Effect of salivary gland extract from *Ixodes ricinus* ticks on the proliferation of *Borrelia burgdorferi* sensu stricto in vivo

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Abstract. Saliva-activated transmission (SAT) of *Borrelia burgdorferi* sensu stricto was demonstrated using real-time PCR and salivary gland extract (SGE) from partially fed *Ixodes ricinus* ticks. C3H/HeN mice were injected intradermally with $1.5 \times 10^7$ spirochetes mixed with 40 µg of SGE per mouse. The control group was inoculated with the same dose of spirochetes without SGE. The accelerating effect of SGE on spirochete proliferation was demonstrated on day 1 post infection, when a 4.2-fold increase in spirochetes was found in the skin and a 10-fold increase in the blood, compared with control mice. The data represent the first direct evidence of a SAT effect of *I. ricinus* SGE on infection with the Lyme disease agent *B. burgdorferi*.


Feeding of ixodid ticks extends over a number of days and even as long as three weeks. The time required for a tick to obtain a blood-meal provides ample opportunity for interaction with the host immune system. Immune responses of naive hosts involve inflammatory reaction, complement activation and cytokine production (Kopecký and Kuthejlová 1998), preceding activation of adaptive immune mechanisms. Ticks have developed a number of mechanisms that operate against the host immune response. Immunomodulatory molecules present in tick saliva or in salivary gland extract (SGE) affect both innate and adaptive immunity. Tick saliva and/or SGE inhibit activation of the alternative pathway of complement (Ribeiro 1987) and anaphylatoxin activity (Ribeiro and Spielman 1986). Phagocytosis, and production of superoxide by neutrophils and nitric oxide by macrophages, are reduced by tick saliva (Ribeiro et al. 1990, Urioste et al. 1994, Kuthejlová et al. 2001). Tick saliva and/or SGE cause a decrease in inflammatory cytokines IL-1, TNF-α, IL-2, IFN-γ and upregulation of anti-inflammatory cytokines IL-4, IL-10 and TNF-β (Ramachandra and Wil kel 1992, Fereira and Silva 1999, Kopecký et al. 1999, Schoeller et al. 1999, Kuthejlová et al. 2000). The immunomodulatory effect of tick saliva was often studied using SGE, due to the difficulties in obtaining the saliva in sufficient amount. It is obvious that SGE contains also cellular proteins that are not secreted in the saliva. Sometimes, these proteins can influence the host immune response, so that it is different from that affected by tick saliva (Severinová et al. 2005).

Accumulating evidence indicates that immunomodulatory molecules in tick saliva or SGE can promote transmission and survival of *B. burgdorferi* spirochetes in mice (Pechová et al. 2002). This phenomenon is supported by the fact that tick saliva and/or SGE polarize the host immune response to a T-helper 2-type cytokine profile which causes inhibition of spirochete killing, consequently promoting their proliferation. Facilitation of tick-borne pathogen transmission, via the actions of tick saliva on the host, has been termed saliva-activated transmission (SAT). This phenomenon was described as the enhancement of Thogoto arbovirus infectivity when it was inoculated into the host together with SGE (Nut tall and Jones 1991). Existence of SAT can be demonstrated by direct and indirect evidence. Observations of efficient transmission of *Borrelia* spirochetes between cofeeding ticks without a systemic infection of the host represent indirect evidence of SAT (Ogden et al. 1997). Directly, SAT is demonstrated by enhanced transmission of spirochetes to ticks feeding on a host syringe-inoculated with the pathogen plus SGE (Pechová et al. 2002).

Non-bacteraemic transmission has been reported for *B. burgdorferi* s.s., *B. garinii* and *B. afzelii* (Cern and Rais 1996, Patrician 1997, Sato and Nakao 1997, Richter et al. 2002). *Borrelia burgdorferi* s.s. strain B31 showed SAT with SGE from *Ixodes scapularis* (its natural vector species in USA), but not with SGE from *I. ricinus* (Zeidner et al. 2002). We investigated the effect of SGE from *I. ricinus* ticks on the proliferation of the CB53...
strain of *B. burgdorferi* s.s. (České Budějovice) in the acute phase of mouse infection. In the Czech Republic, *I. ricinus* represents the natural vector for spirochetes of the strain CB53.

**MATERIALS AND METHODS**

**Mice.** SPF female C3H/HeN mice, 6 to 10 weeks old, were purchased from Charles River, Sulzfeld, Germany.

**Borrelia burgdorferi.** Two strains of *B. burgdorferi* s.s. were used: B31 (from dr. Lise Gern, University of Neuchâtel, Switzerland) and CB53, isolated from *I. ricinus* (Štěpánová-Tresová et al. 1999). Spirochetes were grown in Barbour-Stoerner Kelly-H (BSK-H) medium (SIGMA) supplemented with 6% rabbit serum at 34°C. Bacteria were counted by dark-field microscopy and used at low passage (five or fewer passages).

**Salivary gland extract (SGE).** Twenty *I. ricinus* females (colony at the Institute of Parasitology, Academy of Sciences of the Czech Republic, České Budějovice), negative in *B. burgdorferi* s.l. PCR screening, were fed on guinea pigs. After 5 days, engorged ticks were removed and the salivary glands were dissected and pooled. The 5-day period, for which the ticks had fed on guinea pigs, was chosen due to the best effect of SGE prepared from such ticks on the saliva-activated transmission of TBE virus (Labuda et al. 1993). After washing in phosphate-buffered saline (PBS), the salivary glands were homogenized in 1 ml PBS by sonication and centrifuged at 10,000 g for 10 min. The protein concentration of SGE was determined and aliquots of the SGE were stored at −80°C. Before use, SGE activity was tested on mouse splenocytes stimulated with lipopolysaccharide. The active SGE inhibits production of INF-γ by these cells (Kopecky et al. 1999).

**Experimental infection of mice.** A group of 9 mice (BS) was intradermally injected in the ventral thorax area with 1.5 × 10⁸ spirochetes in 100 µl of BSK-H medium mixed with 40 µg of SGE in 50 µl PBS per mouse. Forty µg of protein can be obtained in SGE from one *I. ricinus* female. A second group of 12 mice (B) was injected with 1.5 × 10⁹ spirochetes in 100 µl of BSK-H medium mixed with 50 µl of PBS only. The control group of 4 mice (C) was inoculated with the same volume of BSK-H medium mixed with PBS. One hour after injection, samples of skin (3 cm²) were taken from the injection site of 3 mice from group B. The skin samples were used to estimate the actual number of inoculated spirochetes.

**Tissue samples.** On days 1, 3 and 7 post infection (p.i.), 3 mice of group B, 3 mice of group BS and 1 mouse from group C were used for sample collection. At each time point, samples of the following tissues were taken: 1) blood from the orbital sinus, 2) skin from the injection site, and 3) two pairs of lymph nodes: lymphonodus axillaris proprius and lymphonodus axillaris accessorius. Two hundred µl of blood were taken into microtube with 20 µl of 2% (w/v) EDTA. Skin was weighted after scraping off the subcutaneous fat and put in PBS. Each pair of lymph nodes was placed in PBS.

**Extraction of DNA.** DNA was extracted using a commercial Jetquick Tissue DNA Spin Kit (GENOMED, Germany). DNA preparation from skin and lymph nodes was performed according to the manufacturer’s instructions. For DNA extraction from the blood the supplier’s protocol was optimized as follows. One hundred µl of anticoagulated blood was mixed with 20 µl of proteinase K (20 mg/ml; GENOMED). Volume was adjusted to 220 µl with PBS and 200 µl of T2 buffer were added. Samples were vortexed and incubated at 70°C for 10 min. The additional steps correspond to Part B of Jetquick Spin Column Procedure provided by the producer.

**Real-time PCR.** For the q-PCR assay, primer pairs and probe for the flagellin gene sequence were used as described by Zeidner et al. (2001). This sequence was later identified as the gene for flagellar filament cap protein FlID (Ge et al. 1997, GenBank accession no. U66699.1), which is localized upstream of the flagellin gene sequence. The dual-labeled probe sequence was 5′-FAM-TGC TAA AAT GCT AGG AGA TTG TCT GTC GCC-TAMRA-3′, where FAM refers to the reporter dye and TAMRA to the quencher dye. The reverse primer sequence was identical as described by Zeidner et al. (2001): 5′-ACT CCT CGG GAA GCC ACA A-3′. Amplification using the forward primer described previously (Zeidner et al. 2001) produced irrelevant quantification results, which did not correspond to standard sample concentrations (data not shown). Therefore we designed a new forward primer for q-PCR: 5′-CTT CGT GTG GAC TTA GAT CTT CTA GAT-3′. The 20 µl q-PCR mixture consisted of 500 nM dual-labeled probe (Generi Biotech), 2 µM each primer (Generi Biotech), 2 units of Thermo Start Taq DNA polymerase (AB Gene), 200 µM dNTP each (TaKaRa), 2 µl of 10 × buffer (AB Gene), 40 ng of extracted DNA; 4.5 mM MgCl₂ and 10 µl of H₂O. Following an incubation step at 50°C for 2 min and 10 min of incubation at 95°C, the samples were submitted to amplification at 95°C for 15 s, at 60°C for 1 min, 50 cycles in a Rotor Gene 3000 (Corbett Research). Each specimen was tested in triplicate. Negative controls containing all PCR reaction components with the exception of template DNA (denoted No template control; NT) were included.

**Borrelia burgdorferi quantification standard.** We tested real-time PCR with two different *B. burgdorferi* strains, B31 and CB53, to standardise real-time PCR method using those spirochete strains. Three types of standard were used for q-PCR assay. The first type was a plasmid containing the partial sequence of flagellar filament cap protein of *B. burgdorferi* s.s. CB53. For cloning of a partial sequence, a new primer pair was designed: forward primer 5′-GGT GCT ACT ATG GGC-3′ and reverse primer 5′-CTG ACT GCT GCT GCT GCT GCT-3′. Following the transformation assay, pCR 2.1-TOPO vector and TOPO TA Cloning Kit (Invitrogen) were used. Ten-fold serially diluted aliquots of both standards mentioned above were prepared. The individual aliquots were used only twice for generation of the standard curve. The third type of standard was the DNA extracted from mouse tissues (skin, blood, lymph nodes) with addition of various concentrations of spirochetes. These standards served as the control of extraction and to test for the potential presence of inhibitors. For converting the concentration values to gene copy number, i.e. number of spirochetes, the following assumption was made: 1 fg of genomic DNA of *B. burgdorferi* corresponds to one spirochete (Johnson et al. 1992).
**Statistical analysis.** Data are presented as number of spirochetes per mg of skin, ml of blood and per one lymph node. Numbers of *Borrelia* spirochetes are presented as the mean ± SEM. The significance of differences in the mean numbers of spirochetes was evaluated by the Student’s *t*-test (*p* ≤ 0.05).

**RESULTS**

The q-PCR assay was used for examination of *Borrelia burgdorferi* infection rate in mouse tissues. As noted in Fig. 1, the amplification plot, in terms of fluorescence intensity versus cycle number, indicates an amplification of serial dilution of *B. burgdorferi* genomic DNA from 10 to 100,000 copies of *FliD* gene (i.e. number of spirochetes). The correlation coefficient was routinely between 0.97 and 0.99 (data not shown) and the detection limit of *B. burgdorferi* DNA was 10 spirochetes per sample. The sensitivity of the real-time PCR was identical when *B. burgdorferi* sensu stricto B31 and CB53 were compared (data not shown).

Using q-PCR, the effect of *I. ricinus* SGE on the infection of C3H mice with *B. burgdorferi* s.s. was tested. Intradermal injection of spirochetes mixed with SGE was compared with injection of spirochetes alone. Analysis of the skin samples one hour p.i. confirmed inoculation of 1.5 × 10^3^ spirochetes/mouse. The enhancing effect of SGE on the proliferation of spirochetes in vivo was detected on day 1 p.i. in the skin and blood (Fig. 2). The average number of spirochetes detected in the skin samples of the BS group (9.6 × 10^6^ spirochetes/mg) was 4.2-fold higher, but not significantly different from those of group B (2.3 × 10^6^ spirochetes/mg). On the other hand, spirochetes in blood samples of the BS group (8.6 × 10^7^ spirochetes/ml) significantly outnumbered (more than 10-fold) those in blood samples of group B (8.5 × 10^7^ spirochetes/ml). At later time points (3 and 7 days p.i.) the differences between both experimental groups were not significant (Figs. 3, 4).

The spirochete distribution in various tissues during the acute phase of infection was similar in both experimental groups. A significant increase in spirochete numbers in the skin samples, resulting from their proliferation, was detected on day 7 p.i. in group B (9.4 × 10^7^ spirochetes/mg) and group BS (7.2 × 10^5^ spirochetes/mg) compared to days 1 and 3 p.i. The decrease in spirochete load in the skin samples (1.1 × 10^5^ spirochetes/mg in group B and 4.4 × 10^5^ spirochetes/mg in group BS) 3 days p.i. presumably reflected dissemination of *Borrelia* spirochetes through the blood into lymph nodes. Spirochetes were present in lymph nodes 3 days, and some were detected 7 days p.i. None of the differences between SGE-treated and untreated groups in spirochete numbers in lymph nodes was significant.

**DISCUSSION**

Tick saliva-activated transmission of various tick-borne pathogens has been demonstrated (Jones et al. 1989, Labuda et al. 1993, Kročková et al. 2003). In the case of *Borrelia* spirochetes, Pechová et al. (2002) reported indirect evidence for increased bacteraemia in mice injected with *B. afzelii* plus *Ixodes ricinus* SGE compared with SGE-untreated control mice. An eight-fold increase in the number of spirochetes was observed after six days of cultivation in BSK-H medium of blood samples taken on day 1 p.i. from both groups. These results are in good agreement with the results reported here that demonstrate the SAT effect of SGE as early as day 1 p.i., but not later. Another observation supporting the idea of an early effect of SGE on spirochete proliferation was reported in the paper by Pechová et al. (2002). When pathogen-free *I. ricinus* nymphs were placed on mice infected with spirochetes plus SGE, 57% of the nymphs became infected compared with no infected nymphs that had fed on mice inoculated with the spirochetes without SGE. This indicates that the
Fig. 2–4. Quantification of spirochetes CB53 in mouse tissues by q-PCR on day 1 p.i. (Fig. 2), day 3 p.i. (Fig. 3) and day 7 p.i. (Fig. 4). The number of spirochetes is presented as the mean ± SEM. LAP – lymphonodus axillaris proprius, LAA – lymphonodus axillaris accessorius. *Statistically significant difference (p ≤ 0.05) in the number of spirochetes.

accelerating effect of SGE occurs during the feeding period, i.e. 3–4 days p.i. The explanation of the very early effect of SGE on the proliferation of spirochetes can reside in its local, short-lasting effect on the skin immune system. At later time points, when the pathogen can be demonstrated in draining lymph nodes, the number of spirochetes was comparable in all examined organs.

In contrast, Zeidner et al. (2002) also demonstrated an enhancing effect of *Ixodes scapularis* SGE on *B. burgdorferi* spirochete load in target organs, and a similar effect of *I. ricinus* SGE on *Borrelia lusitaniae*. However, in this case spirochete numbers were estimated as late as 8 weeks p.i. using real-time PCR.

The enhancing effect of tick saliva on *B. burgdorferi* transmission and proliferation in the host apparently has an immunological basis. The inhibitory effect of tick saliva on early nonspecific immune defences may be responsible for this phenomenon. For example, an inhibitory activity of *I. ricinus* SGE on complement-mediated killing of *B. burgdorferi* spirochetes, as well as inhibition of phagocytosis and killing of spirochetes by macrophages, have been demonstrated (Kuthejlová et al. 2001; Kýčková, Inst. Parasitol. AS CR, České Budějovice – pers. comm.), as has an anti-inflammatory influence of *I. ricinus* SGE (Severinová et al. 2005). These effects may be connected with anaphylatoxin inactivating activity (Ribeiro and Spielman 1986), histamine-binding activity (Paesen et al. 1999), and/or pro-inflammatory cytokine-inhibiting activity (Rama-chandra and Wikel 1992, Pechová et al. 2004). A direct accelerating effect of SGE on the growth of *Borrelia* spirochetes in BSK-H medium was also observed (Rudolf and Hubálek 2003).

The first identification of a SAT factor for *Borrelia* spirochetes was reported by Ramamoorthi et al. (2005). They showed that the previously described immunomodulatory protein Salp 15 from the saliva of *I. scapularis* tick (Anguita et al. 2002) binds specifically to the spirochete outer surface protein C, protects spirochetes from antibody-mediated killing and enhances the *B. burgdorferi* load in both naive and immune hosts. However, it is not clear whether a similar protein exists in the saliva of the European Lyme disease vector, *I. ricinus*.

It is obvious that injection of *Borrelia* spirochetes mixed with tick SGE does not reflect the *in vivo* situation, when the spirochetes are inoculated by the tick in several small doses mixed with tick saliva for a relatively long period (Sauer et al. 2000). One-shot inoculation of SGE apparently leads to a transient effect on the host immune system due to inactivation or degradation of immunomodulatory molecules mediated by SGE. However, repeated injection of SGE into the site of infection would not mimic the natural infection via tick feeding anyway. Consequently, the most marked effect on spirochete multiplication is observed shortly after inoculation of the pathogen plus SGE.
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REFERENCES


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