Roles of LysM and LytM domains in resuscitation-promoting factor (Rpf) activity and Rpf-mediated peptidoglycan cleavage and dormant spore reactivation

Received for publication, April 21, 2020, and in revised form, May 15, 2020. Published, Papers in Press, May 20, 2020, DOI 10.1074/jbc.RA120.013994

Danielle L. Sexton, Francesca A. Herlihey, Ashley S. Brott, David A. Crisante, Evan Shepherdson, Anthony J. Clarke, and Marie A. Elliot

From the 1Michael G. DeGroote Institute for Infectious Disease Research and Department of Biology, McMaster University, Hamilton and the 2Department of Molecular and Cellular Biology, University of Guelph, Guelph, Ontario, Canada

Edited by Gerald W. Hart

Bacterial dormancy can take many forms, including formation of Bacillus endospores, Streptomyces exosporos, and metabolically latent Mycobacterium cells. In the actinobacteria, including the streptomycetes and mycobacteria, the rapid resuscitation from a dormant state requires the activities of a family of cell-wall lytic enzymes called resuscitation-promoting factors (Rpf). Whether Rpf activity promotes resuscitation by generating peptidoglycan fragments (muropeptides) that function as signaling molecules for spore germination or by simply remodeling the dormant cell wall has been the subject of much debate. Here, to address this question, we used mutagenesis and peptidoglycan binding and cleavage assays to first gain broader insight into the biochemical function of diverse Rpf enzymes. We show that their LysM and LytM domains enhance Rpf enzyme activity; their LytM domain and, in some cases their LysM domain, also promoted peptidoglycan binding. We further demonstrate that the Rpf enzymes function as endo-acting lytic transglycosylases, cleaving within the peptidoglycan backbone. We also found that unlike in other systems, Rpf activity in the streptomycetes is not correlated with peptidoglycan-responsive Ser/Thr kinases for cell signaling, and the germination of rpf mutant strains could not be stimulated by the addition of known germinants. Collectively, these results suggest that in Streptomyces, Rpf have a structural role rather than signaling function during spore germination, and that in the actinobacteria, any signaling function associated with spore resuscitation requires the activity of additional yet to be identified enzymes.

A major constituent of the cell wall, in both vegetative and dormant cells, is peptidoglycan. Peptidoglycan polymers are defined by their glycan backbones, composed of alternating N-acetylmuramic acid (MurNAc) and N-acetylatedglucosamine (GlcNAc) residues, and by short peptides extending from the lactyl groups of the MurNAc residues. The peptide stems of different glycan chains can in turn be joined together either directly, or by amino acid linkers of varying lengths. These peptide bridges cross-link parallel strands together, yielding a rigid structure that maintains the integrity of the cell membrane (4). In dormant cells, the peptidoglycan is relatively inert, whereas in actively growing cells, it is highly dynamic (5).

Cell wall cleavage is a critical component of cell growth, being required for the insertion of new peptidoglycan. Muralytic enzymes target the glycan strands of peptidoglycan, and are classified as either hydrolases or lytic transglycosylases. Hydrolases, including the lysozymes and β-N-acetylmuraminidases, hydrolyze β-(1–4) linkages in the glycan strands (6). In contrast, lytic transglycosylases cleave the same bond as lysozymes (between MurNAc and GlcNAc), but they do not require water and instead generate GlcNAc and 1-6-anhydroMurNAc products (7). All of these enzymes can be further subdivided into exo- or endo-acting enzymes, depending on whether they cleave at the ends of glycan strands, or within strands, respectively.

For many dormant cells, a return to active growth requires the breakdown of the thick protective cell wall, and different bacteria have evolved distinct strategies to achieve this. Within the actinobacteria, dormant cells employ a common degradative enzyme that promotes the resumption of vegetative growth. The so-called “resuscitation promoting factor” (Rpf) enzymes share structural homology with lysozyme and lytic transglycosylases (8), and have muralytic activity (9–15). In Micrococcus luteus, a single Rpf enzyme is required for the resuscitation of metabolically quiescent cells (16). Most other actinobacteria encode multiple Rpf enzymes (17, 18), and these collectively stimulate the growth of dormant cells (13, 19–22). In Streptomyces coelicolor, the products of five rpf genes (rpfA-E) promote the rapid germination of dormant spores, and can influence both vegetative growth and sporulation (11, 13). Deleting individual rpf genes results in modest germination deficits in some instances.
Bacteria, Total 2493 1674 335 7129
Rpf, VCBS 26 (1%) 0 0 26 (0.4%)
Rpf, Peptidase, SLT/GEWL 11 (0.4%) 0 0 11 (0.1%)
9172
(11, 13), whereas the loss of all five has the greatest impact on
germation (13).

Resuscitation is a complex process, and how the Rpfs promote resuscitation is not fully understood. Two models have been put forth to explain Rpfl function during the escape from dormancy: 1) Rpfl activity liberates peptidoglycan-derived signaling molecules that activate a regulatory cascade needed for growth resumption, and 2) Rpfl activity relieves the physical constraints imposed by dormant cell walls, allowing cell growth to resume (23). Although these proposals are not mutually exclusive, investigations to date appear to favor a signaling-based mechanism (24).

Resuscitation from dormancy has been best studied in Bacillus, which forms highly resistant endospores (25, 26) and encodes an Rpf-like enzyme (27). Bacillus spore germination can be promoted by the addition of peptidoglycan fragments (muropeptides) (28). These muropeptides bind to PrkC, a eukaryotic-like Ser/Thr kinase located in the spore membrane, initiating a signaling cascade that triggers spore germination (28). PrkC contains tandem PASTA (penicillin-binding protein and Ser/Thr kinase associated) domain repeats, and these domains recognize both nascent peptidoglycan and muropeptides (28–32). A similar situation may exist in Mycobacterium, where emergence from latency can be stimulated by muropeptide binding to PknB, a PrkC homologue (32). It is worth noting, however, that mycobacterial resuscitation via this route is not robust, and the major function of muropeptide binding appears to be in directing the subcellular localization of PknB (32). The molecular basis for Streptomyces resuscitation, and the contribution made by the Rpfl proteins to this process, remains to be determined.

There is considerable diversity in Rpfl enzyme architecture, and a clear understanding of Rpfl function requires not only a full characterization of the enzymes themselves, but also a systematic assessment of the contributions made by the different domains. Here, we show that Rpfl accessory domains make critical contributions to enzyme activity. We establish that the Rpfl function as endo-acting lytic transglycosylases, and further demonstrate that their activity is independent of known signaling cascades associated with germination in other systems. Unlike most systems investigated to date, our data are most consistent with a cell wall remodeling role for the Rpfls in Streptomyces spore germination.

Results
Rpf domain diversity in the actinobacteria

The Rpfl domain is found in proteins throughout the actinobacteria, in association with a variety of different protein domains (17, 18). How these accessory domains influence the biological and biochemical function of different Rpfls remains unclear. To prioritize different architectures for investigation, we searched for Rpfl domain-containing proteins in the streptomycetes, mycobacteria, micrococci, and other actinobacteria (Table 1). We found the RpfASC (protein bearing a signal peptide, and Rpfl and LysM domains) class to be the most widespread in the actinobacteria, followed closely by the short Rpfl class, which have no obvious functional domains beyond their Rpfl domain (and signal peptide).

There were interesting phylogenetic distributions associated with each of the Rpfl domain architectures. Within the mycobacteria, Rpfl domains were most frequently found in conjunction with uncharacterized N- or C-terminal extensions (17, 18) (Table 1). These extended regions lacked any obvious functional domains, and were confined to Rpfl-associated proteins in the mycobacteria. The corynebacteria also encoded a distinct subset of Rpfl proteins associated with an uncharacterized DUF3235 domain (17) (Table 1).

In addition to these Genus-specific subsets, Rpfls were also associated with other functional domains, with two configurations being highly represented: the RpflB subgroup and the LysM-containing groups (Table 1). Members of the RpflB group were found in a range of actinobacterial species, and contained a G5 domain and tandem repeats of the DUF348 domain. Recent structural studies on an Rpfl variant from Mycobacterium tuberculosis revealed an interesting ubiquitin-like fold for the DUF348 domain, and a close physical association between these domains and the G5 domain (33, 34). G5 domains bind to GlcNAc residues and are thought to promote peptidoglycan binding (35), whereas the DUF348 domains facilitate Rpfl dimerization, and appear to negatively affect Rpfl cleavage activity (13). In contrast to the RpflB group, the LysM-containing groups of Rpfls are the predominant form in the streptomycetes.
and micrococci. The LysM domain, like the G5 domain from the RpfB subfamily, binds GlcNAc residues (36), and is proposed to enhance Rpf binding to its peptidoglycan substrate. In the streptomycetes, many LysM domain-containing Rpfs also possess a LytM domain (Pfam: M23 metallopeptidase), which is expected to have endopeptidase activity and thus the potential to cleave either within peptide stems or peptide cross-bridges (37, 38).

LysM and LytM domains enhance Rpf activity

There is currently nothing known about how the Rpf-associated LysM and LytM domains influence Rpf activity. Given that the vast majority (>85%) of Streptomyces Rpf proteins possess one or both of these domains (Fig. 1A), we sought to determine how they influenced the biochemical activity of the Rpfs. To probe the functional contributions made by these domains, we created a truncated version of RpfA lacking the LysM domain (RpfAΔLysM), alongside two RpfD variants: one missing the LysM domain (RpfDΔLysM), and one lacking both the LysM and LytM domains (RpfDΔLysMΔLytM). We overexpressed and purified these proteins, along with their full-length counterparts (minus their SignalP-predicted secretion signals (39)), and evaluated the enzyme activity of each using a fluorescence-based peptidoglycan cleavage assay. The assay employs fluorescein-labeled M. luteus peptidoglycan as a substrate, where the fluorescein labeling is sufficiently dense so as to quench the fluorescent signal. Peptidoglycan cleavage results in the release of fluorescein molecules, leading to increased fluorescence.

Mature versions of the full-length and truncated RpfA and RpfD enzymes were added in equimolar concentrations to the fluorescein-labeled substrate. For both RpfA and RpfD, we found that enzymes lacking the LysM domain had 65–70% of the activity of the full-length variants (Fig. 1B). This suggested that peptidoglycan targeting by the LysM domain may help position the Rpfs on their substrate and enhance their cleavage capabilities. We observed that removal of the LytM domain from RpfD led to a further decrease in activity; RpfD lacking both LysM and LytM domains had only ~30% of the activity of the full-length enzyme (Fig. 1B).

The contribution of the LytM domain to RpfD activity may be enzymatic, as this domain typically has metallopeptidase activity, or it could function as an additional substrate specificity determinant. LytM peptidase activity requires a Zn^{2+} co-factor (40), and thus we tested the activity of all RpfD variants in the presence and absence of EDTA, which would be expected to chelate any associated Zn^{2+} ions. We found that EDTA had no effect on RpfD activity, irrespective of whether the LytM domain was present or not (Fig. 1C). This suggested that the RpfD-associated LytM domain may not function as an enzyme, and may instead provide additional targeting specificity for...
RpfD; although we cannot formally exclude the possibility that 10 mM EDTA was insufficient to remove any associated metal ions from RpfD. We examined the sequence of the LytM domain to determine whether it was lacking any key Zn$^{2+}$-binding or active site residues, as is the case for EnvC and NlpD in Escherichia coli (41). However, all critical residues appeared to be present (Fig. S1), suggesting that the lack of enzyme activity was not due to the inability to bind the Zn$^{2+}$ co-factor, nor to a degenerate active site.

To determine whether the difference in activity was due to a reduced ability to bind to peptidoglycan, we performed peptidoglycan-binding assays using peptidoglycan isolated from Strep-tomyces coelicolor. Surprisingly, we found that removing the LysM domain from Rpfa had little impact on the peptidoglycan-binding capabilities of this enzyme (Fig. 1D, Fig. S2). In contrast, the loss of the LysM domain from RpfD significantly reduced peptidoglycan binding. Removing the LytM domain from the LysM-deficient RpfD further reduced peptidoglycan binding (Fig. 1D, Fig. S2), supporting the proposal that the LytM domain may function to enhance peptidoglycan binding by RpfD.

**Rpf domain functions as an endolytic transglycosylase**

Having established that the LysM and LytM domains impacted Rpf activities in vitro, we next set out to investigate the mechanistic basis underlying peptidoglycan cleavage by the various Rpf enzymes. We opted to assess the activity of all five RpfS from S. coelicolor (Rpfa–E), as these enzymes represented four different structural classes (Rpf alone, RpfB, RpfLysM, and RpfLysM+LytM). For this, we used the assay of Herlihey et al. (42), taking advantage of the fact that hydrolyses require water to break their cognate glycosidic bonds. Thus, in the presence of water labeled with the stable isotope $^{18}$O, hydrolyse products would have an $^{18}$OH incorporated at the C-1 position, altering the isotopic distribution of products detected using MS, relative to those produced in the presence of unlabeled water. In contrast, lytic transglycosylases do not use water in breaking the glycosidic bond in the peptidoglycan backbone, and as a result their 1,6-anhydromuroglycan products would have an unaltered isotopic distribution.

*M. luteus* peptidoglycan, suspended in $^{18}$O-labeled water, was initially used as a substrate for Rpfa–E, and for the control hydrolytic enzyme mutanolysin. Being a highly active and efficient hydrolyse, mutanolysin completely solubilized the peptidoglycan and we detected a variety of muroglycans by LC–MS analysis (Fig. 2). As expected, a number of these muroglycans were enriched with $^{18}$O. Subsequent MS/MS analyses confirmed the association of this $^{18}$O with only muramoyl residues (Fig. 3). In contrast, we detected very few soluble muroglycan products from each of the reactions involving Rpfa, Rpfb, RpfC, and RpfE, whereas none were produced by RpfD (Fig. 2A). MS analyses indicated that the few soluble muroglycans released did not contain $^{18}$O. These data suggested that the RpfS did not function as either muramidases or B-N-acetylglucosaminidases. Instead, tandem MS analyses revealed that the released soluble muroglycans contained GlcNAc-1,6-anhydro-MurNAc (peptides) (Table 2), indicating that each of RpfA/B/C/E functioned as lytic transglycosylases.

We wondered if the lack of soluble products following reaction with RpfD, and the minimal amount produced by the other Rpf proteins, was due to a predominant endo-type lytic activity associated with each, where reaction products would remain cross-linked to the insoluble peptidoglycan sacculus. To analyze the insoluble fraction for any evidence of lysis, we washed the recovered insoluble peptidoglycan products and digested them with mutanolysin; any initial hydrolytic products of Rpf activity would retain their $^{18}$O enrichment, if present, following this secondary digestion. As seen in Fig. 2B, the muroglycan profiles of peptidoglycan incubated with each of the RpfS, including RpfD, followed by mutanolysin digestion were distinct from the control reaction with mutanolysin alone. Tandem MS analysis of the unique muroglycan fractions revealed that none were enriched with $^{18}$O and that the majority were linear oligomers terminating with an anhydromuramoyl residue (Table 2). These data thus suggested that each of the RpfS activity as endo–acting lytic transglycosylases.

Unexpectedly, the muroglycan profiles for each of the five RpfS were similar, and any specificity for glycan chain length, peptide stem composition, or cross-linking was not observed in the soluble or insoluble fractions. We noted, however, that the peptidoglycan composition of *M. luteus* differed slightly from that of *S. coelicolor*, specifically in the third amino acid and interpeptide bridge positions (43). Consequently, we sought to test whether any differences could be detected when using *S. coelicolor* peptidoglycan as substrate. As we saw for the cleavage profiles of the five WT RpfS using *M. luteus* peptidoglycan, the *S. coelicolor* muroglycan profiles for Rpfa, RpfD, and their LysM-LytM mutant derivatives were all similar (Fig. 4, Table 3), although the relative intensity units were different for the two substrates, suggesting greater Rpf affinity for/activity against the *S. coelicolor* peptidoglycan. These data collectively suggested that the different domains associated with the Rpf proteins did not confer any obvious substrate specificity with respect to peptidoglycan cleavage. Given the potential peptidase activity associated with the LytM domain of RpfD, we closely examined the cleavage products of both the *M. luteus* and *S. coelicolor* peptidoglycan for evidence of any hydrolytic activity not associated with the muroglycan backbone, but none were detected. These findings were consistent with our in vitro analyses, suggesting that the LytM domain enhanced RpfD function, not through peptidase activity, but instead by promoting peptidoglycan binding.

**PASTA domain-containing Ser/Thr kinases in *S. coelicolor* inhibit germination and vegetative outgrowth**

One hypothesis put forward to explain the role of RpfS in cell resuscitation involves the release of muropeptide signals, which activate a regulatory cascade leading to the reactivation of metabolism. Such a model would be most consistent with exo-activity of the RpfS, as this would promote the release of muropeptides; however, our results indicated that the RpfS were endo-acting lytic transglycosylases.

In *Bacillus*, and to a lesser extent in *Mycobacterium*, the resuscitation-promoting signaling cascade is mediated through PASTA domain-containing Ser/Thr kinases (28, 32). We
considered two possibilities that could accommodate both the endo-activity of the Rpfs and a role for Ser/Thr kinase signaling. In one, the PASTA domain-containing kinases in *S. coelicolor* may recognize the ends of cleaved peptidoglycan rather than a defined muropeptide. The second involved Rpf-cleaved products serving as a substrate for other cell wall lytic enzymes, resulting in the release of germination-stimulating muropeptides that are recognized by these kinases.

*S. coelicolor* encodes three PASTA domain-containing protein kinases, and we obtained a triple mutant strain (44), here dubbed the 3ΔPASTA strain (Table S1). We expected that this strain would have similar germination rates to that of an *rpf* null mutant if the Rpfs were involved in generating appropriate peptidoglycan ends or germination-promoting muropeptides that were recognized by these kinases. We measured germination rates of the triple mutant strain, and compared these to the WT and *rpf* null strains. We found that germination of the 3ΔPASTA strain was consistently more rapid than either the WT or the *rpf* null strain (Fig. 5A). This suggested that the three PASTA domain-containing Ser/Thr kinases in *S. coelicolor* were not involved in recognizing a product produced directly or indirectly by the Rpfs. Instead, the rapid germination of these strains implied that the activity of these kinases might inhibit germination. We also assessed the growth of the 3ΔPASTA strain in liquid minimal medium, to determine whether it exhibited defects in vegetative growth compared with WT and the *rpf* null strain. Consistent with our germination results, growth of the 3ΔPASTA strain was faster than either comparator strain (Fig. 5B). These results suggested that these Ser/Thr kinases may function to delay germination/growth, given the enhanced rates of both processes in the absence of these enzymes, and further indicated that the effect of the Rpfs on spore germination and vegetative growth was not a result of Rpf-dependent muropeptide signaling, at least through the PASTA domain-containing Ser/Thr kinases.

**Figure 2. Characterization of RpfA-E as endo-lytic transglycosylases by LC-Q-TOF MS analysis of their reaction products.** Samples of *M. luteus* peptidoglycan suspended in [18O]H2O to a final concentration of 1.4 mg/ml were incubated separately with 1 nmol Rpf or 1.1 nmol mutanolysin (positive control). After incubation at 37 °C for 9.5 h, soluble reaction products were separated from insoluble material by centrifugation. The insoluble peptidoglycan pellets from the Rpf digestions were washed with water and then resuspended in 0.1 mM potassium phosphate buffer, pH 6.2, for solubilization by 1.1 μM mutanolysin. Soluble muropeptides from this secondary digestion were recovered by centrifugation. Each soluble and secondary-soluble fraction was subjected to LC-Q-TOF MS analysis. A, analysis of soluble fraction from peptidoglycan alone (PG), and reactions with 1 nmol Rpf–A–E, as indicated. B, analysis of insoluble products following secondary mutanolysin digestion from reaction with 1.1 μM mutanolysin (Mut; positive control); or 1 nmol Rpf–A–E, as indicated. The identities of the numbered muropeptide fractions are listed in Table 2. The solid vertical bar to the left denotes 10,000 and 200,000 intensity units, for panels A and B, respectively.
Rpf activity is required for germination with alternative germinants

An alternative hypothesis to explain how Rpf enzymes promote germination is that their cell wall cleavage activities provide cells with an opportunity to insert new peptidoglycan, thus permitting cell expansion and growth. We predicted that if the role of the Rpf was a physical one, we should be able to stimulate germination of the WT strain, but not an \textit{rpf} mutant, by adding a known germinant. To test this, we incubated spores on minimal medium in the presence or absence of germination-promoting calcium chloride (45). We followed germination over a 7-h time course, and found that calcium chloride effectively stimulated germination of WT spores, but had little effect on the \textit{rpf} null strain (Fig. 6). This supported the proposal that

\begin{figure*}[h]
\centering
\includegraphics[width=\textwidth]{figure3.png}
\caption{Tandem Q-TOF MS analysis of select muropeptides. Example of MS analysis of parent ions for muropeptides recovered from (A) mutanolysin (positive control) and \textit{B} soluble RpfA digests of \textit{M. luteus} peptidoglycan, by LC–MS as described in the legend of Fig. 2, \textit{C} and \textit{D} tandem Q-TOF MS analysis of denoted parent ions from corresponding panels \textit{A} and \textit{B}. The blue spectral line in the MS spectrum of panel \textit{A} denotes the \textit{18}O-containing isotope of the respective muropeptide. The monoisotopic masses \textit{(M + 2H)}$^{+}$ are presented for each of the identified fragments.}
\end{figure*}

\begin{table}[h]
\centering
\caption{LC-Q-TOF analysis of select muropeptides released from insoluble \textit{M. luteus} peptidoglycan by RpfA-E}
\begin{tabular}{|l|l|l|l|l|l|l|l|}
\hline
Fraction No. & Annotation & RpfA & RpfB & RpfC & RpfD & RpfE & \textit{m/z} \\
\hline
1 & G-anhM(Penta) & 468.2050 & 468.2131 & 468.2112 & 468.2103 & NA & 468.2121 & 2 \\
2 & G-anhM(Penta-Ala) & 503.7250 & 503.7322 & 503.7326 & 503.7330 & NA & 503.7309 & 2 \\
\hline
3 & G-anhM(Penta) & 468.2050 & 468.2118 & 468.2117 & 468.2120 & 468.2137 & 468.2123 & 2 \\
4 & G-anhM(Penta-Ala) & 503.7250 & 503.7300 & 503.7309 & 503.7330 & 503.7320 & 503.7306 & 2 \\
5 & G-M-G(Penta)-G-anhM & 616.6200 & 616.9644 & 616.6293 & 616.9655 & 616.6328 & 616.6308 & 3 \\
6 & G-M-G(Penta)-G-anhM(Penta) & 782.0300 & 782.O & 782.O & 782.O & 782.O & 782.O & 69 & 3 \\
7 & G-M-G-G-anhM(Penta-Ala) & 979.9350 & 979.9418 & 979.9393 & 979.9386 & 979.9434 & 979.9394 & 2 \\
\hline
\textsuperscript{a}The muropeptide fractions correspond to those of the RP-HPLC separation presented in Fig. 2, \textit{A} and \textit{B}. \\
\textsuperscript{b}Identification of each muropeptide was made by tandem Q-TOF-MS analysis of each parent ion (data not shown); G, GlcNAc; M, MurNAc; anhM, 1,6-anhydroMurNAc; penta, L-Ala-D-Glu-(Gly)-L-Lys-D-Ala; *, O-acetylation. \\
\textsuperscript{c}NA, not observed.
\end{tabular}
\end{table}

\textit{Rpf} activity is required for germination with alternative germinants

An alternative hypothesis to explain how Rpf enzymes promote germination is that their cell wall cleavage activities provide cells with an opportunity to insert new peptidoglycan, thus permitting cell expansion and growth. We predicted that if the role of the Rpf was a physical one, we should be able to stimulate germination of the WT strain, but not an \textit{rpf} mutant, by adding a known germinant. To test this, we incubated spores on minimal medium in the presence or absence of germination-promoting calcium chloride (45). We followed germination over a 7-h time course, and found that calcium chloride effectively stimulated germination of WT spores, but had little effect on the \textit{rpf} null strain (Fig. 6). This supported the proposal that...
germination was inhibited at the outgrowth step in the rpf null strain (although we cannot exclude the possibility that calcium promotes germination by stimulating Rpf activity).

We also tested the effect of calcium chloride on the 3-3D PASTA mutant strain, and found that like WT, spore germination was enhanced in the presence of calcium chloride (Fig. 6). This further implied that the activity of the PASTA-domain–containing Ser/Thr kinases was independent of Rpf activity in S. coelicolor.

**Discussion**

Here, we demonstrated that Rpfs were endo-acting lytic transglycosylases. We found that Rpf function was enhanced by their associated domains, with LytM domains appearing to promote peptidoglycan association but not peptidoglycan cleavage, and LysM domains increasing enzyme activity, either by enhancing peptidoglycan binding, as appears to be the case for RpfD, or by some other means, in the case of RpfA. Unexpectedly, we found Rpf function was not tied to an obvious signaling cascade in the streptomycetes. Instead our data were most consistent with a physical role for the Rpfs, where these enzymes functioned to structurally alter the germinating spore wall.

**Role of LysM and LytM domains in Rpf activity**

LysM-containing Rpfs represent one of the largest Rpf protein configurations in the actinobacteria, but the role of LysM domains in cell wall lytic enzymes is not universally conserved: some enzymes require these domains for activity, whereas others do not (46–49). We found that for RpfA and RpfD, deleting their LysM domains decreased their cell wall lytic activity. Interestingly, however, this domain only seemed to promote peptidoglycan association for RpfD, not RpfA, at least in vitro. LysM domains do not have catalytic activity of their own, and thus we suggest that the LysM domain may function to either increase the affinity of Rpf proteins for peptidoglycan (as for RpfD), or perhaps impact substrate orientation relative to the Rpf catalytic domain (RpfA). In their natural environments, LysM domains may further serve to anchor the Rpfs to the cell wall, thereby preventing indiscriminate peptidoglycan cleavage by these enzymes, and ensuring that the Rpfs remain a
the mycolic acid-based outer membrane encasing the peptidoglycan in these organisms may prevent the broad dispersal of the Rpfs, obviating the need for this additional LysM domain.

In contrast to the LysM domains, LytM domains are expected to have catalytic activity, and function in cleaving stem/cross-bridge peptides. Intriguingly, the RpfD-associated LytM domain does not appear to have peptidoglycan cleavage capabilities, at least in the assays conducted here. Despite the lack of LytM enzyme activity, removing the LytM domain, together with the LysM domain, significantly reduced the activity and peptidoglycan binding of RpfD beyond what had been observed by simply deleting the LysM domain. This suggests that the LytM domain further enhanced the affinity of RpfD for peptidoglycan through its peptide binding, although it is also possible that this domain increased Rpf enzyme activity (and/or peptidoglycan binding) through allosteric activation, as has been observed for EnvC and NlpD in their activation of AmiA, AmiB, and AmiC in *E. coli* (41).

Unlike EnvC and NlpD, however, the RpfD LytM domain has retained all active site and Zn$^{2+}$-binding residues, suggesting that it may still be enzymatically competent. Some LytM metallopeptidases require additional processing for activation (40, 50). Such processing may occur upon secretion of RpfD to the *Streptomyces* cell surface, where the LytM domain then makes an enzymatic contribution to RpfD activity. Alternatively, this domain may be functionally silent in the context of the RpfD polypeptide, but may be processed in such a way that it acts independently of RpfD. RpfD is unusual among the Rpfs in *S. coelicolor*, in that *rpfD* transcript levels expression peaks later in development, as opposed to during germination as is the case for all other *rpf* genes with detectable transcription (13). It is therefore possible that RpfD function, and that of its LytM domain, may be more important at later stages of development than the other Rpfs in *S. coelicolor*. This would be consistent with the observation that LytM-containing Rpf proteins are found exclusively in the streptomycetes, and thus may function in aspects of development unique to these bacteria.

**Specificity and redundancy in Rpf function**

A key question is why multiple Rpfs are required for cellular resuscitation in many actinobacteria (13, 21). Our peptidoglycan cleavage assays did not reveal unique specificity for any individual Rpf class, although it is notable that the muropeptides released during cleavage of *S. coelicolor* represented those that are most abundant in spore peptidoglycan (the relative proportion of different monomer and dimer products changes throughout development) (51). At this stage, however, we cannot exclude the possibility that these enzymes have differential specificity in vivo. Germination of *Streptomyces* spores occurs at the spore poles, and it is possible that the peptidoglycan architecture in these regions has specific modifications that were not captured at high levels in our peptidoglycan preparations. It is also possible that other proteins function to localize the Rpfs to specific sites within the cell wall of dormant spores. Future protein interaction and localization studies will help to resolve these questions.
Revising the model of Rpf function during germination

How the RpfS promote resuscitation/germination is still being debated, 20 years after the discovery of these proteins. Do they function to liberate a signaling molecule that acts as a germinant? Or is their role more structural, where they permit cell expansion and new peptidoglycan incorporation through their cell wall cleavage activities?

Three lines of evidence support a peptidoglycan remodeling role for the *Streptomyces* RpfS. One, we demonstrated that peptidoglycan-binding kinases, known to influence germination through a muropeptide-mediated signal transduction cascade in *Bacillus* (28) and to a lesser extent in *Mycobacterium* (32), are not associated with Rpf function in *S. coelicolor*. Instead, these kinases appear to negatively influence germination, based on the rapid germ tube outgrowth observed in their absence. Two, the endo-acting lytic transglycosylase activity of the RpfS is more compatible with an architectural role than with a signaling role. Finally, a known germinant for *Streptomyces* (calcium chloride) stimulated germination of WT spores, but had no effect on the germination of an rpf null strain. The fact that alternative germinants could not substitute for the lack of RpfS, suggests that the RpfS may be universally required for efficient germination. It is, however, worth noting that it is not clear how calcium chloride promotes germination, and it will be interesting to determine whether its effects are mediated through Rpf activity, through other cell wall lytic enzyme(s) that act in conjunction with the RpfS, or through some other means altogether.

In *M. tuberculosis*, an equivalent germination-promoting experiment (treating dormant cells with both an Rpf inhibitor and oleic acid, a known germinant) led to metabolic reactivation but delayed cellular outgrowth, again suggesting that the Rpf role may be more structural (52, 53). Taken together, the simplest explanation for these results would be that the RpfS function to remodel the cell wall and promote cell expansion and growth after metabolic reactivation.

Our findings do not, however, definitively rule out an additional signaling role for the RpfS. Indeed, a wide variety of muropeptides clearly enhance the resuscitation of *Mycobacterium* (24, 29, 54). In *M. tuberculosis*, RpfB acts synergistically in association with the endopeptidase RpfA, and a product of their activity (a peptidoglycan-derived disaccharide-dipeptide) has been proposed to promote mycobacterial resuscitation (10, 54). In the streptomycetes, such a signaling phenomenon would require additional glycosidic enzymes, as the endo-lytic activity of the RpfS would not allow for the generation of a disaccharide molecule. We suggest that muropeptide release may be a secondary effect of Rpf activity, and this could be accomplished either directly through the cleavage activity of RpfS and any associated enzymes, or indirectly through cell growth and the associated peptidoglycan shedding that accompanies this process.

**Experimental procedures**

**Bioinformatic analysis**

The HMMER webserver (55) was used to search the UniProtKB database using the Rpf domain from RpfE in *S. coeli-
color* to identify homologues of each Rpf configuration and their taxonomic distribution.

**Bacterial strains and growth conditions**

Bacterial strains used or created in this work are outlined in Table S1. *S. coelicolor* A3(2) strain M145 and its derivatives were grown at 30°C on solid minimal medium (MM) or mannitol soya flour (MS) agar, with antibiotics to maintain plasmid selection where appropriate, or in new minimal medium with phosphate, as described by Kieser et al. (56). Growth curves were generated using dry weight. This involved transferring 1 ml of culture at a given time point, into pre-weighted tubes. The cells were then collected by centrifugation at full speed in a microcentrifuge for 5 min, after which the supernatant was removed. The remaining cell pellet was then dried at 60°C for ~2 days, after which the tubes were weighed again, allowing for calculation of the cell dry weight. All *E. coli* strains were grown at 37°C on LB or nutrient agar (NA) plates (56), or in LB or super optimal broth liquid medium (57, 58) supplemented with antibiotics where appropriate to maintain plasmid selection.

**Spore germination assay**

To assess the germination efficiency of the different strains, spores were plated on MS agar overlaid with cellophane discs and incubated at 30°C for up to 8 h. At the indicated time points, a 1 × 1-cm square was excised from the cellophane disc and examined using light microscopy. Images were acquired at ×1000 magnification using a Nikon Eclipse E600 microscope fitted with DS-Fi1 camera. Image capture was performed using Nikon NIS-Elements software. Spore germination was then assessed, scoring germinated spores (those possessing at least one germ tube) *versus* non-germinated spores, with a minimum of 200 spores being assessed per strain, at each time point, in at least three independent trials. Spore scoring was performed using the cell counter plug-in for ImageJ (59). To test the effects of Ca²⁺ on germination, spores were plated on MM agar with and without 10 mM CaCl₂. Spore germination assays were then conducted as described above.

**Protein overexpression and purification**

RpfSAs–E, excluding their SignalP-predicted signal peptide, were overexpressed and purified as N terminally His-tagged proteins, as described by Sexton *et al.* (13). The equivalent region for each RpfA and RpfD variant (coding sequence, minus signal peptide) was amplified using the primers outlined in Table S2. Overlap extension PCR (60) with the primers described in Table S2 was used to generate rpfDΔLysM. Other mutants were truncations of either RpfA or RpfD, and were generated using the primers indicated in Table S2. Digested PCR products were cloned into the BamHI and Ndel restriction sites of pET15b (Novagen) (Table S1). Construct integrity was confirmed by sequencing using the T7 promoter and terminator primers (Table S2). Each plasmid was freshly transformed into *E. coli* Rosetta 2 cells (Table S2) prior to overexpression. Transformants were grown overnight at 37°C in 5 ml of LB liquid medium supplemented with ampicillin and chloramphenicol. These overnight cultures were used to inoculate 500 ml of
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LB medium, again supplemented with ampicillin and chloramphenicol. Cultures were grown at 37 °C until they reached an optical density at 600 nm (OD600) of 0.6-1.0 (depending on the Rpf variant), at which point 1 mM isopropyl β-D-thiogalactopyranoside was added to induce protein overexpression. Conditions for overexpression are summarized in Table S3. Overexpression of RpfDΔLysM was attempted at an initial OD600 of 0.4-1.2, using 0.25-2 mM isopropyl β-D-thiogalactopyranoside, and induced cultures were grown for 1.5 h to overnight at 16, 30, or 37 °C. Overexpression was also attempted using in vitro translation with the PURExpress kit (New England Biolabs) following the manufacturer’s recommendations. None of the conditions tested yielded the desired protein.

For those proteins where overexpression was observed, cell pellets were resuspended in 5 ml of lysis buffer (50 mM NaH2PO4, 300 mM NaCl, 10 mM imidazole, pH 8.0) containing cComplete Mini EDTA-free protease inhibitor (Roche Applied Science) and lysed using the Constant Systems TS-2 0.75 kW cell disruptor. The lysate was centrifuged at 10,000 × g for 20 min at 4 °C to remove insoluble debris. The clarified lysate was incubated with 1 ml of nickel-nitriolactose acid-agarose (Thermo) for 1 h at 4 °C before being applied to a chromatography column. The column was washed twice with 5 ml of buffer (50 mM NaH2PO4, 300 mM NaCl, pH 8.0) containing 20 and 50 mM imidazole before His-tagged proteins were eluted sequentially with buffers containing 100 mM, 250 mM, 500 mM, and 1 mM imidazole. The success of protein overexpression and the quality of protein purification was assessed by separating purified proteins (and their accompanying washes and crude soluble and insoluble fractions) on a 10% Tricine polyacrylamide gel (61) and staining with Coomassie Brilliant Blue. Protein concentrations were determined using a Bradford assay (62), with BSA as a standard. Each Rpf was dialyzed into storage buffer (50 mM NaH2PO4, 10% glycerol, pH 8) overnight to remove imidazole. Proteins were stored at 4 °C for a maximum of 24 h before all assays.

**Enzyme activity assays**

Quantitative Rpf activity assays—The EnzChek lysozyme assay kit (Molecular Probes) was used to assess the ability of the different Rpfs and RpfD variants to cleave fluorescein-labeled M. luteus cell walls, as described previously (13). Briefly, 1 nmol of purified Rpf protein was added to each reaction, and the volume was brought to 50 μl with storage buffer before adding 50 μl of fluorescein-labeled M. luteus cell wall substrate. One pmol of lysozyme was used as a positive control, whereas a reaction without protein served as a negative control. Reactions were set up in black 96-well-plates (Thermo). Fluorescein release was measured after 1 h using a Cytation 5 plate reader (BioTek) with an excitation wavelength of 494 nm and emission wavelength of 521 nm. Assays were conducted in technical triplicates, using at least two independent protein preparations.

Isolation and purification of peptidoglycan—Insoluble peptidoglycan for use in the enzymatic assays was isolated from overnight cultures of S. coelicolor using the boiling SDS protocol and purification by enzyme treatment (amylase, DNase, RNase, and Pronase), as described by Clarke (63); as Gram-positive bacteria, both M. luteus and S. coelicolor produce peptidoglycan with limited 1,6-anhydromuramoyl content (64). O-Acetyl groups were removed by incubating peptidoglycan in 20 mM NaOH at room temperature overnight, and insoluble peptidoglycan was isolated by centrifugation (9,000 × g, 30 min, room temperature) and washed with water at least three times. Teichoic acids were removed by extracting the peptidoglycan with 10% TCA overnight at room temperature and peptidoglycan was washed four times in water, frozen, lyophilized, and stored at −20 °C.

**Peptidoglycan-binding assays**—A solution containing 1 mg/ml of purified S. coelicolor peptidoglycan in 10 mM Tris, pH 7, was sonicated continuously on ice for 5 min (to provide a relatively even suspension). Two nanomoles of purified Rpf protein were mixed with 150 μl of the peptidoglycan solution and the volume was brought to 300 μl using 10 mM Tris, pH 7. Reactions were incubated at 4 °C for 2 h with gentle shaking. The peptidoglycan-bound protein was separated from unbound protein by centrifugation at 15,000 × g at 4 °C for 30 min, with the peptidoglycan-bound protein associated with the pellet, and the unbound protein present in the supernatant. The supernatant was then transferred to a new tube. All reactions were brought to 400 μl using sample loading buffer. Twenty microliters of each fraction were separated using 10% Tricine-PAGE and stained with Coomassie Brilliant Blue. ImageJ (59) was used to quantify band intensity.

[^18O]H2O-based assay to differentiate between hydrolases and lytic transglycosylases—The[^18O]H2O-based assays were conducted as described by Herlihey et al. (42) using M. luteus and S. coelicolor peptidoglycan as substrates. For each, peptidoglycan was resuspended to a final concentration of 1.4 mg/ml in [18O]H2O and briefly sonicated to homogenize the suspension. To start reactions, 1 nmol of purified Rpf protein was mixed with 100 μl of resuspended peptidoglycan in [18O]H2O, and the reaction was brought to 200 μl with Rpf storage buffer. Reactions were incubated at 37 °C for 9.5 h with gentle shaking and then stopped by rapid freezing. Mutanolysin (1.1 nmol) was used as a positive control, whereas reactions without added protein were used as negative controls. Reaction mixtures were thawed and soluble reaction products were separated from insoluble peptidoglycan by centrifugation (15,000 × g, 15 min, 4 °C) prior to analysis by LC-Q-TOF-MS. The insoluble fractions were washed four to five times with 200-μl volumes of water and recovered each time by centrifugation (15,000 × g, 6 min, room temperature). The washed peptidoglycan pellets were resuspended in 0.1 mM potassium phosphate buffer, pH 6.2, and solubilized by mutanolysin (1.1 μm, final concentration) prior to LC-Q-TOF-MS analysis. LC-Q-TOF-MS was performed by injecting samples into an Agilent 1260 Infinity liquid chromatograph interfaced to an Agilent 6540 UHD accurate mass Q-TOF mass spectrometer as described previously (42). MS analyses were conducted at the Mass Spectrometry Facility at the University of Guelph.

**Data availability**

All data are shown, apart from the tandem Q-TOF-MS analysis of each parent ion (Tables 2 and 3), the data for which are
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