EFFECT OF RpoN, RpoS AND LuxS PATHWAYS ON THE BIOFILM FORMATION AND ANTIBIOTIC SENSITIVITY OF BORRELIA BURGDORFERI

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Borrelia burgdorferi, the causative agent of Lyme disease, is capable of forming biofilm in vivo and in vitro, a structure well known for its resistance to antimicrobial agents. For the formation of biofilm, signaling processes are required to communicate with the surrounding environment such as it was shown for the RpoN–RpoS alternative sigma factor and for the LuxS quorum-sensing pathways. Therefore, in this study, the wild-type B. burgdorferi and different mutant strains lacking RpoN, RpoS, and LuxS genes were studied for their growth characteristic and development of biofilm structures and markers as well as for their antibiotic sensitivity. Our results showed that all three mutants formed small, loosely formed aggregates, which expressed previously identified Borrelia biofilm markers such as alginate, extracellular DNA, and calcium. All three mutants had significantly different sensitivity to doxycycline in the early log phase spirochete cultures; however, in the biofilm rich stationary cultures, only LuxS mutant showed increased sensitivity to doxycycline compared to the wild-type strain. Our findings indicate that all three mutants have some effect on Borrelia biofilm, but the most dramatic effect was found with LuxS mutant, suggesting that the quorum-sensing pathway plays an important role of Borrelia biofilm formation and antibiotic sensitivity.

Keywords: Lyme disease, biofilm, sigma factor, quorum sensing, mucopolysaccharides, alginate, eDNA

Abbreviations: BSA, bovine serum albumin; BSK-H, Barbour–Stoner–Kelly H; DAPI, 4',6-diamidino-2-phenylindole; DDAO, 7-hydroxy-9H-(1,3-dichloro-9,9-dimethylacridin-2-one; DIC, differential interference contract microscopy; EDTA, ethylenediaminetetraacetic acid; EPS, extracellular polymeric substances; FITC, fluorescein isothiocyanate; PBS, phosphate-buffered saline; RT, room temperature; TE, Tris–EDTA

Introduction

Lyme disease is the most common tick-borne illness caused by the species of bacteria belonging to the genus Borrelia [1]. According the Center of Disease Control within the United States, there are approximately 300,000 reported new Lyme disease cases every year while there are 65,000 people per year reported in Europe [2]. Patients diagnosed with Lyme disease are treated with certain antibiotics; however, recent studies demonstrated those antibiotics insufficient in eliminating certain forms of Borrelia in vitro [3–6]. Furthermore, several clinical [7–12] and in vivo studies [13–18] suggested that there are a potential resistant form which can withhold the antibiotic treatments and the attack of the immune system.

Numerous studies demonstrated that Borrelia burgdorferi can adopt diverse morphologies such as spirochete, round bodies (cysts and granules), and cell deficient forms depending on the condition of the environment they are exposed to [19–23]. These forms provide protective environment for Borrelia for adverse environmental conditions such as exposure to antibiotics, starvation, pH changes, or even high temperature [24–31]. In these defensive forms, B. burgdorferi becomes dormant and remains in this morphological state until it finds a more favorable condition when it returns to its spirochete form [27–31].
Recently, we provided evidences that *B. burgdorferi sensu stricto* and *sensu lato* stains are capable of forming another defensive form called biofilm both *in vitro* and *in vivo* [32, 34]. We also suggested that *Borrelia* biofilm formation could result in increased resistance to various antibiotics [3, 35]. Numerous studies showed that one of the most effective hiding places for a bacterial species is biofilm [36, 37]. Bacterial biofilms are organized communities of cells enclosed in a self-produced hydrated polymeric matrix called extracellular polymeric substances (EPS) which are complex mixtures of polysaccharides, lipids proteins, nucleic acids, and other macromolecules [36–39]. Elimination of pathogenic bacteria in their biofilm form is very challenging because these sessile bacterial cells can endure not just the host immune responses but they are much less susceptible to antibiotics or any other industrial biocides than their individual planktonic counterparts [38, 39]. The biofilm resistance is based upon multiple mechanisms, such as incomplete penetration of certain antibiotics deep inside the matrix and/or inactivation of antibiotics by altered microenvironment within the biofilm and a highly protected resistant bacteria population called persisters [39].

When we evaluated the possibilities that observed *Borrelia* aggregates are indeed biofilm structures, its important biofilm traits like structural rearrangements and changes in development on different substrate matrices as well as the different components of the extracellular protective polymeric surface were studied [32, 33]. Our atomic force microscopic results provided evidence that, at various stages of aggregate development, rearrangements take place at multiple levels that lead to a continuously complex rearranging structure [32]. When the EPS of the aggregates was studied for potential exopolysaccharides, both sulfated and non-sulfated/carboxylated substrates were found; however, the majority was non-sulfated polymucosaccharide alginate. We provided evidence that the *Borrelia* EPS matrix also contains calcium and extracellular DNA [32, 33].

The important molecular pathways of *Borrelia* biofilm development are not known; therefore, to better understand the key components, the potential role of the several important signaling pathways was studied using wild-type and mutant *B. burgdorferi sensu stricto* strain 297.

The first pathway targeted was the RpoN–RpoS alternative sigma factor pathways, which can be found in many bacterial species, and they are involved in various cellular functions in response to different environmental stresses such as adverse temperature, high/low pH, high osmolarity, oxidative stress, high cell density, and carbon starvation [40]. They also control important virulence factors, and they are required for successful infection by many pathogenic bacteria [40]. The RpoN–RpoS pathway in *B. burgdorferi* has similar functions and responsible for sensing the environmental cues and activating RpoN encoded for $\sigma^N$ which then regulates another alternative sigma factor, $\sigma^p$ [41–45]. It was shown that this two-component RpoN–RpoS signal-transduction pathway controls the successful transmission of *B. burgdorferi* from the arthropod vector to the vertebrate host by regulating the expression of several known virulence factors such as outer surface proteins (OspA, OspB, OspC), decorin and fibronectin binding proteins as well as other proteins [44–46]. RpoS-mediated adaptive response is directly regulated by RpoN in *B. burgdorferi*, and together, they regulate over 100 different genes important in survival and stress responses as well as genes involved in the infectious cycles of *B. burgdorferi* [44–46].

Another important global regulatory pathway in bacteria is the quorum sensing bacterial intercommunication system that controls the expression of multiple genes in response to population density [47–50]. The system utilizing small signal molecules is called autoinducers. As the cell population density increases, autoinducers accumulate and cause a population-wide change in the expression of different genes involved in biofilm formation [49, 50]. It was previously demonstrated that *B. burgdorferi* utilizes the autoinducer-2 as one of its many ways of communication [51, 52]; however, one study challenged this observation [53]. The importance of this quorum-sensing pathway in biofilm development was shown for several pathogenic bacteria [54, 55]. For example, a luxS mutant of *Streptococcus gordonii*, a major component of dental plaque biofilm, was unable to form a mixed-species biofilm with another pathogen *Porphyromonas gingivalis* [54]. Furthermore, the luxS mutant of *Streptococcus* spp. shows altered biofilm structure [55]. All these data suggest that LuxS quorum sensing system might play an important role in *B. burgdorferi* biofilm formation.

In this study, different *B. burgdorferi* strains including the wild-type 297 and several mutant strains such as RpoN, RpoS, and LuxS mutants were studied to evaluate the effect of the deletion of these genes on biofilm formations and biofilm specific markers as well as antibiotic sensitivity of *B. burgdorferi* in order to better understand the molecular pathways involved in biofilm formation and its antibiotic resistance. Several previously described *Borrelia* biofilm markers such as sulfated/non-sulfated polysaccharides, alginate, extracellular DNA, and calcium were analyzed using immunohistochemistry and different staining techniques such as Spicer–Meyer, Alizarin, and fluorescent, and extracellular DNA staining methods [32, 33]. The obtained results were visualized using various microscopic techniques such as dark and bright field, fluorescent, and differential interference contrast microscopy. In addition, the antibiotic sensitivity of the different forms (spirochete and biofilm) of the wild-type and mutant *Borrelia* strains was also evaluated.

In summary, the aim of this study to provide insight of the potential molecular pathways regulating biofilm formation in *B. burgdorferi* using different mutant cell lines with the final goal to better understand how *B. burgdorferi* forms biofilms. Data from this study might provide molecular targets in the future to the elimination of *Borrelia* biofilms for therapeutic use.
**Materials and methods**

**Borrelia strains and culture conditions**

Low passages (not more than three passages) of the wild-type 297 and mutant strains (RpoN, RpoS, and LuxS) of *B. burgdorferi* were generously provided by Michael V. Norgard’s research group (Department of Microbiology, University of Texas Southwestern Medical Center, Dallas, Texas). All strains were cultured in BSK-I (Borrelia–Stoner–Kelly H) media with 6% rabbit serum (Peele, then twice with PBS pH 7.4 at RT. The specimens were concentrated hydrochloric acid) was used to stain the fuchsine dye, 6% acetaldehyde in 70% ethanol with 1% 5 min. Aldehyde fuchsine solution (Sigma-Aldrich, 0.5% in double distilled water for 1 min followed by staining with 1% Alcian blue 8GX (Sigma-Aldrich, dissolved in 3% acetic acid, pH 2.5) for 30 min at RT. Then, after rinsing the slides with double distilled water for 3 min, they were dehydrated with graded alcohol and then stained with 2% Alizarin Red-S pH 4.2 (Sigma-Aldrich #A5533) (calcium-specific stain) for 4 min at RT and then were washed twice with double distilled water, dehydrated, and mounted with Permount media.

**Spicer & Meyer mucopolysaccharide staining**

The wild-type 297, RpoN, RpoS, and LuxS mutant strains of *B. burgdorferi* were inoculated in four-well chamber glass slides (Thermo Fisher Scientific, Lab-Tek II chamber glass slides) and were incubated for 7 days at 33 °C and 5% CO₂. For biofilm quantitative assays, 1 × 10⁶ cells/ml of wild-type and mutant strains of *B. burgdorferi* were inoculated in a 48-well tissue culture plates without any antibiotics and incubated for different times as described below.

**Extracellular DNA staining**

To evaluate the extracellular DNA on the aggregates and individual spirochetes, 1 × 10⁶ cells/ml of the wild-type 297 and RpoN, RpoS, and LuxS mutant strains of *B. burgdorferi* were grown in four-well chamber slides for 7 days. The resulting aggregates washed twice with 1× TE buffer pH 8.0. The extracellular DNA was visualized by staining the biofilms with 1 μM DDAO [7-hydroxy-9H-(1,3-dichloro-9,9 dimethylacridin-2-one)] for 30 min at 37 °C in dark. The slides were then washed twice in 1× TE buffer pH 8.0 and mounted with PermaFluor aqueous mounting medium (Thermo Scientific).

**Immunohistochemistry**

Anti-alginate rabbit polyclonal IgG antibody (generous gift from G. Pier, Harvard University) was used to detect alginate expression by the aggregates of the wild-type 297, RpoN, RpoS, and LuxS mutant strains of *B. burgdorferi*. Biofilms were established and cultured above on four-well chamber slides for 7 days. The resulting structures were washed twice with PBS pH 7.4 and fixed in −20 °C using 100% methanol for 10 min and washed then twice with PBS pH 7.4 at RT. The specimens were then pre-incubated with 10% normal goat serum (Thermo Scientific) in PBS/0.5% bovine serum albumin (BSA, Sigma) for 30 min at RT to block nonspecific binding of the secondary antibody. Then, the primary alginate antibody (1:100 dilution in dilution buffer: PBS pH 7.40 + 0.5% BSA) was applied and the slides were incubated overnight at 4 °C in a humidified chamber. After washing, specimens were incubated for ½ h with a 1:200 dilution of DyLight 594 conjugated goat anti-rabbit IgG (Thermo Scientific) at RT. The slides were then washed thrice with PBS/0.5% BSA for 10 min, then incubated at 37 °C for 1 h with FITC-labeled *Borrelia*-specific polyclonal antibody (#73005 Thermo Scientific, diluted 1:50 in 1% BSA/1× PBS, pH 7.4). It was followed by further washing of slides thrice with PBS/0.5% BSA for 10 min at RT and counterstaining with 4’,6-diamidino-2-phenylindole (DAPI) for 10 min. After, then after washing it again with PBS pH 7.4 for 5 min at RT, the specimens were mounted using PermaFluor aqueous mounting medium and the obtained images were analyzed by fluorescent microscopy.

**Alizarin calcium staining**

In order to evaluate the presence of calcium on the surface of *B. burgdorferi* biofilms, 1 × 10⁶ cells/ml of wild-type and mutant strains of *Borrelia* were grown in four-well chamber slides for 7 days, washed twice with PBS pH 7.4, fixed with ice-cold acetone for 5 min, and hydrated with graded alcohol and then stained with 2% Alizarin Red-S pH 4.2 (Sigma-Aldrich #A5533) (calcium-specific stain) for 4 min at RT and then were washed twice with double distilled water, dehydrated, and mounted with Permount media.

**Quantification of B. burgdorferi biofilms by crystal violet and total carbohydrate methods**

The overall mass of the wild-type and mutant *Borrelia* biofilms was quantified by crystal violet methods described earlier [3, 34]. Briefly, seven-day-old aggregates (starting culture of 5 × 10⁶ spirochetal cells) were scraped from the slides and collected by centrifugation method (5000g for 10 min at RT) and were stained with 0.01% (w/v) crys-
Effect of different pathways on *Borrelia* biofilm

European Journal of Microbiology and Immunology

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tal violet 10 min at room temperature. The crystal violet
stain was discarded by centrifugation method (5000g for
10 min at room temperature) and washed twice with sterile
double-distilled water to remove all traces of crystal violet
dye; then, 100 microliter 95% ethanol was added and in-
cubated for 15 min at RT, and absorbance was measured at
595 nm using a BioTek spectrophotometer.

Total carbohydrate assay was also used to detect all
forms of carbohydrates, including simple and complex sac-
ccharides, glycans, glycoproteins, and glycolipids as recent
studies utilized successfully this method to measure and
quantify the formation of extrapoly saccharide layer of the
biofilms as described earlier [32]. Briefly, seven-day-old
aggregates (starting culture of $5 \times 10^6$ spirochetal cells)
were scraped from the slides and collected by centrifuga-
tion method (5000g for 10 min at RT). After removing the
supernatant, pellets were resuspended in double-distilled
H$_2$O (0.2 mL) followed by 5% aqueous phenol (wt/vol,
Sigma) and 0.5 ml of concentrated sulfuric acid (H$_2$SO$_4$)
and incubated for 30 min at RT. The developed yellow/or-
ange color was measured at 485 nm using a BioTek spec-
trophotometer.

*Baclight* (Live/Dead) staining of *Borrelia* aggregates

To visualize live and dead cells for *Borrelia* aggregates,
Baclight Live/Dead viability staining kit (Invitrogen) was
used following manufacturer instructions. Briefly, a 1:1
mixture of SYTO9 and propidium iodide stain prepared
above was added directly to the cultures in the four-cham-
er glass slides. The plate was then covered with alumi-
num foil and incubated for 15 min at RT. The four cham-
bers were separated from the slide, washed, and mounted,
and pictures were taken under the Leica microscope using
fluorescent microscopy.

*Spirochete growth inhibition assay*

For all antimicrobial sensitivity assays, early log-phase
spirochete cultures ($5 \times 10^5$ cells/ml) from the wild-type
and mutant strains were prepared in fresh medium with-
out antibiotics and treated with different concentration of
doxycycline for 48 h of treatment. Spirochete count was es-
timated by direct counting of live/motile spirochetes dark
field microscopy. Six replicates were prepared for each
treatment condition and three independent experiments
were performed. Percentage of control was calculated by
normalization of spirochete counts to the control treatment
(no doxycycline) of the same strain.

*Biofilm microbial sensitivity assay*

Seven-day-old aggregates (starting culture of $5 \times 10^6$ spi-
rochetal cells) in 1 mL suspension were treated with dif-
ferent concentration of doxycycline for 48 h days. Six
replicates were prepared for each treatment condition, and
three independent experiments were performed. After the
treatment period, aggregates are quantified by crystal vio-
et and sulfuric acid-phenol digestion total carbohydrate
assay.

**Statistical analysis**

Each assay was carried out in three independent experi-
ments with six replicates/condition, and numerical average
was obtained for data presentation. Unpaired, two-tailed
TTEST of unequal variance sample was performed to de-
termine $p$ values and statistical difference of two distinct
sample sets. Differences were considered statistically sig-
ificant when $p$ values are <0.05.

**Results**

*The growth pattern of* *B. burgdorferi* wild-type
*and mutant strains*

Evaluation of spirochete growth pattern in the wild-type
and mutant *B. burgdorferi* 297 strains from day 0 to day 8
were performed by using $8 \times 10^5$/ml spirochetes and cul-
tured as described in Materials and methods. The results
of the obtained growth curves showed the standard three-
phase growth: lag, exponential/log, and stationary growth
(Fig. 1). Doubling time of all mutants (RpoN$\Delta$, RpoS$\Delta$, and LuxS$\Delta$) was similar to that of the 297 wild-type strain.

![Fig. 1. The dynamic of the spirochetal growth of the wild-type and mutant *B. burgdorferi* 297 strains. Spirochetes at starting concentration at $8 \times 10^5$ cells/ml were cultured for 8 days as described in Materials and methods, and their cellular growth was evaluated by direct counting method of motile spirochetes using dark field microscopy. Each time point was carried out in three independent experiments with six replicates/condition](image-url)
However, RpoN and RpoS mutants exhibited a delay in the transition from exponential/log phase to stationary growth, resulting in a higher maximum spirochete density: $1 \times 10^8$ compared to $5 \times 10^7$ in wild type. Figure 1 demonstrates that the RpoN mutant entered the stationary growth only on day 6, followed by RpoS mutant and wild-type 297 strains on day 4. On the other hand, LuxS mutant displayed a longer log phase compared to other strains but still had maximal spirochete density that was similar to wild-type strain on day 8.

Analysis of aggregates formed by different mutant strains and wild type of B. burgdorferi strains

In the next experiments, the aggregates produced by the wild-type and mutant strains of B. burgdorferi were analyzed by dark field microscopy. Microscopic pictures of Borrelia strains on glass slides of the four-well chambers after 7 days of incubation of $5 \times 10^6$ cells/ml seeding concentration were compared for the presence of potential aggregate presence. The result showed that aggregates were formed in the 297 wild type as well as all three mutant cultures (Fig. 2). Morphologically, mutants seemed to exhibit a higher tendency to form loose, dispersed, and smaller aggregates than wild-type strain, especially LuxSΔ mutant strains, which showed a reticular mesh structure (Fig 2).

The quantification of aggregates formed by different mutant and wild-type strains was performed using crystal violet and total carbohydrate methods (Fig. 3A and B, respectively). When crystal violet method used to measure the total mass of the different aggregates, there was a significant reduction in the aggregate formation by RpoNΔ strain when compared to the wild-type 297 strain (22% reduction, $p < 0.05$). Comparing RpoSΔ and LuxSΔ aggregates to the wild-type strain, there was a 16% reduction and 39% in the total aggregate masses, respectively ($p < 0.05$). Interestingly, when total carbohydrate method was used to quantify the total carbohydrate component of the different aggregates, no significant difference could be found between the different mutant and wild-type strains (Fig. 3B).

Presence of specific biofilm markers on the surface of the aggregates of different B. burgdorferi strains

Wild-type 297 and various mutant strains of B. burgdorferi were tested for the presence of the previously described Borrelia biofilm specific markers such as sulfated/non-sulfated polysaccharides, alginate, calcium, and extracellular DNA.

1. Sulfated/non-sulfated mucins

Spicer & Meyer staining method was used to examine the presence of sulfated and non-sulfated/carboxylated mucins. Fuchsine stains the weakly acidic sulfomucins; purple coloration signifies strongly acidic sulfomucins, and alcian blue stain is specific for non-sulfated/carboxylated mucins. The aggregates formed by the wild-type and the

Fig. 2. Representative images of aggregates formed by 297-WT (A), RpoNΔ (B), RpoSΔ (C), and LuxSΔ (D) strains of B. burgdorferi as depicted by dark field microscopy. 400× magnification, bar: 200 μm
Effect of different pathways on *Borrelia* biofilm

European Journal of Microbiology and Immunology 277

Different mutant strains (RpoNΔ, RpoSΔ, and LuxSΔ) of *B. burgdorferi* were stained as described in Materials and methods and imaged with dark field microscopy (Fig. 4). The colors developed in the center of the all three mutant

**Fig. 3.** Quantitative analysis of the total mass content of the aggregates formed by 297-WT, RpoNΔ, RpoSΔ, and LuxSΔ using crystal violet method (panel A) or total carbohydrate methods (panel B). Each assay was carried out in three independent experiments with six replicates/condition. *p* values < 0.05 indicates statistical significance as related to the 297-WT control strain.

**Fig. 4.** Spicer & Meyer mucopolysaccharide staining by dark field microscopy of the wild-type 297 (A) and the mutant RpoN (B), RpoS (C), and LuxS (D) *B. burgdorferi* strains. Fuchsia color indicates weakly acidic sulfomucins; purple color indicates strongly acidic sulfomucins/sulfated proteoglycans; blue color indicates non-sulfated/carboxylated mucins. 400× magnification, bar: 200 μm.
strains showed some blue staining pattern (non-sulfated mucin), but contrary to wild-type 297, it was very dispersed. The periphery of the biofilms of all mutant strains studied stained mainly fuchsia purple color indicating sulfated mucins similar to the mucins found in the 297 wild-type strain.

2. Alginate
Spicer & Meyer mucopolysaccharide staining results showed the presence of some non-sulfated mucins on all of the mutant strain biofilms. The non-sulfated mucins could indicate the presence of alginate as described previously for both *B. burgdorferi* B31 and 297 strains [32, 33]. The presence of alginate on the surface of the wild-type and the mutant strains of *Borrelia* species was confirmed by performing a previously published and validated method of double immunohistochemical staining with anti-alginate and anti-*Borrelia* antibodies (Fig. 5A–K). All mutant strains, RpoN, RpoS, and LuxS, were strongly stained with anti-*Borrelia* (Fig. 5D, G, and J) and anti-alginate antibodies (Fig. 5E, H, and K) similar to the wild-type 297 strains (*Borrelia* – green staining, Fig. 5A; alginate – red staining, Fig. 5B, respectively). Similarly to the wild-type strain, only the center of the aggregates stained with al-

![Fig. 5. Immunohistochemical staining of wild-type 297, RpoN, RpoS, and LuxS mutant strains of *B. burgdorferi* with *Borrelia*-specific (green staining, panels A, D, G, and J) and alginate antibodies (red staining, panels B, E, H, and K) as described in Materials and methods. DAPI counterstain (blue staining, panels C, F, I, and L) indicates nuclear staining. Yellow arrowheads indicate spirochetes not stained with alginate. 400× magnification, bar: 200 μm](image-url)
Effect of different pathways on Borrelia biofilm
ginate antibody but not the surrounding spirochetes, indicating the specificity of the alginate staining technique as well as that fact that individual spirochetes do not express alginate (yellow arrow) as described in our previous reports [32, 34].

3. Presence of calcium
It was previously reported that alginate is associated with calcium to form insoluble calcium alginate and calcium was found on the surface of both B. burgdorferi B31 and 297 strains. The potential presence of calcium on the surface of different aggregates formed by the wild-type and the different mutant strains was tested by using Alizarin calcium staining method. The aggregates of all mutant strains RpoN, RpoS, and LuxS were stained red color with Alizarin Red-S stain similar to the wild-type 297 strain, indicating the presence of calcium in all strains studied (Fig. 6, panels A, B, C, and D). As previously reported for the wild-type 297 strain [32], the individual spirochetes were not stained with the calcium stain (indicated in Fig. 6 with black arrowheads), suggesting that calcium is specific to the surface of the aggregates formed by the wild-type and all mutants strains studied.

4. Extracellular DNA
Finally, the aggregates formed by the wild-type and mutant strains were examined for the presence of extracellular DNA (eDNA) as it was described for B. burgdorferi biofilm before [32]. For the detection of eDNA, a red fluorescent dye (7-hydroxy-9H-(1,3-dichloro-9,9-dimethyl-acridin-2-one (DDAO)) was used as reported previously [32]. The aggregate surfaces of the wild-type 297 and the RpoN, RpoS, and LuxS mutant strains were found to have significant amount of eDNA (Fig. 7, panels B, D, F, and H, respectively) whereas the surrounding individual spirochetes did not show staining for eDNA (indicated by yellow arrowheads). The DAPI nuclear counter stain images (Fig. 7, panels A, C, E, and G) depicted the size and morphology of the wild-type and mutant aggregates.

Effect of doxycycline on the wild-type and different mutant strains of B. burgdorferi
The goal of the next experiments is to compare the response of the early log phase spirochetal rich and stationary phase aggregate rich cultures of the wild-type and mutant strains to antibiotic stress. The effects of doxycycline on the spirochetal form of B. burgdorferi in the wild-type 297 and mutant strains (RpoNΔ, RpoSΔ, and LuxΔ) were studied using first early log phase spirochetes and direct counting of the live/motile cells with dark field microscope. Wild-type and mutant spirochetes were treated with various concentrations of doxycycline (ranging from published MIC and MBC concentrations) for 48 h, and the number of motile spirochetes in each treatment was counted. Spirochete numbers were normalized to their no antibiotic control of the same strains (0 μg/ml doxycycline) as depicted in Fig. 8.

Fig. 6. Aggregates of 297 (A), RpoN (B), RpoS (C), and LuxS (D) strains of B. burgdorferi stained with calcium specific stain, Alizarine, as described in Materials and methods. Red color indicates calcium analyzed by dark field microscopy. Black arrowheads indicate spirochetes not stained with Alizarine. 400× magnification, bar: 200 μm.
counts in every concentration were studied. More specifically, as doxycycline concentration increased from zero to 1 μg/ml, spirochetes decreased in number from 100% to 15–20% in every strain. Significant differences between the wild-type 297 and all mutant strains (RpoNΔ, RpoSΔ, and LuxSΔ) were found (marked with #) when 0.1 μg/ml of concentration was used for treatment, while 1 μg/ml doxycycline concentration showed more significant results only in RpoNΔ and RpoSΔ strains but not in the LuxSΔ strain. Interestingly, however, 10 μg/ml doxycycline treatment did not produce significantly different results in any of the mutant strains compared to wild-type *Borrelia* strain.

In the next step of experiments of the crystal violet, total carbohydrate assay and BacLight Live/Dead staining methods were used to study the effect of doxycycline on the viability of the stationary phase aggregates formed by the wild-type and mutant *B. burgdorferi* strains. Crystal violet measured the changes in total mass while total

![Image](image_url)

**Fig. 7.** Aggregates of the wild-type 297, RpoN, RpoS, and LuxS strains of *B. burgdorferi* stained eDNA with DDAO fluorescent dye specific for extracellular DNA as described in Materials and methods (panels B, D, F, and H). DAPI nuclear counterstain was used to demonstrate the structure of the different aggregates (panels A, C, E, and G). Red fluorescent color indicates eDNA as depicted by fluorescent microscopy. Yellow arrowheads indicate spirochetes not stained with DDAO. 400× magnification, bar: 200 μm.
carbohydrate assay was used to detect changes in the extracellular polysaccharide layer of the aggregates before and after antibiotic treatment. Baclight Live/Dead staining method directly visualized the effect of the doxycycline by depicting the live (green) and dead (red) population in the aggregates of the different *Borrelia* strains. Seven-day-old aggregate rich cultures (starting culture of $5 \times 10^6$ spirochetal cells) in 1 mL suspension were treated with different concentration of doxycycline (ranging from 0 to 10 $\mu$g/ml concentration) for 48 h. Data from all antibiotic-treated samples were normalized to their no antibiotic control of the same strains (0 $\mu$g/ml doxycycline). When crystal violet method was used to evaluate doxycycline sensitivity of the wild-type and all three mutant strains, there were no significant differences found for any of the experimental condition for any strains studied (data not shown). However, when the total carbohydrate assay was used to assess doxycycline sensitivity, there were significant differences found in all three mutant strains compared to the wild-type control as depicted in Fig. 9. While doxycycline treatment of 297 wild-type strain did not reduce the extracellular polysaccharide layer significantly at any of the antibiotic concentrations, all three mutant strains (RpoN $\Delta$, RpoS $\Delta$, and LuxS $\Delta$) showed reduction of their extracellular polysaccharide layer at 10 $\mu$g/ml (20–60%). The most dramatic effect, however, was found in the doxycycline sensitivity of the LuxS $\Delta$ strain, indicating that all three concentrations of doxycycline (0.1, 1, and 10 $\mu$g/ml) significantly reduced the extracellular polysaccharide layer (Fig. 9). The reduction was significant when it was compared to both the LuxS $\Delta$...
control and 297 wild-type control \((p \text{ value } < 0.01)\). Furthermore, while the effect of 10 \(\mu\text{g/ml}\) of doxycycline was significant for the extracellular polysaccharide layer of the RpoN\(\Delta\) and RpoS\(\Delta\) strains when it compared to their no treatment controls \((p \text{ value } < 0.05)\), it was not significantly different than the wild-type control data with the same doxycycline treatment.

To further confirm the quantitative data of the doxycycline effect on the different strains, Baclight Live/Dead analyses were performed on the 10 \(\mu\text{g/ml}\) doxycycline-treated cultures after 48 h. Figure 10 shows representative microscopic images obtained from the untreated and doxycycline-treated wild-type 297 and mutant \textit{B. burgdorferi} strains (RpoN\(\Delta\), RpoS\(\Delta\), and Lux\(\Delta\)) strains depicting membrane-intact (live) cells in green and membrane-permeable (dead) cells in red. In all of the no treatment controls (Fig. 10, panels A, C, E, and G), majority of the cells show green staining indicating live cells with a small portion of the cells stained red indicating dead cells. In the doxycycline-treated cultures, the majority of the cells stained green for the wild-type 297 and the RpoN\(\Delta\) mutant cells show no significant difference from their no treatment control cultures. For RpoS\(\Delta\) strain, there were a portion of the aggregates (<10%) which showed higher numbers of dead cells, but overall, the difference from the no treatment control was not significant. On the other hand,

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**Fig. 10.** Representative Baclight Live/Dead images of the effect of doxycycline on the aggregate forms of wild-type 297 (panels A and B) and RpoN\(\Delta\) (panels C and D), RpoS\(\Delta\) (panels E and F), and Lux\(\Delta\) (panels G and H) strains of \textit{Borrelia burgdorferi} after 48 h treatment with 0 (panels A, C, E, and G) or 10 \(\mu\text{g/ml}\) (panels B, D, F, and H) of doxycycline as analyzed using fluorescent microscopy. Green color = live cells, red color = dead cells, 400x magnification, bar: 200 \(\mu\text{m}\)
majority (>95%) of the aggregates of the LuxS strains showed mainly red/dead cells after 48 h of antibiotic treatment, suggesting a greater sensitivity to doxycycline than the wild-type 297 or the Rpo gene mutant strains.

Discussion

In this study, the possible biofilm formation of several mutant strains of B. burgdorferi was investigated to better understand the roles of different genes—intracellular pathways in the development and characteristic of Borrelia biofilm.

Building a biofilm community requires intra- and extracellular communication processes and is considered a survival response in bacteria [36–38]. Rpo transcriptional factor genes are known to control bacterial changes in response to environment stimuli [40]; LuxS quorum-sensing roles in biofilm formation and antibiotic response have been established in some research with other organisms [47–50]. This study attempted to explore the possible roles of Rpo genetic regulation and LuxS quorum communication pathways in Borrelia biofilm formation and its antibiotic sensitivity.

Growth patterns of planktonic spirochetes were compared between the wild-type and the different mutant strains. The results showed that RpoN, RpoS, and LuxS mutant strains had some deviations from wild-type strains, including that mutants lacking Rpo genes had longer time to response to the inadequate nutrient environment, resulting in a longer exponential growth period and, ultimately, the higher maximum spirochete density. LuxS mutant reached the stationary phase later, suggesting that LuxS could also have a potential role in the growth of B. burgdorferi.

Furthermore, in spite of lacking important genes responsible for regulating bacterial environmental responses and quorum-sensing communication, all three mutants readily formed biofilm-like aggregates in the stationary phase of growth. Morphologically, all mutants but especially the LuxS mutant seemed to exhibit a higher tendency to form smaller and looser aggregates than the wild-type strain. This result is in good agreement with a previous study in which mutation in rpoS did not eliminate the biofilm formation in Escherichia coli during exponential phase of growth [54, 55].

The size of the aggregates was also quantified in this study using crystal violet staining as well as total carbohydrate quantitation methods. The obtained data indicated that mutant aggregates were significantly smaller than the wild-type ones; however, those smaller aggregates had similar amounts of total carbohydrate, suggesting that the mutants might produce higher amounts of protective mucopolysaccharide layers. This finding is in good agreement with a study, in which Pseudomonas aeruginosa mutant had increased matrix production [56].

Further investigation of specific biofilm markers in the mutant strains showed that all three mutant strains produce significant amounts of biofilm specific markers such as sulfated/non-sulfated polysaccharides, alginate, calcium, and eDNA, similar to what was found and reported previously to the wild-type B. burgdorferi strains [52, 53]. These results strongly suggest that the aggregates formed by the mutant strains studied are indeed biofilms.

It might be surprising at first that Borrelia strains lacking RpoN, RpoS, and LuxS genes still can form biofilms which could have most of the main phenotypes of the wild-type biofilm. However, in earlier studies, it was reported that mutant strains of RpoN in E. coli K12 and Enterococcus faecalis also formed biofilms [57, 58]. RpoS mutant strain was also observed to form biofilm in E. coli ZK126 stains as well as in P. aeruginosa [59, 60]. For LuxS mutant strain in Streptococcus sp., a report suggested that the mutant LuxS mutant could form biofilm; however, it depends on the in vitro culture condition [61, 62].

One of main differences found for the Borrelia mutant biofilms was the reduced size compared to wild-type 297. The RpoN mutant strain of B. burgdorferi formed a biofilm which was 22% less in size than the wild-type 297. Previously, it was reported that the deletion of RpoN gene in E. coli K12 strains increased the biofilm formation by 40–60%, and also in E. faecalis, the RpoN mutant strain was observed to form robust biofilms [57, 58]. In this study, we showed that RpoS mutant strain of B. burgdorferi had 10% reduction in the biofilm formation compared to wild-type similar to the deletion of RpoS gene in E. coli K12 which reduced the biofilm size by 50% [59]. LuxS mutant strain of B. burgdorferi formed biofilm which showed 39% significant reduction in size which agrees with the report on Streptococcus sp. LuxS mutants [61].

Also, there were slight morphological differences observed in the biofilms formed by the RpoN, RpoS, and LuxS Borrelia mutant strains compared to wild-type strain. The biofilm formed by the wild-type 297 strain shows a compact biofilm mass which resembles the biofilm of B31 strain of B. burgdorferi as described in our recently published paper [32, 33]. The biofilm formed by RpoN mutant was relatively less compact compared to wild-type biofilms while RpoS mutant strain formed many loose and dispersed small aggregates unlike the wild-type 297. Interestingly, the biofilm formed by RpoS mutant of P. aeruginosa was denser and thicker than the wild type [60]. The biofilm formed by LuxS mutant strain of Borrelia represents a very loose reticular mesh. Undifferentiated and loosely-connected biofilms were also found to form in LuxS mutant strain of Shewanella oneidensis covering the entire glass surface uniformly unlike the wild type which forms compact biofilms with significant amounts of spaces in between [63].

Biofilms were demonstrated to be responsible for the antibiotic resistance in many species [36–39]. Therefore, besides examining the growth dynamics and biofilm development of the different mutant strains of B. burgdorferi, this research attempted to compare the responses of wild-type and mutant strains to antibiotic stress. Early log phase spirochetes and stationary phase aggregate
rich cultures were exposed to varying concentrations of doxycycline and compared among all strains studies. In a good agreement with several recent studies using wild-type B. burgdorferi 31 strain, only the early log phase spirochetes of the wild-type 297 strains were sensitive to doxycycline treatment but not the aggregate rich stationary phase cells [3–6]. When the doxycycline sensitivity of the early log phase spirochetes of the wild-type strain and all of the three mutant strains was compared, the obtained results indicated significantly higher sensitivity of all three mutant strains to low MIC dose of doxycycline (0.1 μg/ml) than the wild-type strain. Interestingly, however, there was no difference in the doxycycline sensitivity of the early log phase spirochetes among the strains at higher MBC level concentration.

Similarly, doxycycline sensitivity of the stationary aggregate rich cultures of wild-type and Rpo mutant was not significantly different at lower doses. In contrary, however, antibiotic sensitivity of the LuxS Borrelia mutant dramatically greater than the wild-type mutant at all concentrations was studied. These later results were also confirmed by microscopical analyses of the live and dead cells of the mutant strains as demonstrated by BacLight Live/Dead staining. In summary, the antibiotic sensitivity studies of the mutant strains confirmed that all three genes have some effect on antibiotic sensitivity in some extent but LuxS mutant has the most significant effect.

Involvement of LuxS signaling pathway in antibiotic sensitivity was proven for S. anginosus species, in which the mutant demonstrated increased susceptibility to erythromycin and ampicillin [64]. Furthermore, P. aeruginosa mutants defective in interspecies quorum sensing produced weaker biofilms that were more sensitive to detergents, while Streptococcus mutans mutants defective in interspecies cell signaling generated stronger biofilms that were more resistant to detergents [61, 65].

In our recent studies, we also investigate the potential biofilm formation of several other B. burgdorferi mutants such as the Hk1/Rrp1 and recA mutants that regulate the cyclic di-GMP and the recombination events, respectively [38, 66, 67]. Those two molecular pathways were already implicated to be important in biofilm development and characteristic (data not shown).

In summary, our findings strongly suggest that several alternative pathways could regulate B. burgdorferi biofilm formation, a result that indicates that it is a very important survival mechanism for this bacterium. Findings for the potential importance of the LuxS quorum-sensing pathways for Borrelia biofilm development and antibiotic sensitivity merit further investigation of this pathway.

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**Competing interests**

The authors have declared that no competing interests exist.

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European Journal of Microbiology and Immunology
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