence with a lipoprotein signal peptide (LspA) cleavage site and the requisite conserved cysteine for lipid modification. PlcB (Rv2350c, phospholipase C) is a cell-wall-associated protein of \( M. \) tuberculosis shown to function in virulence (Johansen et al., 1996; Raynaud et al., 2002). Unlike Mpt63 and Mpt83, PlcB has a predicted Tat signal sequence including a twin-arginine motif (Dilks et al., 2003). Signal sequences from all three of these proteins were able to promote export of a fused 'BlaTEM-1 reporter on the basis of production of \( \beta \)-lactam resistance. Notably, the ssMpt83-'BlaTEM-1 fusion protein was expressed from the native \( mpt83 \) promoter and the fusion protein included the predicted signal sequence plus 31 amino acids of the mature Mpt83 protein. This demonstrated the ability of the reporter to work with different strength promoters and extended protein sequences. It is important to note that even though variable levels of fusion protein were detected in \( M. \) tuberculosis whole-cell lysates as determined by immunoblot, each exported fusion provided sufficient protection against 50 \( \mu \)g carbenicillin ml\(^{-1} \) while the most abundant 'BlaTEM-1 without an export signal did not confer \( \beta \)-lactam resistance.

Previously, we showed that the PlcB signal sequence is able to drive export of functional 'BlaC in a Tat- and RR-dependent manner (McDonough et al., 2005). In \( E. \ coli \) the 'BlaTEM-1 reporter works with both Sec and Tat signal sequences (Broome-Smith et al., 1990; Stanley et al., 2002). The Sec and Tat pathways appear essential in \( M. \) tuberculosis (Braunstein et al., 2001; Saint-Joanis et al., 2006; Sassetti et al., 2003). Therefore, to investigate the mode of export of the ssPlcB-'BlaTEM-1 fusion protein, it was tested in \( M. \) smegmatis \( \Delta \)blaS and in a \( M. \) smegmatis \( \Delta tatA \Delta \)blaS double mutant. A 93% reduction in \( \beta \)-lactam-resistant colonies was observed in the \( M. \) smegmatis \( \Delta tatA \Delta \)blaS double mutant. Thus, the Tat pathway is involved in the export of ssPlcB-'BlaTEM-1, although other export pathways participate as well. The signal sequence of PlcB may be promiscuous in targeting the Tat or Sec pathway for export, depending on the folded or unfolded nature of a fused reporter element. Similar results were recently shown for some predicted Tat signal sequences in \( E. \ coli \) (Tullman-Ercek et al., 2007).

In addition to working with the Sec and Tat pathways, the 'BlaTEM-1 reporter has been used with type II and type III secretion systems of Gram-negative bacteria (Charpentier & Oswald, 2004; Sauvonnet & Pugsley, 1996). Since substrates of the type III secretion system lack conventional N-terminal signal sequences, it remains possible that the 'BlaTEM-1 reporter will also work with non-conventional exported proteins of \( M. \) tuberculosis.

An interesting category of exported proteins that has been largely overlooked are those proteins only expressed and/or exported during the course of infection. We hypothesize that these are proteins exclusively exported in the host environment, including virulence factors and protective antigens. Furthermore, only a small number of the exported \( M. \) tuberculosis proteins identified in vitro have ever been directly investigated during intracellular growth in host cells (Kurtz & Braunstein, 2005). For most of these studies, immunomicroscopy was used to localize the proteins in \( M. \) tuberculosis-infected macrophages, which required development of suitable antibodies. We reasoned that if \( \beta \)-lactam antibiotics can reach intracellular \( \Delta \)blaC \( M. \) tuberculosis, \( \beta \)-lactamase reporters should additionally work during intracellular growth. \( \beta \)-Lactam antibiotics do not normally accumulate in eukaryotic cells; however, antibiotics of this class freely diffuse in and out of host cells (Tulkens, 1991), and \( \beta \)-lactam antibiotics are used to treat some intracellular bacterial infections (Safdar & Armstrong, 2003). More specifically, \( \beta \)-lactams reach intracellular \( S. \) aureus and \( L. \) monocytogenes and prevent growth of these organisms in THP-1 cells (Barcia-Macay et al., 2006; Carryn et al., 2003). Here we showed that \( \Delta \)blaC \( M. \) tuberculosis in THP-1 cells was also susceptible to carbenicillin. Thus, BlaC is responsible for \( M. \) tuberculosis resistance to \( \beta \)-lactam antibiotics during intracellular growth, indicating that the chromosomal \( blaC \) is a key factor preventing the use of \( \beta \)-lactams to treat \( M. \) tuberculosis infection.

When the set of exported \( \beta \)-lactamase fusion proteins was tested for the ability to protect \( \Delta blaC \) \( M. \) tuberculosis from \( \beta \)-lactam treatment during intracellular growth, all exported fusions conferred resistance. In contrast, the truncated non-exported \( \beta \)-lactamase reporters were not protective. These experiments demonstrated the effectiveness of both 'BlaC and 'BlaTEM-1 reporters to identify \( M. \) tuberculosis sequences that drive export of each reporter during growth within host cells. Because the ssMpt83-'BlaTEM-1 fusion was expressed from the native promoter, our results indicate that Mpt83, a protein of unknown function, is expressed and exported during intracellular infection. This result is
consistent with the reported induction of mpt83 in macrophages (Schnappinger et al., 2003).

Several approaches have described proteins exported by M. tuberculosis in vitro, but a different suite of proteins may be exported during infection of the host. The intracellular β-lactamase reporter system we describe represents a new genetic tool for studying protein export in M. tuberculosis. It can be used to directly test the intracellular export of a protein of interest. We also hope to use it in combination with multiple rounds of infection and selection of β-lactam-resistant clones from a M. tuberculosis fusion library. This should serve to identify the most interesting category of proteins, namely those that are exported during intracellular growth and missed by alternative methods.

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Protein export by intracellular *M. tuberculosis*


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