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Regulatory RNA in Mycobacterium tuberculosis, back to basics

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Abstract

Since the turn of the millenium, RNA-based control of gene expression has added an extra dimension to the central dogma of molecular biology. Still, the roles of *Mycobacterium tuberculosis* regulatory RNAs, and the proteins that facilitate their functions remain elusive, although there can be no doubt that RNA biology plays a central role in the baterium's adaptation to its many host environments. In this review we have presented examples from model organisms and from *M. tuberculosis* to showcase the abundance and versatility of regulatory RNA, in order to emphasize the importance of these 'fine-tuners' of gene expression.

Abstract

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Graphical abstract. aspects of *M. tuberculosis* regulatory RNA discussed in this review.

Keywords

Mycobacterium tuberculosis; post-transcriptional regulation; regulatory RNA; small RNA; riboswitch; riboregulation

Introduction

Bacterial gene expression consists of two tightly coupled processes, transcription and translation. A detailed, systematic and molecular to global characterisation of how these processes are regulated in pathogens is critical for development and improvement of disease interventions. While the regulation of transcription initiation is chiefly protein-based and orchestrated by sigma- and transcription factors, post-transcriptional regulation is to a large extent influenced by regulatory RNA species or 'riboregulators'.

Until the turn of the millenium, riboregulators were few and far between; their discovery had been serendipituous, and in particular small regulatory RNAs (sRNAs) were mostly associated with plasmid copy number control, phages and transposable elements, reviewed in (Wagner et al., 2002). Two publications from 2001 signalled a change in attitude towards regulatory RNA with systematic searches for intergenic sRNAs based on cDNA cloning, tiling arrays and bioinformatics (Argaman et al., 2001, Wassarman et al., 2001). This was soon followed by the identification and characterisation of a number of highly conserved cis-regulatory elements, e.g. (Grundy et al., 2002, Mironov et al., 2002, Nahvi et al., 2002, Winkler et al., 2002). The trend continued at an accelerated pace with the advent of Deep sequencing methods, including variations such as dRNA-seq, ribo-seq, term-seq and grad-seq providing unbiased and comprehensive mapping of global transcriptomes e.g. (Ingolia et al., 2009, Sharma et al., 2010, Dar et al., 2016, Smirnov et al., 2016). Currently, a plethora of riboregulators with steadily increasing complexity and functionality has been revealed in a substantial number of prokaryotic species, reviewed in (Wagner & Romby, 2015), and by now, the number of riboregulators is likely to equal or exceed the number of transcription factors in many bacteria (Ishihama, 2010, Rau et al., 2015, Holmqvist & Wagner, 2017, Smirnov et al., 2017). The more we uncover, the clearer it becomes that just like the processes of transcription and translation are tightly coupled, the functions of their regulators are interwoven and part of each other's regulons (Arnvig & Young, 2010, Beisel & Storz, 2010, Lee & Gottesman, 2016).

RNA-based control of gene expression

The question 'why regulatory RNA?' keeps emerging in scientific discussions. There are several answers to this question that to some extent dependent on the type of element. Judging by their evolutionary conservation, some regulatory RNAs, including certain riboswitches are very old and possibly remnants from the RNA world and thus predate proteinaceous regulators (McCown *et al.*, 2017). This is not the case for sRNAs, which have evolved later than other non-coding RNAs (ncRNAs) (Peer & Margalit, 2014). Many sRNAs are induced by stress and their synthesis is faster and less costly than transcription and translation of a proteinaceous transcription factor, two qualities that may be of significance during stress. In addition, one might argue from first principles that RNA is the obvious interaction partner for RNA - because basepairing is less complicated than evolving RNA-protein interactions. New sRNAs are continuously emerging as a result of single nucleotide polymorphisms (SNPs) leading to spurious promoters e.g. (Rose *et al.*, 2013), and probably continuously evolving, as sRNA-mRNA interactions can be easily modulated by a few SNPs adjusting their basepairing (Updegrove *et al.*, 2015). Whether these SNPs become fixed depends on the resulting fitness gain or -loss.

Several reports indicate that sRNA regulators have different kinetic properties than protein regulators and that (sRNAs as well as asRNAs) may play a role in suppressing transcriptional noise (Levine & Hwa, 2008, Lasa *et al.*, 2011, Holmqvist & Wagner, 2017).

Finally, it is important to keep in mind that regulatory RNAs operate not instead of, but in addition to, conventional protein-based regulation thereby significantly expanding the number of available components and the complexity of cellular regulatory network.

M. tuberculosis differs from model organisms

M. tuberculosis is subject to multiple stresses and environments through the course of transmission, infection, immune response attack, dormancy and resuscitation, and surviving these changes requires extensive rewiring of its gene expression programme. Regrettably, protein-centric gene expression analysis still provides the main point of reference for *M. tuberculosis.* Our knowledge about mycobacterial riboregulation to a large extent still remains 'dark matter', although there is currently no doubt that this is an important aspect of *M. tuberculosis*'s intracellular life. Most of our knowledge about bacterial riboregulation originates from studies of the Gram-negative Escherichia coli and Salmonella typhimurium, and the Gram-positive Listeria monocytogenes, Bacillus subtilis and Staphylococcus aureus. These model organisms have provided extensive and novel insights into the structure, function and mechanisms of different types of regulatory RNA. In addition to integrating 'ribo-knowledge' into our understanding of how *M. tuberculosis* gene expression is controlled, we may ask whether it has anything to offer to the already vast knowledgebase of RNA biology or whether it is all about finding a cure? Should we simply rely on extrapolating findings from significantly more tractable model organisms instead of sweating in the Cat 3 lab playing catch up? The answer is a resounding 'No!' and the reasons will hopefully be clearer after reading this review. In here we provide a brief overview of different types of riboregulation, showcasing pertinent examples from model organisms as well as *M. tuberculosis*. Due to the rapidly expanding field of regulatory RNA, we are not able to provide a comprehensive overview, and we apologize in advance for the ommission of our colleagues' work that has not been included due to space constraints. We refer to more extensive reviews on riboswitches e.g. (Serganov & Nudler, 2013, Sherwood & Henkin, 2016, Quereda & Cossart, 2017) and on sRNAs e.g. (Updegrove et al., 2015, Wagner & Romby, 2015, Smirnov et al., 2017). Our key aim is to highlight and further the appreciation of riboregulators as important operators in the regulation of gene expression in M. tuberculosis.

Cis- and trans-, non-coding and regulatory RNAs

In the early days of the era of riboregulation, classifications were relatively simple. However, as we learn more about the origins and the functions of the different types of regulatory RNA, the boundaries between *cis*-regulatory elements, *trans*-acting RNA, and *cis*- and *trans*-encoded regulatory RNAs and have become increasingly blurred. In BOX 1 we offer a short list of definitions, bearing in mind that there may be overlaps between these.

RNA chaperones and ribonucleases

As riboregulators adopt more prominent roles, the proteins that regulate their expression, stability and/or degradation have attracted more attention as well. Two classes of proteins important for riboregulator function are the RNA chaperones and Ribonucleases (RNases).

RNA chaperones can have a profound influence on the effect that riboregulators can exert, both as molecular matchmakers, but also due to their role in modulating the stability of these regulators, reviewed in (Vogel & Luisi, 2011). For several years, the widely conserved RNA chaperone, Hfq was considered *the* facilitator of sRNA-mRNA interactions, but different species rely on Hfq to different extents. For example, several sRNAs in the otherwise

highly Hfq-dependent *S. typhimurium* and *E. coli* do not bind Hfq; there are conflicting reports on the requirement for Hfq in Gram-positive bacteria, and certain species including *Helicobacter pylori* and *M. tuberculosis* do not encode an Hfq homologue at all (Jousselin *et al.*, 2009, Arnvig & Young, 2012, Oliva *et al.*, 2015).

Recently, the new global RNA chaperone ProQ was discovered by a combination of density gradient fractionation with mass spectrometry- (of proteins) and and RNA-seq analysis of each fraction (Grad-seq) (Smirnov *et al.*, 2016). Similar to Hfq, there is no ProQ homologue in *M. tuberculosis*, but its identification does rekindle the question as to whether *M. tuberculosis* sRNAs require a matchmaker and if so, how does it work?

A third contender as a general RNA chaperone with a wide range of targets is Cold Shock Protein A (CspA), which belongs to a large family of Csp's, and this protein does have a highly expressed homologue in *M. tuberculosis* (Arnvig *et al.*, 2011, Caballero *et al.*, 2018). CspA is an RNA binding protein that facilitates the melting of secondary RNA structures, which are otherwised stabilised at low temperatures (Jiang *et al.*, 1997). Although well-expressed at all times, the *E. coli* cspA mRNA is itself regulated by a thermoswitch that leads to increased translation at low temperatures (Giuliodori *et al.*, 2010). Whether *M. tuberculosis* CspA ia regulated in a similar manner remains to be seen, but certainly an RNA chaperone that specialises in melting highly structured RNA would seem appropriate for *M. tuberculosis*.

Degrading RNA

The abundance of a transcript is a carefully controlled balance between synthesis and degadation, and hence, ribonucleases (RNases) are crucial players in riboregulation.

In addition to encoding an unprecedented number of type II toxins (Ramage *et al.*, 2009, Sala *et al.*, 2014), *M. tuberculosis* (and other mycobacteria) encode a curious complement of RNases, with elements from both Gram negative (*E. coli*) and Gram positive (*B. subtilis*) species. A comprehensive review of the similarities and differences can be found in (Durand *et al.*, 2015), and we will only mention a few important corner stones. Mycobacteria contain functional homologues of both RNase E and RNase J, while RNase E is absent from *B. subtilis* and RNase J is absent from *E. coli* (Even *et al.*, 2005, Taverniti *et al.*, 2011, Durand *et al.*, 2015).

The endonucleolytic activity of *E. coli* RNase E as well as the exonucleolytic activity of *B. subtilis* RNase J are both sensitive to the phosphorylation state of the 5' nucleotide of their substrates, i.e. they both have a strong preference for mono-phosphorylated transcripts as substrates, and the same 'rules' are likely to apply to the *M. tuberculosis* enzymes (Mackie, 1998, Koslover *et al.*, 2008, Li de la Sierra-Gallay *et al.*, 2008, Mathy *et al.*, 2010, Taverniti *et al.*, 2011). RNA 5' monophosphates can be generated either by endonucleolytic cleavage of a transcript or by the removal of a pyrophosphate group from the 5' nucleotide by the action of the Nudix hydrolase, RppH (RNA pyrophosphohydrolase) (Deana *et al.*, 2008). So far, there are no reports of RppH homologues in mycobacteria, although there are some indications that such an enzyme exists (Moores *et al.*, 2017). Regardless of the enzymes involved and the pathways employed, the degradation of RNA is as important as

RNA 5' leaders

RNA leaders serve as hubs for post-transcriptional regulation, which in many cases involves some means of controlling ribosome entry. In its most basic form, the RNA leader is short and simply provides a Shine-Dalgarno (SD) sequence for binding of the 30S ribosomal subunit (Shine & Dalgarno, 1974). Translation efficiency can be modulated by changing the SD sequence to be more or less complimentary to the 16S sequence or by altering the spacing between the SD and the start codon (Vellanoweth & Rabinowitz, 1992). Certain (longer) RNA leaders have the ability to switch between mutually exclusive conformations that are either permissive or non-permissive for downstream gene expression (transcription or translation), and these are referred to as RNA switches. The switching between alternative RNA structures can be triggered by changes in temperature, pH, metal ions and metabolites, but also by RNA and proteins (Babitzke *et al.*, 2009, Nechooshtan *et al.*, 2009, Ferre-D'Amare & Winkler, 2011, Kortmann & Narberhaus, 2012, Sherwood & Henkin, 2016, McCown *et al.*, 2017). Moreover, the leader is often targeted by sRNAs with a variety of outcomes.

Protein binding 5' leaders

Many 5' leaders regulate downstream gene expression by binding specific proteins reviewed in (Babitzke *et al.*, 2009). This is particularly evident in the case of ribosome biosynthesis, which represents a major drain on cellular resources, and therefore has to be tightly regulated. The expression of many ribosomal protein (r-protein) operons is regulated by direct binding of one or more r-proteins encoded in these operons to their cognate mRNA leaders, usually proximal to the translation initiation region (TIR) typically blocking ribosome entry. This has been extensively characterised in *E. coli* and examples include r-proteins S4, S8, L20, and a complex of L10/L12 (Babitzke *et al.*, 2009). The highly conserved organisation of many of these genes between *E. coli* and other bacteria including *M. tuberculosis* suggests that these feedback mechanisms are also conserved (Arnvig *et al.*, 2011).

In parallel, the transcription of ribosomal RNA (rRNA), is likewise heavily regulated by the 5' leader, which is removed by nucleolytic cleavage from the nascent RNA to generate the mature transcript (Deutscher, 2009). Studies in *E. coli* have elucidated how binding of the antitermination factor NusB and the r-protein S10 to the rRNA leader nucleates a conformational change in the transcription elongation complex that leads to an increase in elongation rate and processivity, i.e. antitermination e.g. (Greive *et al.*, 2005). Similar mechanisms are likely to occur in *M. tuberculosis*, although a specific role for the NusB/E heterodimer has not been demonstrated. However, *M. tuberculosis* has contributed to the antitermination story via its NusA protein and one of the earliest investigations on mycobacterial regulatory RNA. The *M. tuberculosis* NusA lacks the C-terminal domain that masks part of the RNA-binding domain in its *E. coli* counterpart (Gopal *et al.*, 2001). This in turn facilitated the identification of a highly specific interaction between the KH domains of NusA and the antitermination site of the *M. tuberculosis* rRNA leader, an interaction that

was also shown to affect RNAP processivity (Arnvig *et al.*, 2004, Beuth *et al.*, 2005). It remains to be seen how NusA in this context changes from a pausing/termination factor to an antitermination factor.

Riboswitches, leaders sensing metabolites

The term 'riboswitch' refers to a subset of RNA switches that sense changes in the concentration of metabolites or ions without the aid of accessory proteins, and these currently make up the largest class of RNA switches (Serganov & Nudler, 2013, Sherwood & Henkin, 2016, McCown *et al.*, 2017). Riboswitch aptamers sense and interact with a variety of small molecules including nucleotides, amino acids and enzyme co-factors with high affinity and specificity, making them potentially ideal drug targets (Sherwood & Henkin, 2016, Dersch *et al.*, 2017). In many cases these ligands are synthesized by enzymes encoded by the genes that are controlled by the riboswitches itself, thereby implementing feedback regulation (Nudler & Mironov, 2004). The expression platform executes the regulatory output triggered by the presence or absence of ligand binding. If ligand binding leads to reduced expression, the switch is classified as an 'Off' switch (Fig. 1); conversely, if ligand binding leads to increased expression, the switch is classified as an 'On' switch.

Similar to other leader-based regulators, many riboswitches function by blocking/unblocking ribosome entry. Whether the default (i.e. ligand free) conformation is 'On' or 'Off' depends on the individual switch, but in all cases ligand binding to the aptamer domain induces the alternative conformation within the expression platform.

Other riboswitches are based on intrinsic (i.e. factor-independent) termination/ antitermination, which sets them apart from other leader-based modes of regulation as access to the SD is not involved. Moreover, while the two conformers of a translationally regulated riboswitch may exist in a dynamic equilibrium a transcriptionally regulated riboswitch cannot, since both transcription termination and readthrough are irreversible events. This adds additional kinetic requirements to transcriptional riboswitches, as the decision between one or the other conformer has to be made after transcribing the aptamer domain and sensing of a cognate ligand, but before reaching the end of the expression platform. This may require the RNAP to pause at specific and functionally critical positions within the riboswitch to allow for correct co-transcriptional folding of the RNA (Steinert *et al.*, 2017).

A curious characteristic of some riboswitches is that conserved aptamer domains, recognising identical ligands, may be associated with different expression platforms in different species, and in some cases even within the same species. For example, the *B. subtilis* RFN element, which senses Flavin Mononucleotide (FMN) uses SD sequestration within the ypaA riboflavin transporter mRNA, but transcriptional termination in the ribDEAHT mRNA, encoding a series of FMN biosynthetic enzymes (Winkler *et al.*, 2002). It is still unclear exactly where and when a translational expression platform is more or less advantageous than a transcriptional expression platform. While the latter requires ongoing RNA synthesis, the former could in theory regulate the translation of extant transcripts provided that these are relatively stable.

Riboswitches in M. tuberculosis

The vitamin B₁₂**-sensing riboswitch**—Currently only a single metabolite-sensing riboswitch has been experimentally validated in *M. tuberculosis*, although several have been predicted by sequence homology and covariance analyses (Warner *et al.*, 2007, Nawrocki *et al.*, 2015). This is the cobalamine or B₁₂-sensing riboswitch upstream of the *metE* gene, encoding a B₁₂-independent methionine synthase. In the presence of B₁₂, expression of the MetE enzyme, which catalyses the conversion of homocysteine to methionine, is repressed, making this an 'Off' switch. In *M. tuberculosis* H37Rv, this reaction is instead carried out by the B₁₂-dependent isozyme, encoded by *metH*. This means that in the absence of B₁₂, *metE* is required, while in the presence of B₁₂, *metH* is required, due to riboswitch-mediated repression of *metE*. However, this gene has been partially disrupted in *M. tuberculosis* CDC1551 with the result that this strain of *M. tuberculosis* has a severe growth defect in the presence of vitamin B₁₂ (Warner *et al.*, 2007).

A second B_{12} -sensing rioswitch is located in the 5' leader of the *PPE2-cobQ1-cobU* operon. PPE2 (Rv0256c) belongs to the family of proteins sharing proline-proline-glutamate (PPE) N-terminal motifs that were identified in the *M. tuberculosis* genome sequence, many of which are found on the cell surface of *M. tuberculosis*. PPE2 was originally predicted to be a vitamin B_{12} transporter (Rodionov *et al.*, 2003, Vitreschak *et al.*, 2003). However, a more recent study demonstrated that Rv1819c, an ABC transporter is the 'sole corrinoid transporter' responsible for vitamin B_{12} uptake in *M. tuberculosis* under standard *in vitro* growth conditions (Gopinath *et al.*, 2013). Expression of *Rv1819c* is not controlled by a B_{12} riboswitch, and the exact function of PPE2 remains obscure, although the presence of the riboswitch and the *cobQ1-cobU* genes does suggest a role in B_{12} uptake/metabolism.

A Cyclic-di-AMP sensing riboswitch regulates rpfA expression—M. tuberculosis encodes five so-called resuscitation promoting factors (RpfA-E). These are cell wall remodelling enzymes critical for the transition between dormancy and resuscitation (Chao & Rubin, 2010, Kana & Mizrahi, 2010, Mukamolova et al., 2010, Turapov et al., 2014). Precise control of Rpf expression is vital as these enzymes are potentially lethal for M. tuberculosis itself, and multiple, at times shared signals converge in the control of rpf transcription (Fig. 2). The rpfA 5' leader is 272 nucleotides in length and harbours a homologue of the ydaO aptamer domain (Block et al., 2010, Arnvig & Young, 2012). Identified almost a decade before its cognate ligand, cyclic di-AMP (c-di-AMP), the ydaO riboswitch regulates genes associated with cell wall metabolism and osmotic stress in a wide range of bacteria (Barrick et al., 2004, Nelson et al., 2013). The prolonged pursuit for the correct ligand illustrates the difficulty of identifying some riboswitch ligands, even after an element has been characterised. A similar element has been identified in the 5' leader of Streptomyces coelicolor rpfA mRNA, where it has been shown to control expression of RpfA in a c-di-AMP-dependent manner (St-Onge et al., 2015, St-Onge & Elliot, 2017). Due to the close relationship between S. coelicolor and M. tuberculosis, we expect the M. tuberculosis ydaO homologue may also respond to c-di-AMP. Curiously, unlike the B. subtilis element, there are no apparent intrinsic terminators, i.e. a stable stem-loop followed by a poly-U tail, associated with neither the Streptomyces nor the M. tuberculosis

riboswitch, suggesting a different expression platform (Nelson *et al.*, 2013, St-Onge & Elliot, 2017) (J. Green and G. Mukamolova personal communication).

It remains to be seen how this element affects *M. tuberculosis* pathogenesis, but adaptation to changing osmolarity does play an important role in *M. tuberculosis*'s lifestyle as well as in phenotypic drug tolerance (Larrouy-Maumus *et al.*, 2016). Moreover, while the rpfA CDS is highly polymorphic in *M. bovis* (Amadio *et al.*, 2005), the ydao element is 100% conserved between *M. tuberculosis* and *Mycobacterium bovis*, suggesting an important role for this riboswitch.

A novel riboswitch candidate regulating expression of rpfB—Remarkably, three of the five rpf mRNAs (encoding RpfA, B and E), have extensive 5' leaders of more than 100 nucleotides in length (Arnvig *et al.*, 2011, Cortes *et al.*, 2013), suggestive of post-transcriptional regulation; the Rpfs encoded by the same three genes are critical players for Rpf-mediated phenomena such as resuscitation of dormant mycobacteria, growth on solid medium and resistance to detergents (Kana & Mizrahi, 2010).

An RNA switch without a known ligand may be considered a riboswitch candidate (Meyer *et al.*, 2011). Similar to rpfA, the 176-nucleotide 5' leader of the rpfB mRNA harbours an RNA switch (or riboswitch candidate), and like rpfA, identification of the rpfB element precedes identification of its ligand. Unlike ydaO however, the rpfB switch has a recognisable intrinsic terminator structure, and also unlike ydaO, the rpfB switch appears to be restricted to a small subset of pathogenic mycobacteria (Schwenk *et al.*, 2018). By extensive genetic and biochemical analysis, this switch has been shown to control *rpfB* transcription via an intrinsic terminator located immediately upstream of the TTG start codon, which was experimentally re-annotated in the same study. The rpfB switch regulates a *tri-cistronic* operon, which also encodes the methyltransferase KsgA, crucial for ribosome biogenesis and IspE, essential for early steps in *M. tuberculosis* cell wall synthesis (Connolly *et al.*, 2008, Schwenk *et al.*, 2018).

This arrangement provides an intriguing, regulatory link between riboswitch co-ordinated resuscitation from dormancy, ribosome maturation and cell wall synthesis. Moreover, as the operon represents two classical drug targets, i.e. cell wall synthesis and ribosome function under one regulatory roof, it is tempting to speculate that this riboswitch candidate may represent a new target for anti-tuberculosis drug development. Identification of the cognate ligand will undoubtedly provide novel insights into coordinated regulation of macromolecular synthesis as well as post-transcriptional regulation of gene expression in *M. tuberculosis*.

A potential RNA switch regulating rpfE—Little is known about the regulation of *rpfE* expression other than it is induced by chloride, and it is not yet clear if this effect is transcriptional or post-transcriptional (Tan *et al.*, 2013). TSS mapping indicates that the rpfE 5' leader is at least 251 nucleotides in length, and overlaps the divergently transcribed Rv2451 (of unknown function). Similar to the rpfB leader, the rpfE leader harbours the potential to form a stem-loop followed by a poly-U tail close to the TIR. However, the rpfE

poly-U tail is short with only three uridine residues, which may be insufficient to confer intrinsic termination without the support of additional factors/ligands.

Across bacterial species it is clear that only a fraction of riboswitches has been identified to date, and rare (i.e. not broadly conserved) riboswitches are unlikely to be identified by genome alignments. Novel, more experimental approaches are required to tackle this conservation bias. One such approach is Term-seq, which provides a genome-wide display of RNA 3' ends facilitating the identification of conditional terminators and potential novel riboswitches (Dar *et al.*, 2016). Finally, it is worth mentioning in this context that transcriptionally terminated riboswitches can act in *trans* as sRNAs, thus blurring the boundaries *cis*-regulatory elements and *trans*-acting RNA (Loh *et al.*, 2009).

Trans-acting RNAs

In contrast to *cis*-regulatory elements, asRNAs and sRNAs are not part of the transcript they regulate and may therefore be considered '*trans*-acting' (Lease & Belfort, 2000, Loh *et al.*, 2009). This class of transcripts include a small number of protein binding RNAs, and a very large number of basepairing RNAs, which again are divided into *cis*- or *trans*-encoded. Due to space constraints we will focus on basepairing RNAs in this review.

Cis-encoded RNAs are transcripts encoded opposite their target mRNAs (i.e.'true' asRNAs). These transcripts have perfect complementarity to their mRNA targets, suggesting that the resulting hybrids are ideal RNase III substrates. asRNAs can be of varying sizes from <100 nucleotides to several kb, and they are likely to have different modes of action depending on their size and location.

Many, smaller asRNAs are encoded opposite the TIR of their mRNA targets, where they function in a manner similar to *trans*-encoded sRNAs, by blocking ribosome entry and translation. An important class of such small asRNAs are those associated with type I toxin-antitoxin (TA) systems (Brantl & Jahn, 2015). Curiously however, while there is an abundance of Type II/protein based TA systems in *M. tuberculosis*, so far no Type I systems have been identified.

Longer asRNAs can be several hundred nucleotides long and in a few cases even several kb (Arnvig *et al.*, 2011, Lasa *et al.*, 2011, Sesto *et al.*, 2013). An example is the asRNA covering *Rv2817-2816c*, encoding Cas1 and Cas2, respectively in the *M. tuberculosis* CRISPR locus, and while this transcript is relatively abundant, there is very little expression of the coding strand under standard *in vitro* growth conditions, suggesting an inverse correlation in abundance between sense and antisense (Arnvig *et al.*, 2011). The function of these asRNAs is still debated, but pervasive antisense transcription may suppress sense transcriptional noise via transcriptional (RNAP) interferences and/or RNase III mediated cleavage of hybridised sense-antisense transcripts (Lasa *et al.*, 2011).

The 5' leaders of divergently transcribed genes or 3' UTRs of convergently transcribed genes, can also act as asRNA on mRNAs transcribed from the opposing strand, once more blurring the boundaries between cis-regulatory elements and *trans*-acting RNA. This phenomenon was first observed in *L. monocytogenes* (Toledo-Arana *et al.*, 2009) where

it has since led to the to the 'Excludon' concept, coined by Pascale Cossart's group. The excludon specifically refers to 'an unusually long asRNA that spans divergent genes or operons with related or opposing functions (Sesto *et al.*, 2013).

In *M. tuberculosis*, converging 3' UTRs make a significant contribution to the overall antisense transcriptome, and these show a striking enrichment of genes associated with cell wall functions (Arnvig *et al.*, 2011, Cortes *et al.*, 2013). Future studies on gene function and expression should reveal if an excludon mechanism is employed in *M. tuberculosis*.

Finally, some *cis*-encoded sRNAs also have the potential to act as *trans*-encoded sRNAs on mRNA targets with similar sequences as the primary targets e.g. (Arnvig & Young, 2009, Jager *et al.*, 2012).

Trans-encoded sRNAs are encoded in different genomic locations to their targets. The majority of these transcripts are induced by stress and therefore often associated with pathogen adaptation to hostile host environments. In the early days of sRNA identification, searches for sRNAs focused on intergenic regions, and hence this class of regulators were perceived to originate primarily from disinct promoters within these regions, e.g. (Argaman *et al.*, 2001, Wassarman *et al.*, 2001, Arnvig & Young, 2009, Dichiara *et al.*, 2010). However, with the accumulation of data from RNA-seq based methods, it has become evident that many sRNAs are in fact derived from mRNAs. As already mentioned, transcriptionally attenuated leaders can act as sRNAs (Loh *et al.*, 2009); and mRNA 3' UTRs are avid sRNA generators either from processing or from internal promoters (Chao *et al.*, 2012, Chao *et al.*, 2017).

Mode of action

Unlike the interaction between *cis*-encoded (as)RNAs and their targets, the interaction between *trans*-encoded sRNAs and their targets proceeds via limited basepairing apart from a short 'seed sequence', which means that in many cases, trans-encoded sRNAs depend on an RNA chaperone to facilitate the interaction with their targets (Vogel & Luisi, 2011). In addtion to the seed sequence, most sRNAs contain another characteristic feature, which is an intrinsic terminator critical for the interaction with the RNA chaperone Hfq (Otaka et al., 2011, Morita et al., 2017). The limited complementarity also means that prediction of targets can be challenging, and several algorithms have been developed to facilitate this, e.g. TargetRNA2, (Kery et al., 2014) and CopraRNA (Wright et al., 2013). Moreover, a number of experimental approaches have been developed, e.g. RIL-seq, which exploits the proximity of sRNAs to mRNA targets on Hfq (Melamed et al., 2016) or MAPS (pull-downs with MS2tagged sRNAs), which does not require a protein (Lalaouna et al., 2017). Both predictive and experimental approaches require further validation, in particular in an organism such as *M. tuberculosis*, where little remains known about sRNA targets. Individual sRNAs can both repress and increase expression of genes in their regulons, depending on the location of the target region.

Repressing interactions—The most commonly known mode of action for *trans*-encoded sRNAs is repression of translation by blocking the TIR, often followed by mRNA degradation (Fig. 3), reviewed in (Wagner & Romby, 2015).

If the TIR is located early within a multi-cistronic operon, this block may also lead to Rho-dependent termination of transcription further downstream (i.e. polarity), (Bossi *et al.*, 2012). The interaction can also take place downstream of the TIR, several codons into the coding region of the mRNA (Pfeiffer *et al.*, 2009). This may be a means of regulating the many leaderless transcripts in *M. tuberculosis* (Cortes *et al.*, 2013).

Activating interactions—sRNA-mRNA interaction can also lead to increased translation either by direct stabilisation of the mRNA, by unmasking of the TIR and/or by interfering with Rho-dependent termination, reviewed in (Papenfort & Vanderpool, 2015).

An example of direct stabilisation has been observed in *Salmonella*, where the RydC sRNA blocks an RNase E cleavage site in the cfa1 mRNA. This interaction leads to stabilisation of the mRNA even in the absence of translation (Frohlich *et al.*, 2013).

A somewhat more sophisticated means of activation involves a so-called 'anti-antisense' mechanism (Majdalani *et al.*, 1998). In this situation, the leader of the target mRNA contains an auto-inhibitory secondary structure that masks the TIR, and which can be unmasked sRNA binding. A well-characterised example is the *E. coli* rpoS mRNA, which encodes the stationary phase sigma factor, Sigma38 (Battesti *et al.*, 2011). The rpoS mRNA harbours a 567-nucleotide 5' leader, which blocks its own TIR (Majdalani *et al.*, 1998, Peng *et al.*, 2014). Upon binding of one of three sRNAs (DsrA, RprA, ArcZ) to the inhibitory region, the SD sequence and start codon are unmasked via the anti-antisense mechanism to permit translation (Battesti *et al.*, 2011).

Recently, it was shown that the same three sRNAs in addition to unmasking the rpoS mRNA TIR, could also inhibit Rho-dependent termination of *rpoS* transcription in *E. coli* by masking one or more Rho binding sites in the rpoS leader, thus making the sRNA activating effect two-pronged. The authors argued that this novel sRNA-regulated antitermination is likely to be widespread in long leaders (Sedlyarova *et al.*, 2016).

To summarise, sRNAs can both repress or promote translation initiation, and repress or promote Rho-dependent termination of transcription. Moreover, the effect of an sRNA can be greatly enhanced if the mRNA target encodes a regulator such as a sigma or a transcription factor. An overview of different regulatory networks, and their evolution can be found in (Beisel & Storz, 2010, Peer & Margalit, 2014).

M. tuberculosis sRNAs

In spite of several *M. tuberculosis* sRNAs being identified and mapped, and their expression patterns investigated e.g. (Arnvig & Young, 2009, Dichiara *et al.*, 2010, Arnvig *et al.*, 2011, Miotto *et al.*, 2012), only few, including MTS2823, ncRv12659, DrrS and Mcr7, have been functionally characterised in any greater detail (Arnvig *et al.*, 2011, Houghton *et al.*, 2013, Solans *et al.*, 2014, Moores *et al.*, 2017). Like their counterparts in model organisms, *M. tuberculosis* sRNAs are often stress induced and some are highly abundant during infection. The evolutionary conservation of *M. tuberculosis* small RNAs is subject to considerable variation. Some sRNAs, such as ncRv12659 are specific for a subset of *M. tuberculosis* strains (Houghton *et al.*, 2013), some are found throughout species of the *M. tuberculosis*

complex, some a little further afield including non-tuberculous, pathogenic mycobacteria and a few M. tuberculosis sRNAs are conserved in Mycobacterium smegmatis and other Actinomycetes e.g. (Arnvig & Young, 2009, Dichiara et al., 2010, Haning et al., 2014). Many M. tuberculosis sRNAs are highly structured, in part due to the high GC content of the bacterium. Furthermore, by comparing results from 5' and 3' RACE, RNA-seq, northern blotting and RNA structure prediction, it is evident that many *M. tuberculosis* sRNAs do not contain conventional intrinsic terminator structures. For some time this lack of conventional terminators was attributed to the presence of so-called I-shaped terminators, i.e. stem-loop structures without a poly-U tail (Mitra et al., 2008). However, more recently, RNA-seq and in vitro transcription experiments using M. bovis RNA polymerase, have demonstrated that in most cases this type of structure is not sufficient for termination of transcription in vivo or in vitro (Arnvig et al., 2011, Czyz et al., 2014). This in turn suggests that many sRNA 3' termini may be generated by processing in *M. tuberculosis*, setting them apart from the well-known Hfq-dependent sRNAs that require a poly-U tail to function (Otaka et al., 2011). The predicted processing also suggests that some sRNAs may exist as different isoforms, as is the case for the DosR regulated sRNA, DrrS (Moores et al., 2017).

The 108-nucleotide DrrS was first identified by RNA-seq and shown to accumulate to high levels during chronic mouse infection (Arnvig *et al.*, 2011). Recently it was shown that DrrS expression is induced by DosR, but it is a combination of DosR-dependent induction and the unrivalled stability of DrrS that determines the overall levels (Moores *et al.*, 2017).

DrrS has a half-life in the order of several hours due to a stable stem-loop structure at its 5' end. The addition of two or more unpaired nucleotides 5' of this stem-loop, reduces stability significantly, suggesting the involvement of a mycobacterial RppH homologue (Fig. 4). Moreover, this structure increases expression of a *lacZ* reporter when added to the 5' end of its mRNA, suggesting that it represents a general stabilising feature (Moores *et al.*, 2017). In addition to elucidating how RNA stability may be modulated in *M. tuberculosis*, DrrS provides insights into sRNA processing. DrrS is transcribed as a longer (>300 nucleotide) precursor, (DrrS⁺) that is rapidly (in *M. tuberculosis* terms) processed to the shorter, stable 108-nucleotide sRNA (DrrS₁₀₈).

While DrrS⁺ levels peak in early stationary phase, DrrS₁₀₈ accumulates continuously for at least three weeks into stationary phase (Moores *et al.*, 2017). The substantial difference in size and maximum expression between DrrS and DrrS⁺ implies that the longer isoform may play a different role than the the shorter isoform. Apart from shedding light on RNA processing and stability, the DrrS example also highlights the importance of thoroughly characterising multiple aspects of an sRNA before defining its regulon. The application of Term-seq to define *M. tuberculosis* 3' ends on a global scale (Dar *et al.*, 2016), is likely to be hugely informative at this stage.

The best characterised *M. tuberculosis* sRNA in terms of biological role is Mcr7. This sRNA was first identified as a 350-400 nucleotide transcript by cloning and sequencing of *M. bovis* BCG cDNA, and in the same study predicted by sequence homology to be conserved throughout the *M. tuberculosis* complex (Dichiara *et al.*, 2010). RNA-seq later confirmed high expression in *M. tuberculosis* H37Rv (Arnvig *et al.*, 2011). Mcr7 is

encoded downstream of Rv2395 and according to TSS mapping, a single promoter drives transcription in the region downstream of Rv2395 and into PE_PGRS41 (Cortes *et al.*, 2013), suggesting that Mcr7 is (part of) the 5' leader of the latter. However, there is more to this locus than a PE_PGRS protein with a long 5' leader.

In 2011 David Russell's group reported the characterisation of the PhoPR-dependent *aprABC* (Acid and Phagosome Regulated) locus encoding the conserved hypothetical proteins, AprA and AprB, as well as PE_PGRS41 (AprC) (Abramovitch *et al.*, 2011). The aprA coding region lies entirely within the boundaries of Mcr7 (62 basepairs downstream of the annotated TSS), with AprB and AprC encoded downstream of Mcr7 (Fig.5).

The proteins have not yet been experimentally validated in *M. tuberculosis*, but *aprA* does have a likely ribosome binding site upstream of its start codon. Moreover, the recombinant protein has been expressed and purified in *E. coli*, suggesting this is a *bona fide*, stable protein (Abramovitch *et al.*, 2011). Was this then an indication that Mcr7 had been wrongly annotated as an sRNA?

In 2014 the *mcr7/aprABC* promoter was identified as one of the major targets of PhoR (Solans *et al.*, 2014). Based on this finding and the assumption that Mcr7 was a post-transcriptional regulator of gene expression, the authors used *in silico* prediction to identify putative targets of Mcr7, one of which was the tatC mRNA. More specifically positions -16 to +19 relative to the annotated GTG start of the tatC mRNA are targeted by the central portion (nucleotides 119 to 151) of Mcr7, i.e. well within the coding region of *aprA*. The prediction suggests that PhoP/R dependent expression of Mcr7 represses the translation of TatC resulting in reduced secretion of TAT-dependent proteins, which was supported by proteomics on culture supernatants on *M. tuberculosis* wildtype and *phoP* mutant. This study therefore strongly supports the notion of Mcr7 being an sRNA that represses translation of TatC, thereby changing the secretome and modifying the host-pathogen interface (Solans *et al.*, 2014).

So, although AprA has not yet been identified in *M. tuberculosis* and a direct interaction between Mcr7 and tatC mRNA has not been experimentally validated, it appears that this sRNA is a prime candidate for a dual function sRNA in *M. tuberculosis*. As there are no additional TSS in this operon, it also suggests that the 5' end of Mcr7 may regulate *aprA* expression via an as yet uncharacterised post-transcriptional mechanism. If all these elements really represents their annotated functions, this operon represents a complex arrangement of a 5' leader that acts as a *trans*-encoded dual function sRNA.

Concluding remarks

Pathogen survival depends on constant monitoring of, and adaptation to, a range of host environments, an adaptation that sometimes requires rapid and drastic changes in gene expression. This is most efficiently achieved by multi-pronged approaches combining several layers of control, such as transcriptional, post-transcriptional and post-translational regulation. A comprehensive insight into all of these mechanisms is necessary to fully understand how a pathogen interacts with its host, and more importantly, how we

might exploit this to our own advantage. Whether the aim is drug discovery or vaccine development, a thorough understanding of the basic molecular mechanisms of the pathogen in question is fundamental.

In this review we have illustrated (i) how riboregulators work, (ii) argued why riboregulation should be considered by the *M. tuberculosis* community, and (iii) why *M. tuberculosis* should be considered by the RNA community. Although some general rules may apply, riboregulation is still full of surprises, and *M. tuberculosis* is different; with its high GC content (>65%), abundance of leader-less mRNA, distinct complement of RNases and lack of Hfq and ProQ chaperones. In summary, *M. tuberculosis* has the potential to greatly advance our knowledge of RNA based control of gene expression.

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Box1

Non-coding RNA (ncRNA) is a transcript or part of a transcript that does not encode protein or peptide; it may or may not be regulatory. Conversely, there are several examples of regulatory RNAs that are coding.

Cis-regulatory elements are part of the RNAs they regulate, and they include RNA leaders (5' of both coding and ncRNA) and 3' untranslated regions (UTRs), both of which may be a source of sRNAs.

Trans-acting RNAs, as opposed to *cis*-regulatory elements, refer to RNAs that are not part of the transcripts they regulate; they include antisense RNA (asRNA) and small regulatory RNA (\$RNA).

sRNAs comprise a small number of protein binding RNAs, and a large number of basepairing RNAs, which again are divided into *cis*- or *trans*-encoded.

asRNAs are encoded opposite their targets and sizes vary significantly.

Dual function sRNAs are transcripts that act both as riboregulators and as templates for protein synthesis.



Fig. 1.

Riboswitch architecture. Top panel illustrates how a ligand induces transcriptional termination in a transcriptionally controlled 'Off' switch; panel below illustrates a translational 'Off' switch.



Fig. 2.

Control of *rpf* expression in *M. tuberculosis*. The figure illustrates how different, sometimes shared, transcriptional regulators contribute to *rpf* regulation in addition to long 5' leaders, which in the case of *rpfA* harbours a riboswitch with a known ligand (c-d-AMP), in *rpfB* a riboswitch candidate, with unknown ligand and in *rpfE*, a so far entirely uncharacterised element.

Repressing sRNA



Fig. 3.

Basic sRNA modes of action. Top half illustrates how an sRNA (*cis-* or *trans-*encoded) can block ribosome entry and translation. Bottom panel illustrates how an sRNA can activate translation by an anti-antisense mechanism; in this situation the mRNA leader itself blocks translation, by masking the TIR, but an sRNA can interact with the leader to unmask the TIR.



Fig. 4.

Stability of DrrS. Large image shows the predicted structure of $DrrS_{108}$, while the schematic representation illustrates how the number of unpaired nucleotides 5' are inversely correlated to transcript stability.



aprABC operon

Fig. 5.

The *mcr7/aprABC* locus in *M. tuberculosis.* The figure illustrates the elements associated with the PhoP/R regulated operon with the ncRNA Mcr7, which contains an open reading frame encoding the acid inducible AprA, and the proposed interaction between Mcr7 and the tatC mRNA.