Studies on the transovarial transmission of *Borrelia burgdorferi* sensu lato in the taiga tick *Ixodes persulcatus*

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**Abstract.** The possibility of vertical transmission of *Borrelia burgdorferi* sensu lato in *Ixodes persulcatus* Schulze, 1930 ticks was studied in the progeny of 20 females collected from the vegetation in an active focus of ixodid tick-borne borrelioses (ITBB) located in the Perm oblast, Russia, where *Borrelia garinii* and *B. afzelii* are circulating. The presence of *Borrelia* DNA was detected by the PCR method after feeding and egg laying in 16 engorged females (80.0%), as well as in 36.5 ± 7.2% samples containing 20 eggs each and in 21.4 ± 4.2% samples containing 10 eggs each. The respective rates of individual egg infection were 0.4–8.0% and 0.5–23.0%. PCR analysis of 370 eggs (one egg per sample) and 781 unfed larvae hatched from the same egg masses (1, 10, 20, 40, and 50 larvae per sample) failed to reveal the presence of *Borrelia* DNA. Negative results were also obtained in experiments on inoculating the BSK II medium with the egg and larval materials. Microscopic analysis of 1,683 smear preparations of eggs and 1,416 preparations of unfed daughter larvae revealed spirochete-like cells in 7 (0.4 ± 0.3%) and 13 (0.9 ± 0.5%) preparations, respectively; typical *Borrelia* cells were found in seven preparations of larvae (0.5 ± 0.4%). Only 1 out of 16 infected females transmitted *Borrelia* vertically, through the eggs to the larval progeny. The infection rate in this progeny was about 7%, and the prevalence of *Borrelia* in individual larvae was 0.4–0.8 cells per 100 microscopic fields. These data do not allow the conclusion that transovarial transmission of *B. burgdorferi* sensu lato in the *I. persulcatus* tick is an established fact. However, they show that, even if such transmission is possible, its probability is very low.

**MATERIALS AND METHODS**

Adult ticks were collected from the vegetation by flagging in the Pre-Ural region of Russia (Tchusovskii district, Perm oblast) in May and June 1999. Long-term observations in this region have shown that *Borrelia garinii* and *B. afzelii* circulate there (Korenberg et al. 1999), and the infection rate among unfed adult ticks (estimated by dark-field microscopic analysis) varies in different years from 29.6 to 60.8% (Korenberg et al. 1999). According to the results of parallel analysis of 125 unfed female ticks by means of dark-field microscopy and PCR (see below), the infection rate in the corresponding biotope at the time of sampling was estimated at 52.0% by the first method and 64.0% by the second method.

Male and female *I. persulcatus* ticks were kept before experiments in separate tubes at 4°C. Twenty females were allowed to feed individually, in the presence of a male, on outbred white mice. After feeding, they were kept in individual tubes with a humidity gradient. Upon completing egg laying, each female was analysed for *Borrelia* infection by the PCR method.

On the whole, the females laid 19 egg masses, and the larvae hatched out of 17 egg masses. The eggs and unfed larvae taken from different parts of each egg mass were studied by several methods (Tables 1, 2).

**Culture method.** The pools of 50 eggs or 10 larvae were washed with 70% ethanol (less than 2 min for the eggs and approximately 5 min for the larvae), sterile distilled water, and a solution of 0.9% NaCl (2 min each) in sterile Petri dishes and were used for inoculating the BSK II medium without...
antibiotics in 2-ml sterile polystyrene tubes filled to three-fourth of their capacity. The eggs were either homogenized in 200 µl of the medium or simply placed in the tube and punctured with a dissection needle. In the larvae, the gnathosoma was aseptically removed under a microscope, and the body was transferred to the cultivation tube (Gorelova et al. 1996). The tubes were closed with airtight screw caps and incubated at 32°C for two months. Every week, the cultures were tested for the presence of *Borrelia* by analysing test samples under a microscope with a dark-field condenser at a magnification of 600× (100 microscopic fields per culture).

**Microscopic analysis of fixed smears.** The contents of an egg crushed with a dissecting needle were smeared on a glass slide. The preparations of unfed larvae and the ovaries of unfed females were made as described by Levin et al. (1993). The smears were dried, flame-fixed over an alcohol burner, and stained using the Giemsa stain according to Romanovskii and a solution of 1% Crystal Violet for 30 min. In each preparation, 250 microscopic fields were examined at a magnification of 600× to determine the concentration of *Borrelia*, which was expressed as the number of spirochetes per 100 microscopic fields.

**PCR analysis.** The biological samples were washed in 70% ethanol and distilled water, dried on absorbent paper, placed in 1.5-ml Eppendorf tubes, and stored before analysis at −70°C. *Borrelia* genomic DNA was isolated using guanidine thiocyanate treatment and phenol-chloroform extraction according to Gushchin and Khalilov (1996) (see Nefedova et al. 2001). When necessary, DNA samples were stored at −20°C. DNA amplification was performed as described by Salinas-Melendez et al. (1995), with some modifications, using the conservative primers BB1 and BB2 flanking a 244-bp fragment of the *B. burgdorferi* 16S rRNA gene (Adam et al. 1990). The reaction mixture (25 µl) in 10 × buffer (67 mM Tris–HCl, pH 8.6; 16.6 mM (NH₄)₂SO₄; 2.5 mM MgCl₂; 0.001% Triton X-100; and 0.12 mg/ml BSA) contained the four deoxynucleoside triphosphates (dATP, dCTP, dGTP, and dTTP) at a final concentration of 0.2 mM, 15 pmoles of each primer, 1.5 units of Taq polymerase, 10 µl of the test DNA sample, and a drop of mineral oil. In the “blank” control, deionized water instead of DNA was added. The procedure of DNA amplification involved five cycles of reaction at 94, 40, and 72°C followed by 30 cycles at 94, 50, and 72°C (1 min at each temperature) in a four-channel ThermoCycler thermocycler (NPO DNK Tekhnologiya, Russia). To estimate the specificity of amplification, a set of DNA samples from different spirochete species was included in analysis. To this end, we used the isolates of spirochetes of different groups kept at the Museum of *Borrelia* Cultures of the Vector Laboratory, Gamaleya Research Institute for Epidemiology and Microbiology, Russian Academy of Medical Sciences:

(a) The *B. burgdorferi* sensu lato group was represented by *B. afzelii*, type strain Ip-21 (derived from an adult *I. persulcatus* tick caught in Leningrad oblast in 1987); *B. garinii*, isolate Ipl-4906 (from an engorged *I. persulcatus* larva caught in Perm oblast in 2000); *B. burgdorferi* sensu stricto, strains B 31 ATCC 35210 (isolated from an *I. dammini* tick in the United States in 1982, kindly supplied by Dr. A. Barbour) and TxGW (isolated from a diseased person in the United States in 1986, kindly supplied by Dr. R. Johnson); and *B. valaisiana*, isolate Ir-4804 (from an *I. ricinus* tick caught in Krasnodar krai in 2000).

(b) Spirochetes of the group causing tick-borne relapsing fevers included *B. persica*, strain Batyr HT-20 (isolated from *Ornithodoros papillipes* ticks in Uzbekistan in 1988); *B. recurrentis* (the blood of mice infected with the culture kindly supplied by Prof. J. Chalupsky, Czech Republic); and *B. hermsii*, serotype C (isolated from the blood of a diseased teenager in the United States in 1990, kindly supplied by Dr. A. Barbour).

(c) The *Leptospira* group was represented by the *L. biflexa* serovar patoc (I) (from the Museum of the Leptospirosis Laboratory, Gamaleya Research Institute for Epidemiology and Microbiology, Russian Academy of Medical Sciences).

The resulting PCR products were compared by electrophoresis, with the GeneRuler™ 100 bp DNA Ladder marker kit (MBI Fermentas, Lithuania) used for determining DNA length. No reaction with samples containing other spirochetes than *B. burgdorferi* sensu lato was observed, confirming that PCR amplification of the test samples was 100% specific. The detection limit of the method was approximately 100 spirochetes per sample. To correctly interpret the results of PCR analysis, we performed additional experiments with the DNA of *B. afzelii* strain Ip-21 (the positive control) and the DNAs of a female *I. ricinus* from the laboratory colony of uninfected ticks (Institute of Zoology, Switzerland; kindly supplied by Dr. L. Gern), its eggs, and a larva hatched from the same egg mass (the negative controls). The products of PCR amplification, including the blank control (see above), were compared by means of agarose gel electrophoresis in horizontal slabs in the presence of ethidium bromide (1–2% agarose in Tris-acetate buffer, pH 8.1; 165 V). The resulting electrophoretic patterns were analysed using a DNA Analyzer videosystem with the Gel-Imager and Gel-Analysis version 1.0 software programs (Russia). The result of PCR was regarded as positive or negative depending on the presence or absence of an amplification product corresponding to the specific fragment of the control DNA preparation.

**RESULTS**

Spirochetes were found in the ovaries of 3 out of 96 unfed females (3.1 ± 3.5%) studied by means of microscopic analysis of fixed smears, and the presence of *Borrelia* DNA was detected by the PCR method in 16 out of 20 engorged females, which was evidence for the possibility of transovarial pathogen transmission. The females laid a total of 19 egg masses. The DNA of *Borrelia* was detected in approximately one-fifth of 10-egg pools and one-third of 20-egg pools obtained from the engorged females, with the difference in infection rate between the pools being statistically significant (Student’s t-test, t = 3.6). The data on egg infection obtained by the group suspension method (Table 1) were used for calculating the probable rate of individual egg infection for an egg mass by the formula proposed by Beklemishev (1963).

The rates calculated from data on the pools of 10 and 20 eggs were 0.5–23.0% and 0.4–8.0%, respectively. However, the results of

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1. $P_e = e^{-\frac{M}{n}}$, $M = -\ln P_e$, where $P_e$ is probable infection rate, $e$ is the base of natural logarithm, and $M$ is the average number of infected ticks (eggs) per batch in the total sample.
Table 1. Rates of Borrelia infection of eggs laid by infected Ixodes persulcatus ticks as estimated by different methods.

<table>
<thead>
<tr>
<th>Method</th>
<th>No. pools / eggs studied</th>
<th>infected/ examined</th>
<th>typical Borrelia</th>
<th>% (P ± 2mp)</th>
<th>spirochete-like cells</th>
<th>% (P ± 2mp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cultivation (pools of 50 eggs)</td>
<td>12 / 600</td>
<td>0</td>
<td>0</td>
<td>5 / 12^1</td>
<td>41.7 ± 28.5 / −</td>
<td></td>
</tr>
<tr>
<td>Microscopy of fixed smears</td>
<td>− / 1,683</td>
<td>0</td>
<td>0</td>
<td>7 / 1,683^2</td>
<td>− / 0.4 ± 0.3</td>
<td></td>
</tr>
<tr>
<td>PCR analysis of individual eggs</td>
<td>− / 370</td>
<td>0</td>
<td>0</td>
<td>−</td>
<td>−</td>
<td></td>
</tr>
<tr>
<td>samples of 10 eggs</td>
<td>378 / 3,780</td>
<td>81 / 378^1</td>
<td>21.4 ± 4.2 / −</td>
<td>−</td>
<td>−</td>
<td></td>
</tr>
<tr>
<td>samples of 20 eggs</td>
<td>178 / 3,560</td>
<td>65 / 178^1</td>
<td>36.5 ± 7.2 / −</td>
<td>−</td>
<td>−</td>
<td></td>
</tr>
</tbody>
</table>

^1examined by pools; ^2examined individually

Table 2. Rates of Borrelia infection of unfed Ixodes persulcatus larvae as estimated by different methods.

<table>
<thead>
<tr>
<th>Method</th>
<th>No. pools / larvae studied</th>
<th>infected/ examined</th>
<th>typical Borrelia</th>
<th>% (P ± 2mp)</th>
<th>spirochete-like cells</th>
<th>% (P ± 2mp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cultivation (pools of 10 larvae)</td>
<td>60 / 600</td>
<td>0</td>
<td>0</td>
<td>16 / 60^1</td>
<td>26.7 ± 11.4 / −</td>
<td></td>
</tr>
<tr>
<td>Microscopy of fixed smears</td>
<td>− / 1,416</td>
<td>7 / 1,416^2</td>
<td>− / 0.5 ± 0.4</td>
<td>13 / 1,416^2</td>
<td>− / 0.9 ± 0.5</td>
<td></td>
</tr>
<tr>
<td>PCR analysis (total)</td>
<td>− / 781</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

^1examined by pools; ^2examined individually

inoculating culture medium with the egg material were negative, and none of 370 eggs individually analysed by the PCR method proved to contain Borrelia DNA (Table 1).

A total of 781 unfed larvae hatched from the infected egg masses were tested for the presence of Borrelia DNA either individually or 10, 20, 40, and 50 larvae in a sample. The results were negative in all cases. The same concerns the results of cultivation experiments (Table 2).

The cultures of immobile spiral forms of Bacillus sp., described earlier by Heidrich et al. (1999), were obtained from the eggs of 3 out of 6 females (i.e., 3 out of 6 egg masses used for inoculation) and the larval progenies of 7 out of 15 females. Spirochete-like cells were also found in some microscopic preparations of eggs, and morphologically similar cells, in the preparations of unfed larvae of the daughter generation (Tables 1, 2). Only 1 out of 16 infected females transmitted Borrelia vertically, through the eggs to the larval progeny. However, spirochete-like cells were not detected in fixed smears prepared from the eggs of this female and the larvae hatched from the same egg mass. The infection rate in this progeny was about 7%, and the prevalence of Borrelia in individual larvae was 0.4–0.8 cells per 100 microscopic fields. Typical Borrelia cells were detected in only about 0.5% of larvae from the progeny of infected female ticks.

DISCUSSION

Transovarial transmission of viable Borrelia burgdorferi sensu lato can be confirmed only by inoculating the BSK medium with the test material. In this context, it should be noted that viable Borrelia isolates were obtained, in very rare cases, from unfed Ixodes ricinus larvae collected in North Africa (Younsi et al. 2001), Amblyomma americanum in the United States (Schulze et al. 1986), and Ixodes persulcatus in Russia (Gorelova, unpublished). Single isolates were also derived from the biopsies or blood of laboratory animals used as hosts for I. ricinus larvae collected in natural foci (Stanek et al. 1986, Hammer et al. 2002) or engorged larvae from the progenies of ticks infected in the laboratory (Krampitz 1986). Using dark-field microscopy, live Borrelia were detected in the unfed larvae of the following tick species: Ixodes scapularis (Bosler et al. 1983, Piesman et al. 1986), A. americanum (Schulze et al. 1986), I. ricinus (Hubálek and Halouzka 1998), and I. persulcatus (Balashov and Grigor’eva 1997, Balashov et al. 1998). These data are difficult to compare and impossible to evaluate quantitatively.

The results of this study showed that the eggs laid by spontaneously infected female I. persulcatus ticks contained the genomic DNA of Borrelia; therefore, the transovarial transmission of Borrelia in these ticks was principally possible. However, the larvae hatched from
these eggs either contained no *Borrelia* DNA or its concentration was too low to be detected by our PCR system. This fact may be evidence for the loss of the pathogen in the course of egg development and the hatching of larvae. Typical *Borrelia* were found in only a few preparations of the larvae hatched from the same egg mass. Of 16 infected female ticks, only the female that laid these eggs was found to transmit *Borrelia* transovarially to its progeny. The infection rate among these larvae was only 7%, with the prevalence of *Borrelia* being also very low: 0.4–0.8 spirochetes per 100 microscopic fields.

Thus, the above data suggest that the transovarial transmission of *B. burgdorferi* sensu lato in the *I. persulcatus* tick – the main vector of this pathogen in Eurasia – is possible, but its probability is very low. This transmission pathway apparently plays no significant role in the maintenance of *Borrelia* circulation and the dynamics of parameters of infection in adult ticks of the next generation.

There are strong reasons to believe that the functioning of natural foci of ixodid tick-borne borrelioses (ITBB) in Russia depends primarily on the horizontal and transstadial pathogen transmission, with the determinative effect on the prevalence of infection among unfed adult *I. persulcatus* ticks belonging to the parameters of infection of engorged nymphs. Thus, studies on the long-term dynamics of epizooctic activity in natural foci revealed consistent and virtually synchronous changes in the population density and infection rate of adult ticks, on the one hand, and in the abundance of forest voles of the genus *Clethrionomys*, on the other. The coefficients of correlation between the abundance of these rodents in the previous year and the rate of tick infection by *Borrelia* in the current year were $r = 0.73$ for nymphs and $r = 0.84$ for adult ticks. These dynamics of the epizooctic activity of natural ITBB foci in the Pre-Ural region eventually determine the long-term dynamics of their epidemic manifestation (Korenberg et al. 2002).

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