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Lyme disease is caused by transmission of the spirochete *Borrelia burgdorferi* from ticks to humans. Although much is known about *B. burgdorferi* replication, the routes and mechanisms by which it disseminates within the tick remain unclear. To better understand this process, we imaged live, infectious *B. burgdorferi* expressing a stably integrated, constitutively expressed GFP reporter. Using isolated tick midguts and salivary glands, we observed *B. burgdorferi* progress through the feeding tick via what we believe to be a novel, biphasic mode of dissemination. In the first phase, replicating spirochetes, positioned at varying depths throughout the midgut at the onset of feeding, formed networks of nonmotile organisms that advanced toward the basolateral surface of the epithelium while adhering to differentiating, hypertrophying, and detaching epithelial cells. In the second phase of dissemination, the nonmotile spirochetes transitioned into motile organisms that penetrated the basement membrane and entered the hemocoel, then migrated to and entered the salivary glands. We designated the first phase of dissemination “adherence-mediated migration” and provided evidence that it involves the inhibition of spirochete motility by one or more diffusible factors elaborated by the feeding tick midgut. Our studies, which we believe are the first to relate the transmission dynamics of spirochetes to the complex morphological and developmental changes that the midgut and salivary glands undergo during engorgement, challenge the conventional […]

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Star M. Dunham-Ems,1 Melissa J. Caimano,1 Utpal Pal,2,3 Charles W. Wolgemuth,4,5 Christian H. Eggers,6 Anamaria Balic,7 and Justin D. Radolf1,8

1Department of Medicine, University of Connecticut Health Center, Farmington, Connecticut, USA. 2Department of Veterinary Medicine, University of Maryland, College Park, Maryland, USA. 3Virginia-Maryland Regional College of Veterinary Medicine, College Park, Maryland, USA. 4Department of Cell Biology and 5Center for Cell Analysis and Modeling, University of Connecticut Health Center, Farmington, Connecticut, USA. 6Department of Biomedical Sciences, Quinnipiac University, Hamden, Connecticut, USA. 7Department of Craniofacial Sciences and 8Department of Genetics and Developmental Biology, University of Connecticut Health Center, Farmington, Connecticut, USA.

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**Introduction**

Almost half of the world’s population is at risk for acquiring a potentially life-threatening vector-borne disease, such as malaria, dengue, Chagas disease, and plague (1). Conventional approaches for decreasing the prevalence of these infections have focused primarily on reducing or eliminating vector populations (2). A number of factors, most notably the emergence of pesticide-resistant vectors, highlight the need for novel control strategies (3). The rational design of preventive measures requires a detailed understanding of pathogen-vector interactions throughout the transmission process. Microorganisms transmitted by hematophagous arthropods have evolved specialized developmental programs that enable them to capitalize on vulnerabilities created by the remodeling of vector tissues during the blood meal in order to gain access to their vertebrate hosts (4–6). Confocal microscopy of fluorescent microbes has provided investigators with a powerful methodology for visualizing the discrete steps by which live organisms disseminate within the arthropod (7–9).

Lyme disease, caused by the spirochete *Borrelia burgdorferi*, is the most common arthropod-borne infection in the United States (10). If untreated, Lyme disease can lead to a wide array of complications typically involving the heart, joints, or nervous system (11). *B. burgdorferi* cycles between 2 physiologically distinct environments, represented by the arthropod vector (*Ixodes* ticks) and mammalian host (11, 12). During larval acquisition, spirochetes colonize the midgut, which then remains infected through subsequent molts into the nymphal and adult stages. The nymphal stage is the most clinically relevant because it is usually responsible for the transmission of spirochetes to humans, an accidental host outside of the enzootic cycle (11, 12). Nymphal feeding induces the exponential expansion of spirochetes and migration of organisms out of the midgut into the hemocoel (13, 14), a process facilitated by the acquisition of plasminogen (15, 16). Once in the hemocoel, Lyme disease spirochetes must contend with the primary mediators of tick innate immunity, hemocytes and defensins, as they navigate toward the salivary glands (15, 17). Following attachment to the surface of the glands, spirochetes penetrate the surrounding basement membrane and are conveyed by the saliva into the dermis of the mammalian host (18–20). While differential gene expression by *B. burgdorferi* and the kinetics of spirochete replication during the nympha1 blood meal have been investigated extensively (12–14, 21–24), the routes and mechanisms that *Borrelia* utilize to disseminate within the tick have received considerably less attention. Here, we imaged live, infectious *B. burgdorferi* expressing a stably integrated, constitutively expressed GFP reporter in order to elucidate the interactions between spirochetes and *Ixodes scapularis* tissues during the nymphal blood meal. Our investigations revealed that *B. burgdorferi*...
progress through the feeding tick via a novel, biphasic mode of dissemination. In the first phase, spirochetes, positioned at varying depths throughout the midgut epithelium at the onset of feeding, replicate extensively, forming networks of nonmotile organisms that advance toward the basolateral surface of the epithelium while adhering to differentiating, hypertrophying, and detaching epithelial cells. This process, which we have designated “adherence-mediated migration,” involves the inhibition of spirochete motility by one or more diffusible factors elaborated by the feeding midgut. The second phase of dissemination is marked by the transition of nonmotile spirochetes into motile bacteria that then complete the penetration of the midgut, traverse the hemocoel, and migrate to and penetrate the salivary glands en route to the mammal. This motile, invasive phenotype continues throughout mammalian infection, ending when spirochetes colonize the midguts of feeding naive larvae to complete their enzootic cycle. Our studies, the first, to our knowledge, to relate the transmission dynamics of Lyme disease spirochetes to the complex morphological and developmental changes that *Ixodes* tick midguts and salivary glands undergo during engorgement, challenge the conventional viewpoint that *B. burgdorferi* dissemination within ticks is exclusively motility driven.

**Results**

**Bb914 stably maintains a GFP reporter while retaining wild-type infectivity for mice and ticks.** Studies by us (25) and others (26) have demonstrated that borrelial shuttle vectors can be lost over time in the absence of selection during in vivo growth. We reasoned that insertion of a *gfp* expression cassette into cp26, a highly stable, endogenous borrelial plasmid (27, 28), would prevent loss of the fluorescent reporter during the extensive replication of spirochetes that occurs following the nymphal blood meal (14, 29), thereby enhancing our ability to track organisms in locales where bacterial burdens are known to be low (e.g., hemolymph and salivary glands) (15, 18, 30, 31). To this end, Bb914 was generated by inserting *gfp* under the control of the constitutive flaB promoter (*P*flaB) into the bbb20-bbb21 intergenic region of cp26 in CE162, a virulent *B. burgdorferi* 297 clone (32) (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI39401DS1). Bb914 and CE162 displayed essentially identical in vitro growth kinetics (data not shown) and protein expression profiles following temperature shift and cultivation within dialysis membrane chambers (DMCs) (Supplemental Figure 2). Of note, the levels of outer surface protein C (OspC) expressed by temperature-shifted and DMC-cultivated Bb914 and CE162 were highly similar (Supplemental Figure 2), indicating that insertion of the *P*flaB cassette did not alter transcription of *ospC* (bbb19), a gene essential for infectivity (26, 33). The fluorescent properties of Bb914 in vitro were compared with those of CE863, a CE162 transformant harboring *P*flaB::gfp on a cp32-derived shuttle vector (34), using flow cytometry. Virtually all Bb914 retained the cp26-borne *gfp* reporter in the absence of antibiotic, whereas CE863 cultured without selection contained a substantial nonfluorescent population that increased with repeated passaging (Figure 1A). Bb914 also was brighter than CE863, as indicated by a significantly greater mean fluorescence intensity value (324 ± 47 and 228 ± 24, respectively, for passage 1; *P* = 0.0227).

Bb914 and CE162 possessed comparable infectivity (ID₅₀ values of 116 and 224, respectively; *P* = 0.9462) as measured by syringe inoculation of C3H/HeJ mice. Importantly, all Bb914 isolated from nymphs used in these studies were generated by feeding naive larvae on syringe-inoculated C3H/HeJ mice as described in Methods. Data represent the means from 3 independent experiments; error bars indicate SD.
ability of GFP-positive *Borrelia* to be acquired and maintained by ticks, naive *I. scapularis* larvae were allowed to feed on either BALB/c SCID or C3H/HeJ mice 2 weeks following intradermal syringe inoculation with a dose of 1 × 10⁴ *Bb914*. All *Bb914*-infected larvae examined at the time of repletion contained numerous fluorescent spirochetes, while quantitative PCR (qPCR) confirmed that *Bb914* survived the larval molt and expanded during the nymphal blood meal as efficiently as its parent, CE162 (Figure 1B). Spirochete burdens were essentially identical in flat (i.e., unfed) and fed nymphs regardless of whether the starting larvae had fed on BALB/c SCID or C3H/HeJ mice infected with *Bb914* (Supplemental Figure 3). Indirect immunofluorescence assay (IFA) studies demonstrated that the percentage of *Bb914* expressing OspC at 72 hours after placement (44.7% ± 6.79%) was highly similar (P > 0.05) to that previously observed for CE162 (49.43% ± 2.25%) (35) (Supplemental Figure 4), further establishing that the insertion in cp26 had no placement (44.7% ± 6.79%) was highly similar (P > 0.05) to that pre-

*Figure 2*

*B. burgdorferi* are distributed throughout the midguts of flat nymphs. (A) Low-power epifluorescence image of an isolated unfed midgut. (B) Composite confocal image showing the distribution of spirochetes through the full thickness (40 μm) of an unfed midgut. (C) Scheme used to acquire optical sections through an unfed *Bb914*-infected midgut and cartoon depiction of the cellular morphology at varying depths within the epithelium (modeled after ref. 46). (D–F) 3-μm composite images of optical sections acquired (D) at the luminal surface, (E) midway through the epithelium, and (F) at the basement membrane. (G–I) Digital enlargements of single, 1-μm optical sections of the boxed area in B acquired (G) at the luminal surface, (H) midway through the epithelium, and (I) at the most distal point where spirochetes were observed. Arrows, arrowheads, and asterisks indicate spirochete aggregates, intercellular spirochetes, and extracellular blebs, respectively; localization of spirochetes and blebs was determined by optical sectioning. Here and elsewhere, red staining denotes FM4-64; green staining denotes GFP. Scale bars: 50 μm (A); 25 μm (B, D–F); 10 μm (G–I).

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Small undifferentiated cells that repeat the digestive cycle (43, 45, 46) reached near confluence on the luminal surface (Figure 4F), while dense aggregates extended from the lumen to form elaborate networks that essentially encased individual epithelial cells (Figure 4G). Identical networks of spirochetes were observed by confocal as well as epifluorescence microscopy regardless of whether the specimens were mounted in Vectashield, immersion oil, CMRL-1066 (CMRL), or Barbour-Stoenner-Kelley-II (BSK-II) medium (Supplemental Figure 5). In contrast to the 48-hour time point, organisms now were observed at the basolateral pole of the epithelial layer (Figure 4H); analysis of consecutive optical sections and orthogonal views revealed that some spirochetes at or near the basement membrane were wedged between epithelial cells in a “head-on” orientation (Figure 4H). In other regions of the midgut, the epithelial layer was substantially thinner (Figure 4, I–L), presumably because it consists mainly of undifferentiated cells (43, 45, 46). Although spirochetes were numerous in these areas, including at the basement membrane (Figure 4L), they did not reach the same level of confluence observed in the hypertrophied regions (Figure 4, compare E and I with the orthogonal views in H and L). Here, too, spirochetes in a head-on orientation could be identified between epithelial cells.

**Spirochetes advance around and between epithelial cells toward the basement membrane.** To further correlate the distribution of spirochetes with epithelial cell growth and differentiation during feeding, we examined silver-stained paraffin sections and FM4-64–labeled cryosections of midguts isolated at 48 and 72 hours after placement. At the 48-hour time point, hypertrophy of the epithelial cells was prominent, with some larger differentiated cells budding into the lumen (Figure 5, A and D). Spirochetes were distributed as individual organisms and as focal aggregates attached to the luminal surface and also wedged between epithelial cells, without any obvious preference for cell type. Consistent with the long axis confocal images of 48-hour-fed midguts (Figure 4, A–D), cryosectioning clearly demonstrated that organisms deeply penetrated the intercellular spaces but generally stopped short of the basement membrane (Figure 5, A and D). As noted above, by 72 hours after placement, the midgut epithelium either was markedly disrupted and sloughing cells into the lumen (Figure 5B) or consisted primarily of a thin layer of undifferentiated cells (Figure 5, C and E). In thicker areas of the midgut, spirochetes outlined many of the hypertrophied and detached epithelial cells in a manner reminiscent of the networks depicted by the long axis optical sections (compare Figure 4G and Figure 5B). In regions consisting predominantly of undifferentiated cells, the epithelial layer was markedly thinner and the spirochete densities were substantially lower (Figure 5, C and E). Cryosections of these areas highlighted the perpendicular orientation of spirochetes directly adjacent to the basement membrane (Figure 5F).

**Spirochetes within intact flat and fed midguts are nonmotile but respond differently when the midgut environment is breached.** *B. burgdorferi* is known for its vigorous motility in vitro (47, 48). During routine monitoring of intact 48-hour- and 72-hour-fed midguts by epifluorescence microscopy, as well as during the confocal analyses described above, we were surprised that the spirochetes did not appear to be motile. The lack of motility within the lumen and at varying depths...
throughout the epithelial layer was confirmed by time-lapse confocal imaging of numerous intact (i.e., excluding FM4-64) midguts isolated at 48 and 72 hours after attachment (data not shown and Supplemental Video 1, respectively) and was observed regardless of the type of mounting medium used (data not shown). After the time-lapse imaging was completed, we assessed the viability of spirochetes within the imaged specimens by crushing the midguts in CMRL or BSK-II medium; the resulting suspensions contained numerous, highly motile organisms that were cultivatable in BSK-II and formed colonies on solid-phase pBSK medium (data not shown).

It is generally presumed that spirochetes within unfed midguts are nonmotile (49); we confirmed this by time-lapse confocal imaging at various depths through multiple Bb914-infected, flat nympha midguts (Supplemental Video 2). Given the marked differences in both the midgut environment (43, 46) and the metabolic states of spirochetes before and during feeding (50–54), we conjectured that organisms in unfed and fed midguts would differ in their capacities to become motile following exposure to culture medium. To test this idea, unfed midguts immersed in CMRL or BSK-II were nicked and examined by time-lapse epifluorescence microscopy. Nearly all spirochetes attached to the midgut epithelium near the nick site remained nonmotile throughout a 30-minute viewing period (Supplemental Video 3). The few organisms released into the surrounding medium were nonmotile when first observed and exhibited only sluggish motility, at best, after 30 minutes (Supplemental Video 3). Seventy-two-hour-fed mid-

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**Figure 4**

*B. burgdorferi* progress through the nympha midgut during feeding in the form of epithelial cell–associated networks. Representative confocal micrographs of midguts isolated from Bb914-infected nympha at (A–D) 48 and (E–L) 72 hours after placement; E–H and I–L, respectively, show hypertrophied and undifferentiated regions of 72-hour-fed midguts. Composite images in A, E, and I depict the full thickness of the respective epithelial layers. 3-μm composite images show spirochetes (B, F, and J) at the luminal surface (C, G, and K) midway through the epithelium and (D, H, and L) at the basement membrane. Orthogonal views (y–z and x–z axes) for the 72-hour-fed midguts in H and L show that some spirochetes near the basement membrane (arrows) are in a head-on orientation. The positions of the lumen (Lu) and basement membrane (BM) in each orthogonal view are indicated. The asterisk in the y–z axis view of H denotes a spirochetal network. Scale bars: 25 μm.
guts were similarly examined. Unlike what was observed with the nicked, unfed midguts, many of the organisms attached to epithelial cells near the nick site (≤ 1.5 mm) were flexing at their unattached ends as soon as they were imaged (Supplemental Video 4) and became increasingly vigorous over a 30-minute viewing period (data not shown). In contrast to the motile organisms within the lumens of intact, fed midguts (Supplemental Video 1), spirochetes released from the luminal contents (indicated by the presence of blood remnants) into CMRL or BSK-II displayed vigorous motility (Supplemental Video 4). The majority of spirochetes more distal to the nick site (i.e., within the stomach, ~5 mm away), however, remained nonmotile (Supplemental Video 4).

**Fed midguts inhibit spirochete motility in a gelatin matrix-based assay.** To clarify the above findings, we devised an experimental system to examine the effect of the midgut environment on spirochete motility. Consistent with the well-documented propensity of spirochetes to localize in collagen-rich tissues (55, 56), we found that a substantial percentage (56% ± 12%) of input Bb914 readily penetrated a 5% gelatin matrix and were retained following extensive washing. Organisms within the matrix remained actively motile for at least 6 hours at room temperature. In contrast to BSK-II, in which nearly all organisms displayed translational motility characteristic of *B. burgdorferi* in a low-viscosity milieu (57, 58), spirochetes within the gelatin matrix exhibited 3 distinct motility patterns: motile/translational, motile/nontranslational, and nonmotile (Table 1; see Supplemental Video 5 for representative examples). Installation of a small aliquot (1–2 μl) of mouse blood into a sample well in a corner of a matrix resulted in a marked and progressive increase in both the overall percentage and degree of motility of the organisms over a 15-minute viewing period; this effect was noticeably less pronounced with increasing distance from the well (Table 1). In stark contrast, the vast majority of spirochetes (~92%) in proximity to the sample well were nonmotile soon after the addition of a naive, 72-hour-fed midgut to the sample well and remained so throughout the entire 15-minute viewing period (Table 1). Contrary to what was observed with blood, the inhibitory effect of the fed midgut diminished as the distance from the well increased (Table 1). Notably, neither unfed midguts nor salivary glands isolated at 72 hours after placement had any discernible effect on spirochete motility. Trajectory plots depicting the motility patterns of spirochete populations exposed to the various samples in this model system illustrate the contrast between the stimulatory effect of fresh mouse blood and the potent inhibitory effect of 72-hour-fed midguts (Figure 6).

**Penetration of *Borrelia* into the hemocoel is a rare event.** Spirochetes were not observed in hemolymph until 72 hours after placement (data not shown), suggesting that this time point marks a critical period during which organisms complete the traversal of the midgut. Despite careful examination of the basolateral surfaces of fifty 72-hour-fed midguts, we found only a single spirochete protruding through the basement membrane into the hemocoel (Figure 7A). The ostensible rarity of this event was in accord with the small number of organisms (~3–4 per tick) observed in isolated hemolymph (Figure 7B) by epifluorescence microscopy and the minuscule number detected by qPCR of hemolymph as compared with fed midguts (477 ± 322 versus 5.88 × 10^6 copies per tick, respectively) at 72 hours after placement. Interestingly, the robust motility of organisms in hemolymph (Supplemental Video 6) was in marked contrast to the lack of motility of spirochetes in intact unfed and fed midguts. Hemocytes containing GFP-positive material, presumably phagocytosed spirochetes, also were visualized in hemolymph (Figure 7C).

**Borrelia attach to all acini types but invade only types II and III.** Staining with FM4-64 allowed unambiguous identification, using established morphologic criteria (43, 59, 60), of all 3 acini types in salivary glands isolated from flat (Figure 8A) and (B, C, E, and F) 72 hours after placement on naive C3H/HeJ mice. (D) Digital enlargement of the boxed area in E. Representative examples of differentiated (dc) and undifferentiated cells (uc) are indicated. Arrows indicate spirochete aggregates on the epithelial surface (A–C) or aggregates within the intercellular spaces (D). In stark contrast, the vast majority of spirochetes within the gelatin matrix exhibited 3 distinct motility patterns: motile/translational, motile/nontranslational, and nonmotile (Table 1). Installation of a small aliquot (1–2 μl) of mouse blood into a sample well in a corner of a matrix resulted in a marked and progressive increase in both the overall percentage and degree of motility of the organisms over a 15-minute viewing period; this effect was noticeably less pronounced with increasing distance from the well (Table 1). In stark contrast, the vast majority of spirochetes (~92%) in proximity to the sample well were nonmotile soon after the addition of a naive, 72-hour-fed midgut to the sample well and remained so throughout the entire 15-minute viewing period (Table 1). Contrary to what was observed with blood, the inhibitory effect of the fed midgut diminished as the distance from the well increased (Table 1). Notably, neither unfed midguts nor salivary glands isolated at 72 hours after placement had any discernible effect on spirochete motility. Trajectory plots depicting the motility patterns of spirochete populations exposed to the various samples in this model system illustrate the contrast between the stimulatory effect of fresh mouse blood and the potent inhibitory effect of 72-hour-fed midguts (Figure 6).

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The nymphal midgut undergoes complex physiological and morphological changes during feeding (43, 45, 46). The influx of interstitial fluid during the preparatory phase primes the epithelial cells for nutrient uptake and stimulates activation of the digestive machinery (43, 45, 46). Toward the end of the preparatory phase, midgut epithelial cells begin to secrete components of the peritrophic membrane, a semipermeable layer composed of chitin, proteoglycans, and proteins that protects the epithelium from ingested pathogens and abrasive food particles (65, 66). As feeding progresses, salivary enzymes degrade formed elements in the blood, the remnants of which are avidly phagocytosed and digested to completion within lysosomal food vacuoles (43, 45, 46). The uptake of nutrients promotes the growth and differentiation of epithelial cells, some of which eventually slough from the basement membrane and disintegrate, releasing their contents into the lumen; replacement by adjacent undifferentiated cells maintains the integrity of the epithelial layer (43, 45, 46). Our results highlight the remarkable degree to which spirochetes accommodate to, and even exploit, this rapidly changing and potentially dangerous milieu. For instance, *B. burgdorferi* escape digestion within the lumen by being situated in the ectoperitrophic space, a protected position presumably attained when spirochetes colonize the midgut at the time of larval acquisition (67). Another example was provided by our observation that spirochetes shed copious blebs at the outset of feeding. The contrast between the avid internalization of blebs by epithelial cells and the exclusively extracellular localization of intact organisms was impressive and suggests that *Borrelia* employ an unknown mechanism to evade uptake and certain destruction while maintaining an intimate association with highly active digestive cells. It is widely believed that the logarithmic expansion of the spirochete population in response to the blood meal occurs in parallel with the detachment of individual organisms from the luminal surface and their rapid migration through the midgut epithelium (13, 14, 37, 40, 68). We found instead that replicating spirochetes coalesce from scattered foci distributed on and deeply within the epithelium into networks of nonmotile organisms that advance toward the basement membrane by surrounding individual epithelial cells. According to this alternative model, which we call adherence-mediated migration, the strategic positioning of spirochetes at multiple points around individual cells prior to feeding facilitates midgut traversal by enabling the

### Table 1

<table>
<thead>
<tr>
<th>Experimental sample</th>
<th>Motile/ nontranslationalA</th>
<th>Motile/ translationalA</th>
<th>Nonmotile</th>
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<tr>
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<td>46.78 ± 12.59</td>
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<td>Salivary glands, 0 mmC, D</td>
<td>12.45 ± 9.15</td>
<td>56.44 ± 6.12</td>
<td>31.11 ± 15.07</td>
</tr>
</tbody>
</table>

Values denote mean percentages ± SD calculated using three 5-minute viewing intervals taken over a 15-minute observation period. A minimum of 500 organisms were scored for each sample. Highly similar results were obtained for each sample in 3 independent experiments.

A Organisms that exhibited net displacement in the x and/or y axis. B Organisms that exhibited obvious signs of motility (i.e., flexing) but no net displacement in either axis. C Measurement denotes distance from the well containing the experimental sample. D Similar values were obtained at the 5-mm distance (not shown).

### Discussion

Motility has long been believed to be essential for dissemination of *B. burgdorferi* from the site of inoculation and subsequent invasion of mammalian host tissues (61, 62). The linkage between motility and infectivity for mammals recently was strengthened by intraval imaging of blood-borne spirochetes in mice; following attachment to the vascular endothelium, *B. burgdorferi* move toward and then rapidly penetrate endothelial cell junctions (63, 64). Previous attempts to elucidate the mechanisms of spirochete dissemination within the arthropod have employed fixation and microscopic techniques that preclude analysis of the dynamic interactions between pathogen and vector (19, 20, 37).

In this study, we utilized a *B. burgdorferi* transformant harboring a stable, constitutively expressed GFP reporter to track live spirochetes during transmission by *I. scapularis* nymphs. Our investigations revealed, quite unexpectedly, that *Borrelia* progress toward the basement membrane of the midgut as nonmotile aggregates that are closely associated with the epithelial cells. Only well into the feeding process do spirochetes transition to the motile, invasive phenotype in which they conjoin their journey within the nymphal tick, infect a mouse, and, ultimately, transit back into a naive vector during larval feeding.

The nymphal midgut undergoes complex physiological and morphological changes during feeding (43, 45, 46). The influx of interstitial fluid during the preparatory phase primes the epithelial cells for nutrient uptake and stimulates activation of the digestive machinery (43, 45, 46). Toward the end of the preparatory phase, midgut epithelial cells begin to secrete components of the peritrophic membrane, a semipermeable layer composed of chitin, proteoglycans, and proteins that protects the epithelium from ingested pathogens and abrasive food particles (65, 66). As feeding progresses, salivary enzymes degrade formed elements in the blood, the remnants of which are avidly phagocytosed and digested to completion within lysosomal food vacuoles (43, 45, 46). The uptake of nutrients promotes the growth and differentiation of epithelial cells, some of which eventually slough from the basement membrane and disintegrate, releasing their contents into the lumen; replacement by adjacent undifferentiated cells maintains the integrity of the epithelial layer (43, 45, 46). Our results highlight the remarkable degree to which spirochetes accommodate to, and even exploit, this rapidly changing and potentially dangerous milieu. For instance, *B. burgdorferi* escape digestion within the lumen by being situated in the ectoperitrophic space, a protected position presumably attained when spirochetes colonize the midgut at the time of larval acquisition (67). Another example was provided by our observation that spirochetes shed copious blebs at the outset of feeding. The contrast between the avid internalization of blebs by epithelial cells and the exclusively extracellular localization of intact organisms was impressive and suggests that *Borrelia* employ an unknown mechanism to evade uptake and certain destruction while maintaining an intimate association with highly active digestive cells. It is widely believed that the logarithmic expansion of the spirochete population in response to the blood meal occurs in parallel with the detachment of individual organisms from the luminal surface and their rapid migration through the midgut epithelium (13, 14, 37, 40, 68). We found instead that replicating spirochetes coalesce from scattered foci distributed on and deeply within the epithelium into networks of nonmotile organisms that advance toward the basement membrane by surrounding individual epithelial cells. According to this alternative model, which we call adherence-mediated migration, the strategic positioning of spirochetes at multiple points around individual cells prior to feeding facilitates midgut traversal by enabling the
expanding population of *Borrelia* to reach the base of the epithelium faster than the ballooning epithelial cells can transport them toward the lumen. Residual organisms left behind by detaching digestive cells are likely to be ideally situated for penetrating the thinner regions of the epithelium comprised predominantly of undifferentiated cells. It is interesting to note that spirochetal networks are readily discernible, though were not recognized as such, in previously published confocal immunofluorescence micrographs of fed nymphal midguts (14, 24, 68); the significance of network formation most likely was not appreciated in these studies because the specimens were fixed prior to imaging.

It is a central tenet of Lyme disease pathogenesis that *Borrelia* transition between 2 dramatically different transcriptional programs with the onset of tick feeding, with motility being a salient, distinguishing phenotypic trait. Live imaging has revealed that this conception is an oversimplification. That *Borrelia* within unfed nymphal midguts are nonmotile is consistent with the quiescent metabolic state of organisms in this milieu (50–54) and is in line with observations made with crushed midguts dating back to the discovery of the Lyme spirochete as a tick-borne pathogen (49).

On the other hand, our finding that the fed midgut is populated with replicating, but nonmotile, spirochetes was unexpected and runs counter to the prevailing notion, which equates metabolic activity with motility (58). Evidence presented herein, however, indicates that the ostensibly identical motility phenotypes in flat and fed ticks are mechanistically distinct. First, we found that spirochetes in unfed midguts displayed, at best, sluggish motility even after prolonged exposure to CMRL or BSK-II, while *Borrelia* in fed midguts became highly active within minutes following exposure to the same medium. Second, in a gel matrix model in which the effect of exogenous substances on spirochete motility could be assessed qualitatively and quantitatively, organisms in proximity to fed, but not flat, midguts rapidly lost motility. To explain these observations, we hypothesize that the physiologic adaptation of spirochetes to the nutritionally deficient milieu of the unfed midgut restricts their capacity to import and/or utilize nutrients needed to power the flagellar motor, whereas the flagellar motors of spirochetes actively replicating within the fed midgut are reversibly disengaged in response to one or more diffusible factors elaborated by the midgut epithelium. Indeed, the lack of motility displayed by spirochetes in the lumens of fed midguts was remarkable in light of the strong chemoattractant properties of blood in the gel matrix model and other in vitro systems (54, 69). These results, although counterintuitive at first blush, make sense; if blood in the midgut acted as a chemoattractant, spirochetes would never disseminate during feeding. The rapidity of the transition to motility once the midgut environment was breached argues strongly for a posttranscriptional form of control and is consistent with a growing number of reports that bacteria can quickly and reversibly modulate their state and degree of motility via protein–protein interactions involving the flagellar motor and chemosensory apparatus (70, 71).

The midgut epithelium is well recognized as an anatomical barrier to the migration of vector-borne pathogens following ingestion by hematophagous arthropods (4–6). Three complementary lines of evidence suggest that the midgut of *I. scapularis* functions in a similar capacity to impede the dissemination of *B. burgdorferi*. One is the marked reduction in spirochete numbers in the hemolymph relative to the fed midgut; it is noteworthy that the spirochete burdens in hemolymph determined here by qPCR are highly similar to those reported by Coleman et al. (15), who enumerated spirochetes by IFA. A second comes from our micrographs, which revealed a tapering in the density of networks toward the basolateral pole of the epithelium, where some spirochetes were wedged between cells, assuming a head-on orientation. The last was the rarity of actually observing spirochetes “caught in the act” of penetrating through the midgut into the hemocoel. Contrary to an earlier report (40), we did not find any evidence for the intracellularility of intact spirochetes as they traverse the midgut. Our results imply that the intercellular junctions and basement membrane form a series of anatomical barriers that *Borrelia* must cross in order to gain access to the hemocoel. A key prediction of our model is that adherence-mediated migration positions spirochetes in proximity to the intercellular junctions as they become weakened by remodeling and the basement membrane is thinned by luminal disintegration (43, 45, 46). Negotiation of these structural impediments presumably requires the resumption of motility and is likely facilitated by the binding of host-derived molecules, such as plasminogen, to the bacterial surface (15, 16). Conceivably, chemoattractants originating from the hemocoel and/or stimuli resulting from the interaction of spirochetes with midgut components at or near the intercellular junctions override the inhibitory factor(s) secreted by the midgut epithelium. While the factors and environmental cues promoting the transition to motility remain to be determined, it is important to note that the motile phenotype consistently displayed by organisms within the hemolymph is a biophysical necessity for translation within the fluid-filled hemocoel (58, 72). This point in the dissemination process, therefore, can be thought of as marking the convergence of our model with in vitro (73, 74) and in vivo (63, 64) studies depicting highly motile organisms penetrating cellular junctions and extracellular matrices.

The miniscule numbers of organisms observed on and within salivary glands is in line with reports by others (18, 19, 29, 33) and is consistent with studies indicating that mammalian infection is initiated by a small inoculum of highly virulent organisms delivered to the feeding site (31, 75). Undoubtedly, the paucity of gland-
associated spirochetes reflects, at least in part, the small percentage of organisms that successfully penetrate the midgut and enter the hemolymph. Although I. scapularis hemolymph is poorly borrelial-cidal (76), elimination of spirochetes by cellular innate defenses within the hemocoel, shown herein by us and others (15), may further limit the number of organisms that reach their salivary gland target. Because salivary glands are a well-recognized anatomical barrier for vector-borne pathogens (6, 77–79), it is also plausible that this organ impedes the migration of B. burgdorferi. Along these lines, our visualization of spirochetes in association with the acinar surface as well as with multiple intraglandular structures raises the possibility that, as postulated for the midgut, the salivary glands comprise a succession of obstacles that the spirochete must negotiate via a multistep process in order to attain its goal of accessing the salivary stream. The similar time frames at which spirochetes are detected in the salivary glands and are transmitted to murine tissue, 72 hours here and 60 hours in the extensive time-course analyses conducted by Ohnishi et al. (31), suggest that thinning of the acinar basement membrane may be integral to the spirochete’s strategy for penetration of salivary glands. Invasion of types II and III acini also may be driven by chemotactic factors that are not produced by type I acini (80).

The transcriptional and translational changes that the Lyme disease spirochete undergoes in response to mammalian host- and arthropod-derived signals have been the subject of intensive investigation (12, 21, 23, 24). Abundant evidence has emerged indicating that the Rrp2/RpoN/RpoS pathway serves as a master regulator of the genetic programs required for infection of the mammalian host (19, 35, 51, 81–83). The imaging studies presented here provide a structural framework for interpreting the transcriptional events mediated by RpoS during tick-to-mammal transmission. For example, we (35) and others (31, 44) have demonstrated that the majority of spirochetes within fed nymphal midguts continue to express substantial amounts of both ospA transcript and protein throughout the blood meal. Our model for adherence-mediated migration holds that the protracted down-regulation of ospA mediated by RpoS (51) would maintain spirochetes in close contact with epithelial cells via interaction with TrosPA (37), and possibly other surface receptors, until the midgut barriers weaken. Among those genes upregulated by RpoS in vivo (51), 4 (cheW-2, mcp1, mcp4, and mcp5) encode proteins related to chemotaxis (84), and a fifth (oppA5) may be involved in nutrient sensing (85); expression of one or more of these genes may be

Figure 8
B. burgdorferi are present on salivary glands in low numbers and are observed only within types II and III acini. Representative micrographs of salivary glands isolated from (A) flat or (B–D) fed nymphs at 72 hours after placement on C3H/HeJ mice. Spirochetes (arrowheads) adhering to the surfaces of (B) type III acini and type I (inset). (C) Representative 10-μm composite confocal image showing spirochetes on the surfaces of and within types II and III acini. Spirochetes on the surface (arrowheads), below the surface near the basement membrane (filled arrows), between epithelial cells (open arrows), and within ducts (asterisks) are indicated. (D) Single view taken from a 3D reconstruction showing spirochetes on the surface of (arrowheads), penetrating into (asterisks), and within (arrows) a type III acinus. Note the size difference between type I acini (B, inset), which do not hypertrophy, and the hypertrophied types II and III acini in B–D. Scale bars: 25 μm (A–D); 5 μm (B, inset).

Table 2
Attachment to and invasion of salivary gland acini by B. burgdorferi

<table>
<thead>
<tr>
<th>Time after placement</th>
<th>Acini with surface-associated Bb</th>
<th>Acini containing Bb</th>
<th>Total acini positive for Bb</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Type I</td>
<td>Type II</td>
<td>Type III</td>
</tr>
<tr>
<td>0 h</td>
<td>0 (19)</td>
<td>0 (21)</td>
<td>0 (20)</td>
</tr>
<tr>
<td>24 h</td>
<td>0 (14)</td>
<td>0 (21)</td>
<td>0 (19)</td>
</tr>
<tr>
<td>48 h</td>
<td>0 (12)</td>
<td>0 (53)</td>
<td>0 (40)</td>
</tr>
<tr>
<td>72 h</td>
<td>3 (43)</td>
<td>4 (49)</td>
<td>7 (93)</td>
</tr>
</tbody>
</table>

Values denote number of acini (total examined), AAcini with surface-associated spirochetes following multiple washes, BAcini containing spirochetes near the basement membrane, between epithelial cells, or within acinar ducts. CP < 0.01; CP < 0.003 vs. type I.
critical for the transition from the nonmotile to motile phenotype of organisms that penetrate the hemocoele. The complexity of the spirochete's RpoS-dependent, as well as RpoS-independent, regulatory networks results in an extraordinary degree of transcriptional and antigenic heterogeneity within the feeding midgut (29, 31, 35, 44, 86). A key question for future investigations is whether *Borrelia* reach the hemocoele as a result of a purely stochastic process dependent on the deposition of large numbers of organisms near the basement membrane or by selection for spirochetes expressing an "invasion-competent" transcriptional profile.

**Methods**

*B. burgdorferi* strains, culture conditions, and growth curves. CE162, a virulent clonal derivative of strain *B. burgdorferi* 297 (32), CE863, a CE162 transformant harboring *P* flgB on a pCP32-derived shuttle vector (34), and Bb914, described below, were cultivated in BSK-II medium (87) supplemented with 6% normal rabbit serum (Pel-Freeze Biologicals). Bb914 and CE863 were grown in the presence of gentamycin (50 μg/ml) and kanamycin (400 μg/ml), respectively. Borellial cultures were passaged no more than 3 times prior to experimental manipulations. The plasmid content of each isolate was monitored by PCR as previously described (88). To obtain temperature-shifted organisms, *B. burgdorferi* clones were grown at 23°C to a density of 2 to 5 × 10^8 spirochetes/ml, diluted into fresh BSK-II to a density of 1 × 10^7 spirochetes/ml, and incubated at 37°C until mid-log phase (~5 × 10^7 spirochetes/ml). For determination of growth curves following temperature shift, spirochetes were enumerated over 10–12 days by dark-field microscopy with a Petroff Hauser counting chamber (Hauser Scientific); growth curves for each isolate were performed in triplicate in at least 3 independent trials. Mammalian host-adapted spirochetes were obtained by cultivation of organisms in DMCs implanted into the peritoneal cavities of rats as previously described (89).

Generation of Bb914. The strategy used to generate pMC1916, a suicide vector for insertion of a constitutively expressed *P* flgB reporter and a gentamycin-resistance cassette into the endogenous cp26 plasmid of strain 297, is presented in Supplemental Figure 1A (see Supplemental Table 1 for primers). First, approximately 1 kb each of the upstream and downstream sequences flanking the bb20-bb21 intergenic region were amplified from CE162. The resulting upstream and downstream region amplicons were separately cloned into pCR2.1-TOPO (Invitrogen), yielding plasmids pMC1624 and pMC1626, respectively. The cp26 "crossover region" was generated by ligating the insert from pMC1626 into linearized pMC1624, both digested with AatII and XbaI, yielding plasmid pMC1636. To facilitate subsequent cloning steps, the cp26 crossover region was subcloned out of pMC1636 by digestion with BclIII and ligated into similarly digested pBluescript II (Stratagene), yielding plasmid pMC1661. The polynkinker from the pBluescript II SK vector was amplified using primers containing AatII restriction enzyme sites and then subcloned into AatII-digested pMC1661, yielding plasmid pMC1667. GFP under the control of the borrelial *flaB* promoter was subcloned from pPSB-gfp/pCE320 (88) by digestion with XbaI and Xhol and ligated into similarly digested pMC1667. A fragment containing *P* ospC-aacC1 encoding resistance to gentamycin, was amplified from pBSV2G (90) using primers containing flanking SacI restriction enzyme sites and cloned into SacI-digested pMC1667, yielding the final suicide plasmid construct, pMC1916. Orientations of the *P* ospC-aacC1 and *P* flgB cassettes within pMC1916 were confirmed by sequencing, using primers BB220 junc F and BB221 junc R, respectively. Competent CE162, prepared using the method described by Samuels et al. (91), was electroporated with approximately 20–30 μg of CsCl-purified pMC1916; transformants were selected in 96-well plates in BSK-II containing gentamycin (83). Supplemental Figure 1B depicts the double-crossover event between pMC1916 and the endogenous cp26, which resulted in Bb914. Insertion and orientation of the *P* ospC-aacC1 and *P* flgB cassettes in Bb914 were confirmed by PCR using primer pairs BB220upsp 5′-BssHII plus flgB-aacC1 3′-Sacl and BB222downsp 5′-BssHII plus GFP601-R, respectively.

Effect of FM4-64 on spirochete motility and viability. One ml aliquots of Bb914, cultured to mid-log phase (~5 × 10^7 spirochetes/ml) following temperature shift, were incubated for 24 hours at 37°C in BSK-II in the absence or presence of graded concentrations (2 ng/ml to 100 μg/ml) of FM4-64 (Invitrogen). Following incubation, organisms were assessed for motility by dark-field microscopy and for viability by solid-phase plating in pBSK (91).

SDS-PAGE. Cultures of Bb914 and CE162, before and after temperature shift and following cultivation within DMCs, were harvested by centrifugation at 8,000 g for 15 minutes at 4°C, and the resulting pellets were washed twice with PBS. Whole-cell lysates from 1 × 10^8 organisms were boiled in Laemmli sample buffer (Bio-Rad), separated in 12.5% SDS-polyacrylamide gels, and stained with silver as previously described (32).

**Flow cytometry.** *B. burgdorferi* strains Bb914 and CE863, grown to mid-log phase following temperature shift, were analyzed on a FACSCalibur flow cytometer (BD) as previously described (92). In brief, *B. burgdorferi* cells were stained with the nucleic acid stain SYTO59 (Invitrogen) before fixation in 1% paraformaldehyde in FA buffer (Difco). For each sample, data were collected for 100,000 SYTO59 positive events and analyzed using WinMidi (version 2.8). Three independent experiments were performed for each construct and condition.

Determination of ID₃₀ values for CE162 and Bb914. Groups of 5- to 3-week-old female C3H/Hej mice (Jackson Laboratories) were inoculated intradermally with 1 × 10^4, 1 × 10^5, or 1 × 10^6 temperature-shifted Bb914 or CE162. Infection was assessed at 4 weeks after inoculation by cultivation of ears, joints, hearts, and bladders in BSK-II containing *Borrelia* antibiotic cocktail (Sigma-Aldrich). The percentage of infected animals in each group was used to determine the ID₃₀ values using IDSD0 software, available from the NIH (ftp://ftp.ncbi.nih.gov/pub/spouge/web/software/IDSD0_5.0_Windows), as previously described (93, 94). All experimental procedures using mice were approved by and performed in accordance with the guidelines of the University of Connecticut Health Center Institutional Animal Care and Use Committee.

**Tick acquisition and transmission of B. burgdorferi.** *I. scapularis* larvae (Department of Entomology, Oklahoma State University) and nymphs were stored in an incubator at 22°C over saturated potassium sulfate (95%–97% relative humidity) with a 16-hour light/8-hour dark cycle. Naive larvae were fed to repletion on BALB/c SCID or C3H/Hej mice (Jackson Laboratories) 2 weeks after syringe inoculation with either Bb914 or CE162 (1 × 10^8 spirochetes). Engorged larvae were crushed into 10 μl of PBS for examination by epifluorescence microscopy (GFP filter-475/510 nm), processed for qPCR (see below), or allowed to molt into nymphs. For all transmission experiments, flat nymphs infected with Bb914 or CE162 were fed on naive C3H/Hej mice and processed for microscopy or qPCR as described below; murine infection was confirmed at 4 weeks by cultivation of ears, joints, hearts, and bladders in BSK-II. Nymphs used in these experiments typically attached and established a feeding site within the first 24 hours following placement on mice and were replete by 96–108 hours. Because the actual time of attachment for individual ticks varies, hours indicated are after placement.

**Analysis of OspC expression in Bb914-infected ticks by IFA.** Expression of OspC and FlaB within flat nymphs and nymphs fed for 72 hours on naive mice was determined by double-label indirect immunofluorescence using polyclonal, monospecific rabbit and rabbit anti-serum directed against OspC and FlaB, respectively, as previously described (35). Midguts were mounted in Vectashield (Vector Laboratories) and viewed on a BX41 epifluorescence microscope (Olympus) equipped with FITC and rhodamine.
filters, ×40 (0.65 numerical aperture [NA]) objective, and a Retiga EXi CCD camera (Q Imaging). All epifluorescence images (here and below) were acquired at 696 × 520 pixel resolution. At least 7 unfed and 7 fed nymphs each were examined. The mean percentage (± SD) of OsPCI labeling was calculated based on the number of Flab-labeled organisms.

**Enumeration of spirochetes by qPCR.** DNA was isolated from engorged larvae (pools of 10), unfed nymphs (pools of 15), and replete nymphs (pools of 5) using the Puregene Yeast/Bact. Kit B (QiAGEN) according to the manufacturer’s instructions. Spirochete burdens within infected ticks were assayed using a TaqMan assay directed against flaB [hb0147; Suplemental Table 1] performed using iQ SuperMix (Bio-Rad) in an iCycler thermal cycler (Bio-Rad). Purified plasmid DNA containing the target gene amplicon cloned into pCR2.1 TOPO (Invitrogen) was used to generate a standard curve (10^7–10^2 copies/μl). For each experiment, the standard curve and negative controls were analyzed in duplicate, whereas experimental samples were analyzed in quadruplicate. Copies of flaB present in each sample were calculated using the iCycler post-run analysis software (version 3.1) based on internal standard curves. The means (± SD) for each stage were derived from 3 independent pools.

**Confocal microscopy to determine the distribution of live Bb914 within freshly isolated, nymphal midguts and salivary glands.** Bb914-infected flat nymphs and infected nymphs removed from naive C3H/HeJ mice at 48 and 72 hours after placement were gently washed 3 times with CMRL or BSK-II to remove unbound FM4-64 (2 ng/ml) for 3 minutes at room temperature. The glands then were carefully dissected from the cuticle; specimens showing gross evidence of disruption (i.e., visible leakage of luminal contents) were discarded. Isolated midguts then were transferred to a clean microscope slide and immersed in CMRL or BSK-II containing FM4-64 (2 ng/ml). The tip of a single diverticulum was gently nicked with a no. 11 scalpel and placed on a microscope slide containing 10 μl of CMRL or BSK-II. Midguts then were carefully dissected from the cuticle; specimens showing gross evidence of disruption (i.e., visible leakage of luminal contents) were discarded. Isolated midguts then were transferred to a clean microscope slide and immersed in CMRL or BSK-II containing FM4-64 (2 ng/ml). The tip of a single diverticulum was gently nicked with a no. 11 scalpel and the entire midgut was incubated for 3 minutes at room temperature to allow for uptake of the dye. The midguts were gently washed 3 times with CMRL or BSK-II to remove unbound FM4-64 and transferred to a clean microscope slide containing a drop of Vectashield (Vector Labs). Specimens were viewed on a LSM-510 confocal microscope (Zeiss) equipped with argon and HeNe lasers using a ×40 (1.3 NA) oil-immersion objective within 20 minutes of sample preparation. Optical sections were acquired at 1-μm intervals through the full thickness of midguts isolated from flat and fed nymphs 24 and 48 hours after placement. The large amount of blood combined with the marked expansion of the luminal space, however, precluded imaging through the full thickness (>90 μm) of midguts isolated at 72 hours after placement. Consequently, individual diverticula from 72-hour-fed midguts were incised gently to expose the lumen, and optical sections were acquired from the luminal surface toward the basement membrane. All images were acquired at 512 × 512 pixel resolution and analyzed using ImageJ (95). Composite images, here and elsewhere, comprise serial Z-series images collected in both red and green channels. A minimum of 20 nymphal midguts were examined for each time point.

**Intact salivary glands were carefully dissected from the same cohort of ticks used to obtain midguts and immersed in CMRL or BSK-II containing FM4-64 (2 ng/ml) for 3 minutes at room temperature. The glands then were gently washed 3 times with CMRL or BSK-II to remove unbound FM4-64 and transferred to a clean microscope slide containing a drop of Vectashield; glands were examined within 20 minutes of specimen preparation. Confocal microscopy and image analysis were performed as described above for midguts. A minimum of 20 pairs of salivary glands were examined for each time point. Three-dimensional reconstructions of salivary glands were generated using IMARIS (Bitplane AG) imaging software.**

**Time-lapse confocal imaging of live spirochetes within freshly isolated, intact midguts.** Midguts from Bb914-infected flat nymphs and infected nymphs forcibly removed from naive C3H/HeJ mice at 48 and 72 hours after placement were carefully dissected as above and immersed in CMRL or BSK-II containing FM4-64; for these experiments, specimens were not nicked. After 3 minutes of incubation, the midguts were washed gently 3 times with CMRL or BSK-II to remove unbound FM4-64, transferred to clean, dry microscope slides, and mounted in Vectashield, immersion oil, CMRL, or BSK-II. Confocal microscopy was performed as described above using a ×63 (1.4 NA) oil-immersion objective; only those midgut specimens that excluded FM4-64 were considered to be intact and used for time-lapse imaging. Images (512 × 512 pixel resolution) were acquired at 1-minute intervals for a total of 1.5 hours. A minimum of 7 intact midguts were examined for each time point. Image analysis was performed using ImageJ (95) and QuickTime Pro (Apple) software.

**Time-lapse epifluorescence imaging of live spirochetes within freshly isolated, intact, and nicked midguts.** To assess the motility of spirochetes within the undisturbed midgut environment, midguts from Bb914-infected flat nymphs and infected nymphs forcibly removed from naive C3H/HeJ mice at 72 hours after placement were carefully dissected and immersed, without nicking, in CMRL or BSK-II containing FM4-64. After 3 minutes of incubation, the midguts were washed gently 3 times with CMRL or BSK-II to remove unbound FM4-64, transferred to clean, dry microscope slides, mounted in CMRL or BSK-II, and viewed on a BX41 epifluorescence microscope using a ×40 (1.3 NA) oil-immersion objective; only specimens that excluded FM4-64 were used for time-lapse imaging. Images (696 × 520 pixel resolution) were acquired using a Retiga EXi CCD camera at 8-minute intervals for a total of 30 minutes. Image analysis was performed using ImageJ (95) and QuickTime Pro (Apple) software. A minimum of 7 unfed and 7 fed midguts were examined in either CMRL or BSK-II.

To assess the effect of a breach in the midgut environment on spirochete motility, midguts from Bb914-infected flat nymphs and 72-hour-fed nymphs were carefully dissected and immersed in CMRL or BSK-II. The tip of a diverticulum was nicked with a no. 11 scalpel to expose spirochetes to culture medium, a cover glass was applied, and the specimen was viewed on a BX41 microscope using a ×40 (1.3 NA) oil-immersion objective. Images (696 × 520 pixel resolution) were acquired using a Retiga EXi CCD camera at 5-minute intervals for a total of 30 minutes. The distance of spirochetes from the nick site was determined using an optical reticle (Olympus). Image analysis was performed using ImageJ (95) and QuickTime Pro (Apple) software. A minimum of 10 unfed and 10 fed midguts were examined in either CMRL or BSK-II.

**Gelatin matrix-based assay for spirochete motility.** Gelatin matrices (~1 mm thick) were prepared by adding type A gelatin from porcine skin (5% w/v; Sigma-Aldrich) to prewarmed (56°C) PBS. 200 μl of the liquefied gelatin solution was added to each chamber of a clean-8 well chamber slide (Lab-Tek). The slides were cured overnight at 4°C in a humidified chamber and warmed to room temperature prior to use. 150-μl aliquots of BSK-II containing approximately 5 × 10^5 in vitro–cultivated Bb914 (harvested during logarithmic phase of growth) were added to each chamber, and the slides were incubated for 1 hour at room temperature. The residual medium was gently removed and saved for the enumeration of organisms that did not penetrate or attach to the matrices. Each chamber subsequently was rinsed 3 times with sterile PBS to remove any loosely bound spirochetes. A well (~1 mm diameter) was gently introduced into one corner of each matrix using a glass pipette, and the chambers were washed 3 times with sterile PBS. Each matrix then was gently separated from its plastic chamber using a sterile scalpel and transferred to a new microscope slide. The wells were filled with approximately 1–2 μl of one of the following materials: (a) whole blood, freshly isolated from an uninfected C3H/HeJ mouse; (b) minced midgut from a flat, uninfected nymph; (c) minced midgut from an uninfected nymph 72 hours after placement; or (d) salivary glands isolated from an uninfected nymph 72 hours after placement.
placement. Individual matrices then were covered with a cover glass and viewed on a BX41 epifluorescence microscope using a ×40 (1.3 NA) oil-immersion objective. For each sample, images (696 × 520 pixel resolution) were acquired using a Retiga EXi CCD camera at 5-minute intervals immediately adjacent to (0 mm) and 5 mm away from the periphery of the sample well, for a total of 15 minutes at each distance. Individual spirochetes within a given field of view were scored visually as nonmotile, motile/nontranslational (i.e., movement with no net displacement in the x-y axis), or motile/translational (i.e., movement with net displacement in the x and/or y axes) (see Supplemental Video S for examples). For each sample, a minimum of 500 spirochetes at the 0 and 5 mm distances were categorized; the percentage of spirochetes in each category represents the average of three 5-minute viewing intervals taken over the 15-minute observation period for each distance. A minimum of 3 experiments were performed for each sample. The x-y trajectory plots were generated from time-lapse epifluorescence images during the first minute of viewing using the Multi-Track plug-in of ImageJ (95) software.

Parafin embedding and silver staining of nymphal midguts. Bb914-infected nymphs were placed on naive C3H/HeJ mice and then forcibly removed at 48 and 72 hours after placement. Intact midguts were dissected from the cuticle as described above and fixed overnight at 4°C in 4% paraformaldehyde (pH 7.4) containing FM4-64 (2 ng/ml). The following day, midguts were washed in PBS, soaked in toluene and embedded in paraffin. 10-μm sections were rinsed with PBS and embedded in 1% agarose. Once solidified, excess agarose was trimmed, and the samples were placed into histology cassettes (Shandon Lipshaw). Following dehydration in ethanol, the samples were rinsed in toluene and embedded in paraffin. 10-μm sections were stained using the Warthin-Starry procedure (96) and viewed on a BX41 epifluorescence microscope using a ×40 (1.3 NA) oil-immersion objective. Images were acquired using a Retiga EXi CCD camera at 696 × 520 pixel resolution and analyzed using ImageJ (95).

Cryosectioning of nymphal midguts. Bb914-infected nymphs were placed on naive C3H/HeJ mice and then forcibly removed at 48 and 72 hours after placement. Intact midguts were dissected from the cuticle as described above and fixed overnight at 4°C in 4% paraformaldehyde (pH 7.4) containing FM4-64 (2 ng/ml). The following day, midguts were washed in PBS, soaked overnight at 4°C in 30% sucrose, and then gently placed in Cryomatrix freezing compound (Thermo Shandon). Five-μm cryosections were cut on a Leica SM1900 Cryostat (Leica) equipped with a CryoJane Frozen Sectioning Kit (Instrumedics Inc.) (97). Images were acquired on a LSM-510 confocal microscope using a ×40 (1.3 NA) oil-immersion objective at 512 × 512 pixel resolution and analyzed using ImageJ (95).

Collection of hemolymph. Bb914-infected nymphs, forcibly removed from naive C3H/HeJ mice at 24, 48, and 72 hours after placement, were surface sterilized by successive washes in sterile water, 0.5% bleach, and 70% ethanol.

The front legs were amputated at the distal joints, and the exuded fluid was collected onto a clean, dry microscope slide as previously described (98). Hemolymph was inspected for contamination with blood and/or midgut material by widefield microscopy prior to being imaged by confocal microscopy. The motility of spirochetes within hemolymph was followed over time on a LSM-510 confocal microscope using a ×63 (1.4 NA) oil-immersion objective. Time-lapse images (512 × 512 pixel resolution) were acquired at 5-second intervals for a minimum of 3 minutes per specimen and analyzed using ImageJ (95). To assess spirochete burdens within hemolymph, 15 nymphs were placed in a sterile microfuge tube immediately after the legs were severed at the distal joints and then centrifuged at 8000 g for 5 minutes to collect the hemolymph. A small aliquot of the hemolymph (0.5 μl) was removed and examined by widefield microscopy for the presence of blood and/or midgut contaminants. The remaining hemolymph (~4 μl) was transferred into a sterile microfuge tube for qPCR as described above.

Statistics. The statistical significance of the quantitative differences between flow cytometric sample groups, ID390 calculations, and the localization of spirochetes in salivary gland acini was determined by unpaired Student’s t test with 2-tailed P values using Prism (version 5; GraphPad software). P values of less than 0.05 were considered statistically significant.

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Address correspondence to: Justin D. Radolf, Department of Medicine, University of Connecticut Health Center, 263 Farmington Avenue, Farmington, Connecticut 06030-3715, USA. Phone: (860) 679-8480; Fax: (860) 679-1358; E-mail: jradolf@up.uconn.edu.

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