Genomic Confirmation of *Borrelia garinii*, United States

Appendix

Rodents, Locations, and Borrelia Cultures

Borrelia cultures analyzed in this report were a part of collection of southeastern United States isolates. Two samples under investigation, SCCH-7 and SCGT-19, were isolated from the ear clip tissues of cotton mice (*Peromyscus gossypinus*) and eastern woodrat (*Neotoma floridana*), respectively. The ear clips represented a small triangular pieces of tissue from the very peripheral tip of the external pinna of each animal. Before the use of ear clip for culture seeding, the ear clips were cleaned with 95% ethanol, and tissues were sliced into smaller pieces, than washed once in 95% ethanol, followed by a rinse in a 1:1 mixture of 10% Clorox and 95% ethanol (*1,2*). *P. gossypinus* was trapped in Mt. Pleasant, Charleston County, South Carolina (USA) in 1995 and *N. floridana* was trapped in Georgetown County, South Carolina (USA) 1 year later. *Borrelia* isolates from the ear clip tissues were cultured in Barbour-Stoenner-Kelly (BSK) H medium that contained 0.15% agarose (Seakem; FMC Bioproducts, Rockland, ME), antimicrobial drugs (rifampin and phosphomycin), and fungicide (amphotericin B). The cultures were incubated in 5% CO₂ at 33°C–34°C. When the cultures reached a cell density of 2×10^6 spirochetes/mL, they were stored at $-80^{\circ}C$ (3).

Co-infection

The total DNA of all samples was purified with a DNeasy Blood and Tissue kit (QIAGEN) according to the manufacturer's recommendations. A *MasterTaq* Kit (Eppendorf, Germany) containing a special $5 \times$ TaqMaster PCR enhancer was used for the amplification of spirochete sequences. All PCRs were set up in a dedicated area, and precautions were taken to limit contamination (supplies, equipment and personal safety items, and pre- and post-amplification activities). In all steps of the analysis of reported samples, the DNA of *B. burgdorferi* sensu stricto strain B31 was used as a positive control. Initial PCR analysis of isolates from sample depository included amplification of the 5S–23S intergenic region (4), followed by sequencing. The cultures in which the presence of more than one spirochete species

was confirmed (overlapping chromatograms) were further analyzed by PCR amplification with species-specific ospA-targeting primers (5) that enabled detection of spirochetes from B. burgdorferi sensu lato complex (SL primers) and enabled differentiation of B. burgdorferi sensu stricto (GI primers, PCR product 543 bp), B. garinii (GII primers, PCR product 344 bp) and B. afzelii (GIII primers, PCR product 189 bp). PCR products of the expected size (344 bp) were cut out of the gel, purified with a Gel Extraction kit from QIAGEN, and sequenced in both directions with the same primer set as used for amplification. Further control of cultures that produced the GII amplicon (B. garinii) included amplification of the partial *flagellin* gene from the total DNA purified from co-infected culture, cloning of total PCR products into the pCR4-TOPO cloning vector, transformation of Escherichia coli, plating of recombinants into LB agar plates, picking of individual clones, and growth in 100 microliters of LB/ampicillin medium. Ninety-six well plates with recombinants were submitted for sequencing to the University of Washington (Seattle). Sequencing was conducted in both directions. The obtained sequences were used for similarity searching in GenBank. Co-infected cultures in which the presence of *B. garinii* was confirmed through the 2 above steps proceeded to separation of spirochete species by cultivation on solid BSK medium.

Nucleotide Sequence Accession Numbers

Sequences determined in this study have been deposited into GenBank with the following accession numbers for SCCH-7/SCGT-19, respectively: *flagellin* EU220774/EU220773; 5S–23S IGR–KP795350/KP795363; 16S-23S ITR–KP795352/KT285872; *ospA* KP795349/KP795362; *ospC* KP795351/KT285871 and *p66* KP795348/KP795361. The accession numbers for the eight housekeeping genes for SCCH-7/SCGT-19 are *clpA*, KP795360/KT285880; *clpX*, KP795358/KT285878; *nifS*, KP795357/KT285877; *pepX*, KP795359/KT285879; *pyrG*, KP795356/KT285876; *recG*, KP795355/KT285875; *rplB*, KP795353/KT285873; and *uvrA*, KP795354/KT285874. The genome assembly of SCCH-7 has been deposited in GenBank under BioProject PRJNA431102 with the BioSample accession SAMN26226110.

Unique loci detected in both South Carolina isolates were submitted to PubMLST database (https://pubmlst.org/) as well and were assigned unique alleles numbers as follows: isolate SCCH-7 *clpX* gene-allele number 272; *uvrA* gene- allele number 278. Isolate SCGT-19 *clpA* gene- allele number 311, *clpX* gene- allele number 273. Unique ST numbers in pubMLST database for SCCH-7/SCGT-19 are 1049 and 1050, respectively.

Whole-Genome Sequencing and Genome Assembly

Genome sequencing was performed by using the Pacific Biosciences Sequel II system. Total genomic DNA was isolated from SCCH-7 cells by using DNeasy Blood and Tissue Kit (QIAGEN) according to the manufacturer's instructions. The gDNA sample contained relatively short fragments mostly less than 5 kb, so no shearing was performed. A random library was prepared by using the PacBio SMRTbell express template kit 2.0 according to the manufacturer's instructions. Sequencing was performed by using one Sequel II cell which generated 178 Gb total sequence and 1.37M High Fidelity (HiFi) reads totaling 2.4Gb with mean length of 1757bp and mean quality of QV60. To facilitate assembly, the subset of HiFi reads longer than 5,000 bp was generated, which yielded 210-Mb sequence in 32.6 k reads with mean quality of QV42.

Genome assembly was performed with the Genome Assembly tool in PacBio SMRTLink 10.2 by using 150 Mb of the HiFi reads greater than 5kb (100× down sample with 1.5-Mb expected genome size). The assembly was polished by the PacBio Arrow algorithm, and the telomere ends were examined manually by comparison to multiple individual HiFi reads.

Phylogenetic Analysis

The sequences of 8 housekeeping loci (*clpA*, *clpX*, *nifS*, *pepX*, *pyrG*, *recG*, *rplB*, and *uvrA*) of *Borrelia* isolates were obtained from the PubMLST database (https://pubmlst.org/), and sequences of several additional relevant isolates were added from our collection. The resulting dataset was aligned with the MUSCLE aligner (*6*) implemented in SeaView 4 (*7*). To ensure the frame integrity of codon information, we translated the sequences into amino acids first, then aligned and back-translated them into nucleotides. Phylogenetic analysis with the maximum likelihood method was performed in RAxML (*8*) under the General time-reversible substitutional matrix, with nucleotide frequencies estimated from the dataset and four gamma-corrected rate site classes (GTR + G4 + F). The best topology and the branching support values were calculated by using the rapid bootstrap inferences from 1,000 replicates followed by a thorough ML search (fa parameter) in RAxML.

Phylogeographic analysis of diffusion on discrete space as implemented in BEAST (9,10) was performed under the GTR + G4 matrix, constant-size coalescent tree prior, and symmetric substitution model with BSSVS enforced. To make the analysis more feasible, we simplified the location coding into following geographic categories: Europe, Asia, United States and Canada.

To obtain sufficient effective sample sizes for the estimated parameters, we ran Monte Carlo Markov Chains (MCMC) for 100 million generations subsampling every 10,000 trees. The MCMC convergence was then inspected, and burnin value was selected in Tracer (11). The final topology, including the reconstruction of distribution of *B. garinii* strains was summarized in TreeAnnotator (9), and the topology was visualized in FigTree. The approximately unbiased test (12) as implemented in CONSEL (13) was used to test alternative phylogenetic topologies.

Prevalence of B. garinii in Samples from the Southeastern United States

During 2005–2017, we have studied a group of spirochete cultures that was a part of southeastern depository of *Borrelia burgdorferi* sensu lato strains in former James H. Oliver Institute of Arthropodology and Parasitology (Statesboro, Georgia, USA). Spirochetes from analyzed group were isolated from environmental samples, collected in 9 different localities of the southern region of the United States. Samples from the studied group were collected in South Carolina, Georgia, and Florida during 1993–1997. Spirochetes were cultured from 3 rodent species, cotton mouse *Peromyscus gossypinus*, eastern woodrat *Neotoma floridana*, and cotton rat *Sigmodon hispidus* (78 cultures), 3 tick species, *Ixodes affinis, I. minor*, and *I. scapularis* (40 cultures), and 9 bird species, Carolina wren (*Thryothorus ludovicianus*), Downy woodpecker (*Picoides pubescens*), White eyed vireo (*Vireo griseus*), Swainson's thrush (*Catharus ustulatus*), American redstart (*Setophaga ruticilla*), Northern waterthrush (*Parkesia noveboracensis*), Pine warbler (*Setophaga pinus*), Northern cardinal (*Cardinalis cardinalis*), and Eastern towhee (*Pipilo erythrophthalmus*) (13 cultures) (Appendix Table 4).

Recultivation of spirochetes, DNA purification and PCR amplification of selected genomic loci were conducted as described above (3,14). The results of analyses revealed the presence of multiple spirochete species in the samples of the studied group: 16 strains represented new *Borrelia* species that were described as *Borrelia* carolinensis (3,15); another 7 isolates, that represented unknown spirochete species, were described as *Borrelia* americana (16,17); 53 isolates were identified as *Borrelia* burgdorferi sensu stricto (14,18); 36 cultures represented rather diverse group of *Borrelia* bissettiae. The remaining 19 isolates represented cultures with more than one spirochete species present (co-infected). All 13 cultures isolated from birds contained >1 spirochete species.

Of 131 cultures analyzed, *B. garinii* was detected by PCR amplification targeting the *ospA* gene with species-specific primers (5) in 5 co-infected cultures. The detected prevalence of *B. garinii* (3.8%) is based on results of the studied group only (not the whole collection of depository strains). Isolation of monoclonal populations of *B. garinii* was successful only for SCCH-7 and SCGT-19 isolates (Appendix Table 5).

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Appendix Table 1. Borrelia strains included into phylogenetic analysis based on 8 housekeeping genes*

Strain name	Sample location	Species
164/11g	Serbia	Borrelia garinii
20047	France	Borrelia garinii
IPT140	France	Borrelia garinii
IPT130	France	Borrelia garinii
IPT139	France	Borrelia garinii
IPT165	France	Borrelia garinii
MDH1	China	Borrelia garinii
VH4	China	Borrelia garinii
VL2	China	Borrelia garinii
70576B	UK	Borrelia garinii
6910BT	UK	Borrelia garinii
61030BT	UK	Borrelia garinii
70531B	UK	Borrelia garinii
		Borrelia garinii
	•	Borrelia garinii
		Borrelia garinii
	•	Borrelia garinii
		Borrelia garinii
		Borrelia garinii
•		Borrelia garinii
Np189		Borrelia garinii
J-37 (human)		Borrelia garinii
05105781	Italy	Borrelia garinii
05106051	Italy	Borrelia garinii
02203541	Italy	Borrelia garinii
rebun_20	Japan	Borrelia garinii
konnai 1	Japan	Borrelia garinii
konnai 15 clone5		Borrelia garinii
EU9-22	The Netherlands	Borrelia garinii
Ekb701-11	Russia	Borrelia garinii
		Borrelia garinii
akt7		Borrelia garinii
	-	Borrelia garinii
		Borrelia garinii
	•	Borrelia garinii
akt43	Norway	Borrelia garinii
anito	norway	Donena yanını
akt/6	Norway	Borrelia garinii
akt46 akt52	Norway Norway	Borrelia garinii Borrelia garinii
	164/11g 20047 IPT140 IPT130 IPT139 IPT165 MDH1 VH4 VL2 70576B 6910BT 61030BT 70531B 62303L 61209L 75803L 80201G 82805G TS1 DR46 LV5M DV12 b34 DV46 DV13 LA9 LA34 MNG61 HKIP2 HP1 HP3 NP4 NP76 NP81 NT25 NT31 Ip-90 Np189 J-37 (human) 05105781 05106051 02203541 rebun_20 konnai_1 konnai_15_clone5 EU9-22 Ekb701-111 Ekb712-111 akt7 akt10 akt11 akt19	164/11g Serbia 20047 France IPT140 France IPT130 France IPT139 France IPT165 France MDH1 China VL2 China 70576B UK 6910BT UK 61030BT UK 62303L Latvia 61209L Latvia 61209L Latvia 61209L Latvia 80201G Germany 82805G Japan UK DV12 UK DV13 UK LA34 UK MNG61 Mongolia HP1 Japan NP4 Japan NP56 Japan </td

pubMLST ID	Strain name	Sample location	Species
1707	PKuf (human)	Germany	Borrelia garinii
1743	Ekb151-2012	Russia	Borrelia garinii
1758	Ekb783-2013	Russia	Borrelia garinii
1770	Nsk1400-2013	Russia	Borrelia garinii
1790	Tmsk1125-2013	Russia	Borrelia garinii
1791	Tmsk1128-2013	Russia	Borrelia garinii
1792	Tmsk1130-2013	Russia	Borrelia garinii
1793	Tmsk1187-2013	Russia	Borrelia garinii
1794	Tmsk1188-2013	Russia	Borrelia garinii
1795	Tmsk1189-2013	Russia	Borrelia garinii
1798	Tmsk1193-2013	Russia	Borrelia garinii
1800 1808	Tmsk1218-2013 Tmsk2148-2014	Russia Russia	Borrelia garinii Borrelia garinii
1824	PCoo (human)	Germany	Borrelia garinii
1826	PMag (human)	Germany	Borrelia garinii
1827	PMek (human)	Germany	Borrelia garinii
1829	PUI (human)	Germany	Borrelia garinii
1860	Tom5202	Russia	Borrelia garinii
1863	Tom2903	Russia	Borrelia garinii
1864	Tom1805	Russia	Borrelia garinii
1866	Tom8705	Russia	Borrelia garinii
1901	NL11-021	Canada	Borrelia garinii
1904	NL12-114	Canada	Borrelia garinii
1910	NL12-334C	Canada	Borrelia garinii
1911	NL12-340F	Canada	Borrelia garinii
1913	NL13-029	Canada	Borrelia garinii
1918	NL13-245	Canada	Borrelia garinii
1919	NL13-440	Canada	Borrelia garinii
1921	NL13-534	Canada	Borrelia garinii
1923	NL14-1000	Canada	Borrelia garinii
1962 1981	PBe (human)	Germany	Borrelia garinii Borrelia garinii
1992	PNel (human)	Germany Germany	Borrelia garinii Borrelia garinii
2034	PStg (human) HN13	South Korea	Borrelia garinii Borrelia garinii
2452	EuTu104	Czech Republic	Borrelia garinii
2454	EuTu185	Czech Republic	Borrelia garinii
2457	EuTu251	Slovenia	Borrelia garinii
2459	EuTu347	Sweden	Borrelia garinii
2460	EuTu352	Sweden	Borrelia garinii
2463	EuTu451	The Netherlands	Borrelia garinii
2467	EuTu490	Finland	Borrelia garinii
2471	EuTu488	Finland	Borrelia garinii
2473	EuTu456	Finland	Borrelia garinii
2475	EuTu519	Estonia	Borrelia garinii
2479	EuTu476	Finland	Borrelia garinii
2515	DNQ100	Slovakia	Borrelia garinii
2523	DNQ9	Slovakia	Borrelia garinii
2583 2669	451_UA 132DIVN1	Ukraine Slovakia	Borrelia garinii Borrelia garinii
2698	NE5245	Switzerland	Borrelia garinii Borrelia garinii
2090	17-58N4 wgs	Norway	Borrelia garinii
2875	85DIVN13	Slovakia	Borrelia garinii
2906	9-22-27	Latvia	Borrelia garinii
3068	8/1/2029	Latvia	Borrelia garinii
3070	1-31LT	Latvia	Borrelia garinii
3083	2-27LT	Latvia	Borrelia garinii
3107	3-17-10	Latvia	Borrelia garinii
3113	10-26-29	Latvia	Borrelia garinii
3140	NE4906	Switzerland	Borrelia garinii
3177	MaN25417/86	Slovakia	Borrelia garinii
3178	MaN25417/149	Slovakia	Borrelia garinii
3179	MaN111017/219	Slovakia	Borrelia garinii
3181	KF3517/45	Slovakia	Borrelia garinii
3182	KF3517/47	Slovakia	Borrelia garinii
3184	ZSF25417/69	Slovakia The Netherlands	Borrelia garinii
NA	NLD124_2011	The Netherlands	Borrelia garinii
NA	NLD128_2011	The Netherlands	Borrelia garinii Borrelia garinii
NA NA	NLD132_2010 NLD135 2011	The Netherlands The Netherlands	Borrelia garinii Borrelia garinii
	NED 135_2011		Donella yallilli

pubMLST ID	Strain name	Sample location	Species
NA	NLD145 2010	The Netherlands	Borrelia garinii
NA	NLD146_2010	The Netherlands	Borrelia garinii
NA	NLD149_2010	The Netherlands	Borrelia garinii
NA	NLD212_2009	The Netherlands	Borrelia garinii
1049	USA233_1995	USA	Borrelia garinii
1050	USA234_1996	USA	Borrelia garinii
NA	ISL367_2010	Iceland	Borrelia garinii
NA	ISL369_2010	Iceland	Borrelia garinii
NA	ISL370_2010	Iceland	Borrelia garinii
NA	ISL371_2010	Iceland	Borrelia garinii
NA	ISL372_2010	Iceland	Borrelia garinii
NA	ISL373_2010	Iceland	Borrelia garinii
NA	ISL375_2010	Iceland	Borrelia garinii
NA	ISL376_2010	Iceland	Borrelia garinii
198	NMK6	China	Borrelia bavariensis
202	JW3	China	Borrelia bavariensis
203	VH1	China	Borrelia bavariensis
204	VH2	China	Borrelia bavariensis
205	VH3	China	Borrelia bavariensis
209	VH19	China	Borrelia bavariensis
214	JLHCH	China	Borrelia bavariensis
216	HQ	China	Borrelia bavariensis
1060	MNG14	Mongolia	Borrelia bavariensis
1061	MNG24	Mongolia	Borrelia bavariensis
1076	HkIP1	Japan	Borrelia bavariensis
1082	N346	Japan	Borrelia bavariensis
1087	NT24	Japan	Borrelia bavariensis
1092	Mp7	Russia	Borrelia bavariensis
1095	HkCR3	Japan	Borrelia bavariensis
1104	FsAE1	Japan	Borrelia bavariensis
1112	ChYAE2	China	Borrelia bavariensis
1118	J-15	Japan	Borrelia bavariensis
1119 1122	J-16 J-20T	Japan	Borrelia bavariensis Borrelia bavariensis
1122	J-201 HH1	Japan	Borrelia bavariensis
1313	enkichi 32	Japan China	Borrelia bavariensis
1333	konnai 14	Japan	Borrelia bavariensis
1340	takamine As 5	Japan	Borrelia bavariensis
1431	Prm7564-11	Russia	Borrelia bavariensis
1440	Ekb166-10	Russia	Borrelia bavariensis
1440	Alt763-11	Russia	Borrelia bavariensis
1446	Arh976-12	Russia	Borrelia bavariensis
1459	PScf	Germany	Borrelia bavariensis
1735	Arh913-2012	Russia	Borrelia bavariensis
1741	Ekb1421-2014	Russia	Borrelia bavariensis
1745	Ekb169-2012	Russia	Borrelia bavariensis
1802	Tmsk1253-2013	Russia	Borrelia bavariensis
1804	Tmsk1613-2014	Russia	Borrelia bavariensis
1839	Mng4702	Mongolia	Borrelia bavariensis
1843	Tom1003	Russia	Borrelia bavariensis
1853	Tom4606	Russia	Borrelia bavariensis
1859	Tom5007	Russia	Borrelia bavariensis
1902	NL11-061	Canada	Borrelia bavariensis
2488	Om16-103-lapr	Russia	Borrelia sp.
1283	PoTiBtur10	Portugal	Borrelia turdi
1285	PoTiBtur12	Portugal	Borrelia turdi
1458	Ya501	Japan	Borrelia turdi
2075	T2084	Portugal	Borrelia turdi
2076	TPT2017	Portugal	Borrelia turdi
*NA, not available.			

*NA, not available.

Appendix Table 2. Results of approximately-unbiased (au) topology test comparing the original most-likely topology as seen on Figure 1 (row 1), with the alternative topologies with enforced Canadian-US *B. garinii* monophyly (rows 2–11). The p-AU column denotes the statistical significance (p values) of the test for individual topology

Tree	logL	deltaL	p-AU
1	-22917.7	0	0.995
2	-23054.4	136.65	0.0002
3	-23090.9	173.17	0.000748
4	-23143.6	225.85	6.44E-05
5	-23201.7	283.95	1.48E-36
6	-23030.6	112.85	0.0192
7	-23181.5	263.79	3.62E-07
8	-23159.1	241.34	0.000139
9	-23099.5	181.79	0.00125
10	-23051.1	133.35	0.00791
11	-23055.2	137.41	0.00345

Appendix Table 3. Distribution of single nucleotide variants (SNVs) among the housekeeping genes

MLS	ST locus/posit	tion*			cl	oA (579	nt)				cl	oX (624	nt)			nifS	(564nt)				pyrG	(603 nt	:)		recG	i (651 i	nt)			rpIB	(624 n	t)			uvrA	(570 ו	nt)		
												382-																											
Strain	Origin	ID	57	201	207	342	355	413	444	192	342	384	548	558	13	336	425	447	78	405	468	477	524	585	249	303	351	474	546	15	69 4	420 ·	468	585	267	330	370	477	[′] 49
20047	France	CP028861	Α	Α	С	Т	Α	С	С	С	Α		Α	Т	A	Α	А	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	Ç
20047	France	CP018744	Α	Α	С	Т	Α	С	С	С	Α		Α	Т	A	Α	Α	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	C C
20047	France	153	Α	Α	С	Т	A	С	С	С	Α		Α	Т	A	Α	Α	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	Ç
PCoo	Germany	1824	Α	Α	С	Т	Α	С	С	С	Α		Α	Т	A	Α	Α	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	C C
DNQ100	Slovakia	2515	Α	Α	С	Т	Α	С	С	С	Α		Α	Т	A	Α	Α	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	ç
1-31LT	Latvia	3070	Α	Α	С	Т	Α	С	С	С	Α		Α	Т	A	Α	Α	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	C C
SCCH-7**	USA	1049	Α	Α	С	Т	A	С	С	С	Α	AGA	Α	Т	A	Α	Α	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	Ç
SCGT-19	USA	1050	Α	G	С	С	G	С	С	С	Α	AGA	G	С	A	Α	Α	G	Т	С	С	Т	G	A	Т	Т	Т	G	Т	A	A	G	А	A	Т	Т	G	Т	Ç
HP1	Japan	1078	Α	Α	С	Т	A	С	С	С	Α		Α	Т	A	Α	Α	G	С	Т	Т	С	G	G	Т	Т	Т	G	Т	A	A	G	А	A	Т	С	Α	С	7
NT31	Japan	1089	Α	Α	С	Т	Α	С	С	С	Α	AGA	G	С	С	Α	Α	G	Т	Т	С	Т	G	G	С	С	С	Α	С	G	G	A	G	G	Т	С	Α	С	7
rubun20	Japan	1323	Α	Α	С	Т	A	С	С	A	G		G	С	С	Α	Α	G	С	Т	Т	С	G	G	С	С	С	Α	С	G	G	A	G	G	Т	С	Α	С	7
NL11-021	Canada	1901	G	G	Т	С	G	С	Т	A	G	AGA	G	С	С	G	G	A	С	Т	Т	С	Α	G	С	С	С	Α	С	A	G	A	G	G	Т	С	Α	Т	7
NL12-334C	Canada	1910	Α	G	С	С	G	Т	Т	A	G	AGA	G	С	С	G	G	A	С	Т	Т	С	Α	G	С	С	С	Α	С	G	G	A	G	G	С	С	Α	С	7
NL12-340F	Canada	1911	Α	G	С	С	G	Т	Т	A	G	AGA	G	С	С	G	G	A	С	Т	Т	С	Α	G	С	С	С	Α	С	G	G	A	G	G	С	С	Α	С	7
NL13-029	Canada	1913	Α	G	С	С	G	Т	Т	A	G	AGA	G	С	С	G	G	Α	С	Т	Т	С	Α	G	С	С	С	Α	С	G	G	A	G	G	С	С	Α	С	7
NL13-245	Canada	1918	G	G	Т	С	G	Т	Т	С	Α	AGA	G	С	С	А	А	G	С	Т	С	Т	G	G	С	С	С	Α	С	Α	G	A	G	A	С	С	Α	С	7
NL13-440	Canada	1919	G	G	Т	С	G	С	Т	A	G	AGA	G	С	С	G	G	A	С	Т	Т	С	A	G	С	С	С	Α	С	A	G	A	G	G	Т	С	A	Т	7
NL13-534	Canada	1921	G	G	Т	С	G	С	Т	С	Α	AGA	G	С	С	G	G	A	С	Т	Т	С	А	G	С	С	С	Α	С	A	G	A	G	G	Т	С	A	Т	7 🗾
NL14-1000	Canada	1923	G	G	Т	С	G	Т	Т	С	А		A	С	С	G	G	Α	С	Т	Т	С	А	G	С	С	С	Α	С	A	Α	Α	Α	Α	С	С	A	С	7

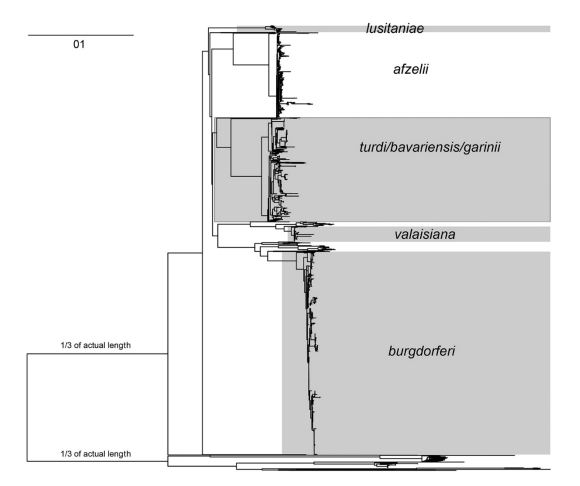
*Position of SNP conflicts is shown on respected housekeeping genes, which size was adjusted according to the MLST scheme (19). **Sequences of respective genes of strain SCCH-7 were extracted from genome sequences.

Appendix Table 4. Composition of the analyzed group of samples from strains depository, United States

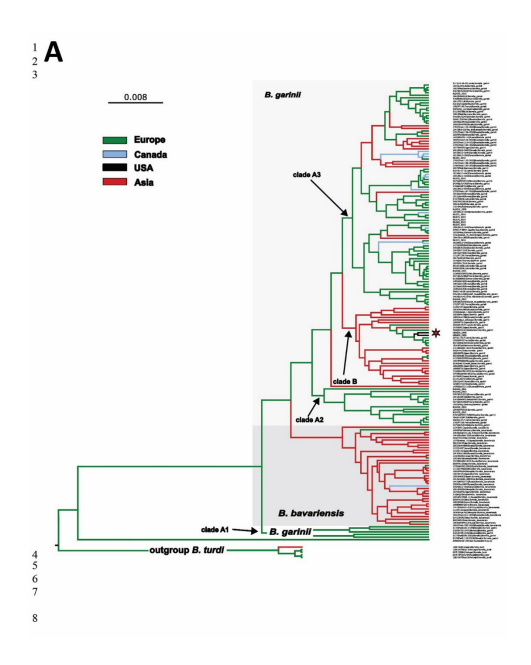
Location	Σ	P. gossypinus	N. floridana	S. hispidus	I. scapularis	I. affinis	I. minor	Birds
Georgia	21	9	4	2	0	6	0	0
Florida	3	1	0	2	0	0	0	0
South Carolina	94	32	20	8	4	18	12	13
Total	118	42	24	12	4	24	12	13
			Rodents 78			Ticks 40		

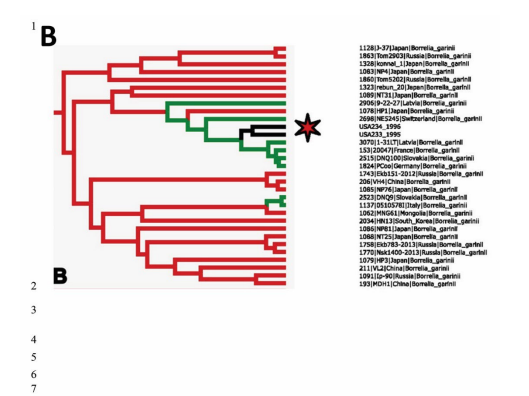
Appendix Table 5. Sample description and current state of *B. garinii*-positive cultures, detected in strains depository.

Sample	Collection date/site	Host	Source	Culture	Original status	Present status
SCCH-7	February 1995/SC USA	Cotton mouse	Ear clip	Cultured	Co-infection	Monoclonal
SCGT-19	January 1996/SC USA	Eastern woodrat	Ear clip	Cultured	Co-infection	Monoclonal
SCI-8	November 1997/SC USA	Downy woodpecker	Skin	Cultured	Co-infection	Co-infection
BUL-12	November 1997/GA USA	Cotton rat	Ear clip	Cultured	Co-infection	Co-infection
GAC-6	January 1998/GA USA	Cotton mouse	Ear clip	Cultured	Co-infection	Co-infection

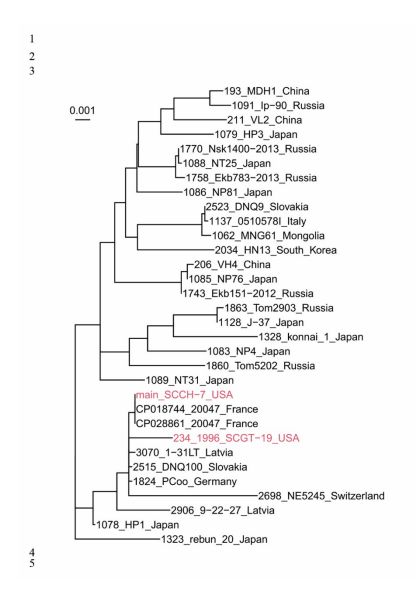


Appendix Figure 1. Unrooted maximum likelihood phylogeny of *Borrelia* based on analysis of 8 MLST genotyping loci (see above for details) under the GTR + G4 model implemented in RAxML 8. The final alignment comprises 2,977 taxa and 4,815- nt positions. Length of basal branches was reduced for formatting reasons.





Appendix Figure 2. A) Phylogeographic analysis of *B. garinii/B. bavariensis* rooted with *B. turdi* by using the diffusion on the discrete space algorithm implemented in BEAST under the GTR + G4 matrix, constant-size coalescent tree prior, and symmetric substitution model with BSSVS enforced. The isolates were clustered into 4 categories according to their geographic origin, coded in form of branch colors according the scheme in the upper right part of the tree on the topology. The position of two USA isolates is indicated by an asterisk. B) Ancestrally Asian clade. Maximum-likelihood phylogeny of *B. garinii* based on analysis of the partitioned dataset of 8 MLST genotyping loci (see Methods for details) under the GTR + G4 model (for each partition) in RAxML 8. Subset of results of diffusion on the discrete space showing the estimated geographic origin of the inner branches for the clade, on which USA-originated *B. garinii* were allocated. *B. garinii* type strain 20047 is indicated by the arrow. Full topology is shown in panel A.



Appendix Figure 3. Maximum-likelihood tree of 32 closely related *B. garinii* isolates based on sequences at the 8 housekeeping loci. The maximum-likelihood tree was inferred with IQTREE. All branches shown are supported by a bootstrap value \geq 80%. Two US isolates of *B. garinii* are highlighted in red. The SCCH-7 MLST sequence, grouped with the 2 previously sequenced genomes of strain 20047 (CP028861 and CP018744), was derived from the genome sequence.