Small RNAs of *Borrelia burgdorferi*: Characterizing Functional Regulators in a Sea of sRNAs

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**Review**

*Borrelia (Borreliella) burgdorferi* and closely related genospecies are the causative agents of Lyme disease, the most common tick-borne disease north of the equator. The bacterium, a member of the spirochete phylum, is acquired by a tick vector that feeds on an infected vertebrate host and is transmitted to another vertebrate during subsequent feeding by the next tick stage. The precise navigation of this enzootic cycle entails the regulation of genes required for these two host-specific phases as well as the transitions between them. Recently, an expansive swath of small RNAs has been identified in *B. burgdorferi* and likely many, if not most, are involved in regulating gene expression. Regardless, with only a few exceptions, the functions of these RNAs are completely unknown. However, several state-of-the-art approaches are available to identify the targets of these RNAs and provide insight into their role in the enzootic cycle and infection.

**INTRODUCTION**

Lyme disease is an emerging infectious disease that is the most common tick-borne zoonosis in the Northern hemisphere [1,2]. Human infection typically first presents as a skin rash at the site of the tick bite and can disseminate to multiple organ systems resulting in rheumatic, cardiac, and neurologic manifestations. The etiologic agent of Lyme disease is the spirochete *Borrelia (Borreliella) burgdorferi*, and closely related genospecies, which belong to a morphologically serpentine and deeply branching phylum of bacteria [3-5]. *B. burgdorferi* is maintained in nature in an enzootic cycle that requires a hard tick vector and a vertebrate host [1,6,7]. Larval ticks acquire the spirochetes by feeding on infected reservoir hosts; the spirochetes persist in the midgut through the molt into nymphs, emigrate to the salivary glands when the nymph feeds, and transmit to a vertebrate. *B. burgdorferi* adapts to the species-specific environments, including nutrient resources and immune responses, encountered as it traverses its enzootic cycle by sensing its surroundings and substantially shifting its gene expression via several regulatory systems [1,8,9].

The genes of *B. burgdorferi* constitute a Gordian genome that is highly segmented and predominantly linear [10,11]. A sparse set of recognized orthologs of

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†Abbreviations: sRNA, small RNA; RNA-seq, RNA sequencing; UTR, untranslated region; RBP, RNA-binding protein; asRNA, anti-sense RNA; dsRNA, double-stranded RNA; intrRNA, intragenic RNA.

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transcriptional regulators suggests that post-transcriptional mechanisms (including curious post-translational modifications [12]) contribute to controlling gene expression [1,8,10,13]. We identified the first regulatory RNA, DsrABb, and delineated its role in activating the regulon required for transmission and vertebrate infection [14]. Moreover, we demonstrated that this small RNA (sRNA†), like most sRNAs, requires the RNA chaperone Hfq for its presumed base-pairing; the \textit{B. burgdorferi} Hfq is an oddball, but it can complement a heterologous \textit{hfq} mutant, and it is required for vertebrate infection [15]. Two other RNA-binding proteins have been found in the spirochete: CsrA [16-18] and Bpur [19,20], although neither their function nor RNA targets (other than Bpur binding to its own mRNA) have been determined. The other players in regulating the levels of mRNA are the ribonucleases that degrade RNA and, again, \textit{B. burgdorferi} encodes a limited quiver, including RNase III, M5, Y, and Z [21,22]; the kinetics of decay have been assayed for several mRNAs and half-lives vary from a minute to almost an hour [22].

The number of putative regulatory sRNAs in \textit{B. burgdorferi} has exponentially expanded by the publication of several high-throughput transcriptomes [23-25]. The challenge now for molecular borreliologists is to decipher the function of this glut of sRNAs in the physiology of the spirochete and the pathogenesis of Lyme disease. Therefore, we set a course of experimental approaches, including new methodologies to identify potential targets, to reveal the role of sRNAs in gene regulation.

**EXTENSIVE UNCHARACTERIZED sRNAs IN \textit{B. BURGDORFERI}**

Recently, several strand-specific transcriptome-wide RNA-seq studies have reported large numbers of unannotated transcripts in \textit{B. burgdorferi} [23-25]. Specifically, Lybecker and colleagues identified over 1000 sRNAs with sizes ranging from 50 to 450 nucleotides [25]. These sRNAs were classified based on their genomic location and included sRNAs encoded between (intergenic sRNAs), within (intragenic sRNAs), and opposite (antisense sRNAs) annotated open reading frames (Figure 1). In addition, some sRNAs overlapped the 5′ end of an annotated ORF and were classified as 5′ UTR-associated sRNAs [25]. Small regulatory RNAs influence gene regulation via base-pairing with target mRNAs, affecting their stability, translation, transcription, or processing [26-29]. \textit{Trans}-encoded sRNAs imperfectly base-pair with target mRNAs, encoded in a different genomic location, while antisense RNAs base-pair with complete complementarity to their cognate mRNA. In addition, some sRNAs regulate cellular processes via binding and sequestering target proteins. While high-throughput RNA-sequencing techniques have identified sRNAs in many different bacteria, determining sRNA function has proven far more challenging. Global transcriptional profiling of sRNA mutant strains has been used to map sRNA regulons, but due to functional redundancy, deleting a single sRNA often does not disturb the transcriptome; furthermore, transcriptional changes may not be a direct target of the sRNA. Therefore, many researchers have focused on identifying the direct protein and RNA targets of sRNAs (the sRNA-targetome). Ascertaining the regulons and functions of the newly identified sRNAs in \textit{B. burgdorferi} will reveal the mechanisms of post-transcriptional gene regulation as well as the role of sRNAs in bacterial physiology and virulence.

**IDENTIFYING sRNA-TARGETOMES AND -INTERACTOMES**

Several algorithms predict potential RNA targets of \textit{trans}-encoded sRNA, although the output can include a plethora of false positives [30]. Therefore, high-throughput techniques are necessary to identify direct sRNA-targets (sRNA-targetome). An \textit{in vivo}
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MS2 affinity purification coupled with RNA sequencing (MAPS) technology was developed to globally identify RNA-binding partners of sRNAs [31-33]. The technique is based on the strong interaction between the coat protein of bacteriophage MS2 and the MS2 operator RNA hairpin, an aptamer that can be fused to any sRNA of interest. The sRNA and its binding partners can be co-purified via the MS2 protein and identified by RNA-seq (Figure 2). MAPS has been applied to various classes of RNAs to successfully identify the targetomes of specific RNAs. Recently, two novel RNA-seq-based methods, RIL-seq (RNA interaction by ligation and sequencing) and CLASH (crosslinking, ligation, and sequencing of hybrids), were developed that globally map sRNA-target interactions (sRNA-interactome) [34,35]. However, both methods rely on the interaction of sRNAs with a known RNA-binding protein (RBP). The RNA-binding proteins (Hfq and RNase E) were epitope-tagged and bound RNA was co-immunoprecipitated (co-IP) after in vivo crosslinking. The co-IP RNAs were digested and ligated together to fuse the sRNA and target RNA, generating chimeras, which were subjected to RNA-seq (Figure 3).

IDENTIFYING FUNCTIONAL ANTISENSE sRNAs

Antisense sRNAs (asRNAs) are transcribed opposite to protein-coding strands and have complete complementarity to the corresponding sense RNA. Although asRNAs are ubiquitously found in all organisms, their functions remain mostly elusive [40-42]. The majority of functionally characterized asRNAs are encoded on plasmids and opposite phage, transposon and toxin genes. Antisense transcripts regulate gene expression via base-pairing with their cognate sense RNA and can affect transcription, transcript stability, and translation. asRNAs were initially considered to be transcriptional...
ELUCIDATING THE FUNCTION OF ANTI-SENSE sRNAs AND INTRAGENIC sRNAs

The reverse genetic approach is commonly wielded to dissect gene function in *B. burgdorferi* [11,46]; however, mutating asRNAs or intragenic sRNAs (intraRNAs), which are encoded opposite to or within an annotated ORF, respectively, is technically challenging: deleting the sRNA gene will have a severe polar effect on the overlapping ORF. Several groups have overexpressed asRNAs and intraRNAs to study their function in gene regulation. However, there are inauspicious caveats to this approach, particularly related to overproducing the sRNA, which can sequester crucial RNA-binding proteins, such as Hfq, resulting in pleiotropic dysregulation of gene expression. To generate asRNA-deficient strains, we are currently characterizing the asRNA promoters and then mutating a few nucleotides in the −35 and −10 regions to inhibit transcription of the asRNA while maintaining the coding region of the complementary sense RNA. IntraRNAs can be generated either by processing the full-length mRNA or from an internal promoter within the ORF. Adams et al. globally identified transcriptional start sites and processed ends of transcripts in *B. burgdorferi* [24]. IntraRNAs with internal promoters can be mutated via the same site-directed mutagenesis as the asRNA promoters, which will deplete the cell of only the intraRNA. However, a different approach is required to study intraRNAs that are generated by RNA processing of the mRNA. Chao et al. successfully uncoupled the function of a 3′ UTR intraRNA from its cognate gene by mutating the start codon, effectively blocking translation of the protein, but allowing for sRNA production [47]. Of crucial importance for employing a reverse genetic approach to elucidate the function of genes in *B. burgdorferi* microbiology and Lyme disease pathogenesis, sRNAs that are encoded within an ORF targeted for disruption must be considered in the mutagenesis strategy, as inadvertently deleting an sRNA may have profound pleiotropic consequences.

ROLE OF sRNAs IN PATHOGENESIS

Non-coding RNAs are key components in intricate networks regulating the development of bacterial disease [29,48-52]. Discovering the sRNAs that are important in
the enzootic life cycle of *B. burgdorferi* and the pathogenesis of Lyme disease may reveal novel virulence factors and, potentially, new therapeutic targets. Several culture conditions have been used to simulate the environments of the tick vector and vertebrate host, as well as transmission and acquisition between the two; however, Iyer et al. clearly demonstrated that these conditions are poor mimics of the physiologically relevant milieu [33]. Therefore, a more direct approach to identifying sRNAs involved in virulence or required for the enzootic cycle would be to apply RNA-sequencing of *B. burgdorferi* in both the tick vector and the vertebrate host. Moreover, recent studies have used dual RNA-sequencing to simultaneously sequence host and pathogen transcriptomes in co-culture and from tissue [54-57]. However, there are technical limitations of applying these RNA-seq methods to the tick-mouse experimental model of *B. burgdorferi* infection, particularly due to the low numbers of spirochetes in the vertebrate.

The function of an sRNA in the enzootic cycle of *B. burgdorferi*, including persistence in the tick vector and transmission to the vertebrate host, can be assayed using sRNA mutant strains in the tick-mouse model. However, the deletion of a single sRNA in other pathogenic microorganisms often has a modest phenotype, likely because most sRNAs serve as fine-tuners of gene regulation [58]. In addition, mRNAs can be redundantly regulated by multiple sRNAs, so the mutation of a single sRNA may not have a drastic effect. Therefore, identifying and characterizing sRNAs and their interactomes prior to employing animal models is paramount toward understanding their roles in infection and disease pathogenesis.

**CONCLUSION**

*B. burgdorferi*, the etiologic agent of Lyme disease, encodes a vast sRNA transcriptome [25] that the spirochete almost certainly avails to regulate gene expression during the course of its life cycle as it encounters disparate environmental conditions and host-specific responses [1,8,9,13]. Adaptation to each phase of the enzootic cycle is associated with a sea change in the expressed regulon and response to environmental and metabolic cues; one signal transduction pathway features DsrABb in response to environmental and metabolic cues; one is associated with a sea change in the expressed regulon [1,8,9,13]. We hypothesize that sRNAs are responsible for fine-tuning gene regulation throughout the enzootic cycle of the spirochete and we hope we have provided a beneficial guide to tease out the functional regulators from the sea of sRNAs.

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**REFERENCES**

1. Radolf JD, Caimano MJ, Stevenson B, Hu LT. Of ticks, mice and men: understanding the dual-host lifestyle of Lyme disease spirochaetes. Nat Rev Microbiol. 2012;10:87-99.
2. Steere AC, Strle F, Wormser GP, Hu LT, Branda JA, Hovius JW, et al. Lyme borreliosis. Nat Rev Dis Primers. 2016;2:16090.
3. Steere AC, Grodzicki RL, Kornblatt AN, Craft JE, Barbour AG, Burgdorfer W, et al. The spirochetal etiology of Lyme disease. N Engl J Med. 1983;308:733-40.
4. Burgdorfer W, Barbour AG, Hayes SF, Benach JL, Gunnarldt E, Davis JP. Lyme disease—a tick-borne spirochetosis? Science. 1982;216:1317-9.
5. Benach JL, Bosler EM, Hanrahan JP, Coleman JL, Habicht GS, Bast TF, et al. Spirochetes isolated from the blood of two patients with Lyme disease. N Engl J Med. 1983;308:740-2.
6. Lane RS, Piesman J, Burgdorfer W. Lyme borreliosis: relation of its causative agent to its vectors and hosts in North America and Europe. Annu Rev Entomol. 1991;36:587-609.
7. Piesman J, Schwan T. Ecology of borreliae and their arthropod vectors. In: Samuels DS, Radolf JD, editors. Borrelia: Molecular Biology, Host Interaction and Pathogenesis. Norfolk UK: Caister Academic Press; 2010.
8. Samuels DS. Gene regulation in *Borrelia burgdorferi*. Annu Rev Microbiol. 2011;65:479-99.
9. Caimano MJ, Drecktrah D, Kung F, Samuels DS. Interaction of the Lyme disease spirochete with its tick vector. Cell Microbiol. 2016;18:919-27.
10. Fraser CM, Casjens S, Huang WM, Sutton GG, Clayton R, Lathigra R, et al. Genomic sequence of a Lyme disease spirochete, *Borrelia burgdorferi*. Nature. 1997;390:580-6.
11. Brisson D, Drecktrah D, Eggers CH, Samuels DS. Genetics of *Borrelia burgdorferi*. Annu Rev Genet. 2012;46:515-36.
12. Miller MR, Miller KA, Bian J, James ME, Zhang S, Lynch MJ, et al. Spirochaete flagella hook proteins self-catalyse a lysinoalanine covalent crosslink for motility. Nat Microbiol. 2016;1:16134.
13. Samuels DS, Radolf JD. Who is the BosR around here anyway? Mol Microbiol. 2009;74:1295-9.
14. Lybecker MC, Samuels DS. Temperature-induced regulation of RpoS by a small RNA in *Borrelia burgdorferi*. Mol Microbiol. 2007;64:1075-89.
15. Lybecker MC, Abel CA, Feig AL, Samuels DS. Identification and function of the RNA chaperone Hfq in the Lyme disease spirochete *Borrelia burgdorferi*. Mol Microbiol. 2010;78:622-35.
16. Sanjuan E, Esteve-Gasset MD, Maruskova M, Seshu J. Overexpression of CsrA (BB0184) alters the morphology
and antigen profiles of Borrelia burgdorferi. Infect Immun. 2009;77:5149-62.

17. Sze CW, Li C. Inactivation of bb0184, which encodes carbon storage regulator A, represses the infectivity of Borrelia burgdorferi. Infect Immun. 2011;79:1270-9.

18. Ouyang Z, Zhou J, Norgard MV. CsaA (BB0184) is not involved in activation of the RpoN-RpoS regulatory pathway in Borrelia burgdorferi. Infect Immun. 2014;82:1511-22.

19. Jutras BL, Chenail AM, Carroll DW, Miller MC, Zhu H, Bowman A, et al. Bpur, the Lyme disease spirochete's PUR domain protein: identification as a transcriptional modulator and characterization of nucleic acid interactions. J Biol Chem. 2013;288:26220-34.

20. Jutras BL, Jones GS, Verma A, Brown NA, Antonicello AD, Chenail AM, et al. Posttranscriptional self-regulation by the Lyme disease bacterium's BpuR DNA/RNA-binding protein. J Bacteriol. 2013;195:4915-23.

21. Hargreaves M. Novel ribosome biogenesis in the Lyme disease spirochete Borrelia burgdorferi [Dissertation]: The University of Montana; 2013.

22. Archambault L, Borchert JS, Bergeron J, Snow S, Schlax PJ. Measurements of mRNA degradation in Borrelia burgdorferi. J Bacteriol. 2013;195:4879-87.

23. Arnold WK, Savage CR, Brissette CA, Seshu J, Livny J, Stevenson B. RNA-seq of Borrelia burgdorferi in multiple phases of growth reveals insights into the dynamics of gene expression, transcriptome architecture, and noncoding RNAs. PLoS One. 2016;11:e0164165.

24. Adams PP, Flores Avile C, Popitsch N, Bilusic I, Schroeder R, Lybeck- er M, et al. In vivo expression technology and 5′ end mapping of the Borrelia burgdorferi transcriptome identify novel RNAs expressed during mammalian infection. Nucleic Acids Res. 2017;45:775-92.

25. Popitsch N, Bilusic I, Rescheneder P, Schroeder R, Lybecker M. Temperature-dependent sRNA transcriptome of the Lyme disease spirochete. BMC Genomics. 2017;18:28.

26. Waters LS, Storz G. Regulatory RNAs in bacteria. Cell. 2009;136:615-28.

27. Beisel CL, Storz G. Base pairing small RNAs and their roles in global regulatory networks. FEMS Microbiol Rev. 2010;34:866-82.

28. Storz G, Vogel J, Wassarman KM. Regulation by small RNAs in bacteria: expanding frontiers. Mol Cell. 2011;43:880-91.

29. Caldelari I, Chao Y, Romby P, Vogel J. RNA-mediated regulation in pathogenic bacteria. Cold Spring Harb Perspect Med. 2013;3:a010298.

30. Li W, Ying X, Lu Q, Chen L. Predicting sRNAs and their targets in bacteria. Genomics Proteomics Bioinformatics. 2012;10:276-84.

31. Carrier MC, Lalouana D, Massé E. A game of tag: MAPS catches up on RNA interactomes. RNA Biol. 2016;13:473-6.

32. Desnoyers G, Massé E. Noncanonical repression of translation initiation through small RNA recruitment of the RNA chaperone Hfq. Genes Dev. 2012;26:726-39.

33. Lalouana D, Prévost K, Eyrud A, Massé E. Identification of unknown RNA partners using MAPS. Methods. 2017;117:28-34.

34. Waters SA, McAteer SP, Kudla G, Pang I, Deshpande NP, Amos TG, et al. Small RNA interactome of pathogenic E. coli revealed through crosslinking of RNase E. EMBO J. 2017;36:374-87.

35. Melamed S, Peer A, Faigenbaum-Romm R, Gatt YE, Reiss N, Bar A, et al. Global mapping of small RNA-target interactions in bacteria. Mol Cell. 2016;63:884-97.

36. Vogel J, Luisi BF. Hfq and its constellation of RNA. Nat Rev Microbiol. 2011;9:578-89.

37. Corcoran CP, Rieder R, Podkaminski D, Hofmann B, Vogel J. Use of aptamer tagging to identify in vivo protein binding partners of small regulatory RNAs. Methods Mol Biol. 2012;905:177-200.

38. Said N, Rieder R, Hurwitz R, Deckert J, Urlaub H, Vogel J. In vivo expression and purification of aptamer-tagged small RNA regulators. Nucleic Acids Res. 2009;37:e133.

39. Smirnov A, Förstner KU, Holmqvist E, Otto A, Günster R, Becher D, et al. Grad-seq guides the discovery of ProQ as a major small RNA-binding protein. Proc Natl Acad Sci U S A. 2016;113:11591-6.

40. Thomason MK, Storz G. Bacterial antisense RNAs: how many are there, and what are they doing? Annu Rev Genet. 2010;44:167-88.

41. Georg J, Hess WR. cis-antisense RNA, another level of gene regulation in bacteria. Microbiol Mol Biol Rev. 2011;75:286-300.

42. Lybecker M, Bilusic I, Raghavan R. Pervasive transcription: detecting functional RNAs in bacteria. Transcription. 2014;5:e944039.

43. Lybecker M, Zimmermann B, Bilusic I, Tukhtubayeva N, Schroeder R. The double-stranded transcriptome of Escherichia coli. Proc Natl Acad Sci U S A. 2014;111:3134-9.

44. Lastra I, Toledo-Arana A, Dobin A, Villanueva M, Ruiz de los Mozos I, Vergara-Irigaray M, et al. Genome-wide antisense transcription drives mRNA processing in bacteria. Proc Natl Acad Sci U S A. 2011;108:20172-7.

45. Lioloiu E, Sharma CM, Caldelari I, Helfer AC, Fechter P, Vandenesch F, et al. Global regulatory functions of the Staphylococcus aureus endoribonuclease III in gene expression. PLoS Genet. 2012;8:e1002782.

46. Groshong AM, Blevins JS. Insights into the biology of Borrelia burgdorferi gained through the application of molecular genetics. Adv Appl Microbiol. 2014;86:41-143.

47. Chao Y, Papenfort K, Reinhardt R, Sharma CM, Vogel J. An atlas of Hfq-bound transcripts reveals 3′ UTRs as a genomic reservoir of regulatory small RNAs. EMBO J. 2012;31:4005-19.

48. Pérez-Reytor D, Plaza N, Espejo RT, Navarrete P, Bastias R, García K. Role of non-coding regulatory RNA in the virulence of human pathogenic vibrios. Front Microbiol. 2016;7:2160.

49. Pitman S, Cho KH. The mechanisms of virulence regulation by small noncoding RNAs in low GC gram-positive pathogens. Int J Mol Sci. 2015;16:29797-814.

50. Oliva G, Sahr T, Buchrieser C. Small RNAs, 5′ UTR elements and RNA-binding proteins in intracellular bacteria: impact on metabolism and virulence. FEMS Microbiol Rev. 2015;39:331-49.

51. Papenfort K, Vogel J. Small RNA functions in carbon metabolism and virulence of enteric pathogens. Front Cell Infect Microbiol. 2014;4:91.

52. Toledo-Arana A, Repoila F, Cossart P. Small noncoding
RNAs controlling pathogenesis. Curr Opin Microbiol. 2007;10:182-8.

53. Iyer R, Caimano MJ, Luthra A, Axline D, Jr., Corona A, Iacobas DA, et al. Stage-specific global alterations in the transcriptomes of Lyme disease spirochetes during tick feeding and following mammalian host adaptation. Mol Microbiol. 2015;95:509-38.

54. Nuss AM, Beckstette M, Pimenova M, Schmuhl C, Opitz W, Pisano F, et al. Tissue dual RNA-seq allows fast discovery of infection-specific functions and riboregulators shaping host-pathogen transcriptomes. Proc Natl Acad Sci U S A. 2017;114:E791-E800.

55. Westermann AJ, Gorski SA, Vogel J. Dual RNA-seq of pathogen and host. Nat Rev Microbiol. 2012;10:618-30.

56. Baddal B, Muzzi A, Censini S, Calogero RA, Torricelli G, Guidotti S, et al. Dual RNA-seq of nontypeable Haemophilus influenzae and host cell transcriptomes reveals novel insights into host-pathogen cross talk. MBio. 2015;6:e01765-15.

57. Rienksma RA, Suarez-Diez M, Mollenkopf HJ, Dolganov GM, Dorhoi A, Schoolnik GK, et al. Comprehensive insights into transcriptional adaptation of intracellular mycobacteria by microbe-enriched dual RNA sequencing. BMC Genomics. 2015;16:34.

58. Barquist L, Vogel J. Accelerating discovery and functional analysis of small RNAs with new technologies. Annu Rev Genet. 2015;49:367-94.