

Re: Borrelia genetics

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Date: 02/27/05

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eugeneshapiroisapig wrote:

> *Hey guys, I could spend a lot of time at the library looking this up on my own, or I could save time and ask here. What I am interested in now are basic facts about the molecular biology of Borrelia species other than Borellia burgdorferi. By which I mean hermsii, etc., you know the stuff that causes relapsing fever and things like this. In particular, I am interested now in the genetic organization of these bugs. Burgdorferi has I believe 9 circular plasmids, 12 linear plasmids, and a linear chromosome. What about other borrelia? Also interested in worldwide distribution of hermsii. Any other information about comparative cell biology or evolution would also be appreciated.*

Thanks.

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Comparative analysis of the *Borrelia garinii* genome — Glöckner et al.
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Comparative analysis of the *Borrelia garinii* genome
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ABSTRACT

Three members of the genus *Borrelia* (*B.burgdorferi*, *B.garinii*, *B.afzelii*) cause tick-borne borreliosis. Depending on the *Borrelia* species involved, the borreliosis differs in its clinical symptoms. Comparative genomics opens up a way to elucidate the underlying differences in *Borrelia* species. We analysed a low redundancy whole-genome shotgun (WGS) assembly of a *B.garinii* strain isolated from a patient with neuroborreliosis in comparison to the *B.burgdorferi* genome. This analysis reveals that most of the chromosome is conserved (92.7% identity on DNA as well as on amino acid level) in the two species, and no chromosomal rearrangement or larger insertions/deletions could be observed. Furthermore, two collinear plasmids (*lp54* and *cp26*) seem to belong to the basic genome inventory of *Borrelia* species. These three collinear parts of the *Borrelia* genome encode 861 genes, which are orthologous in the two species examined. The majority of the genetic information of the other plasmids of *B.burgdorferii* is also present in *B.garinii* although orthology is not easy to define due to a high redundancy of the plasmid fraction. Yet, we did not find counterparts of the *B.burgdorferi* plasmids *lp36* and *lp38* or their respective gene repertoire in the *B.garinii* genome. Thus, phenotypic differences between the two species could be attributable to the presence or absence of these two plasmids as well as to the potentially positively selected genes.

INTRODUCTION

The genus *Borrelia* comprises 19 species of which 10 belong to the *Borrelia burgdorferi sensu lato* complex (1). Only three species of this complex cause the multisystem disorder Lyme borreliosis, *B.burgdorferi sensu stricto*, *B.garinii* and *B.afzelii*. In the United States, *B.burgdorferi sensu stricto* is the only causative agent for this disease (2), whereas in Europe *B.garinii* as well as *B.afzelii* are major contributors to the reported case numbers (3,4).

All these borreliae live in the gastro–intestinal tract of ticks (*Ixodes spec.*) and are able to infect multiple hosts via tick bite. The host range is thought to be defined by gene variations on a group of redundant plasmids (5–7). Furthermore, the disease patterns observed in humans are dependent on the particular *Borrelia* species involved. *B.garinii* is primarily associated with neuroborreliosis (3), *B.afzelii* with acrodermatitis chronica atrophicans (a chronic skin disease) (8), whereas *B.burgdorferi sensu stricto* was found to be prevalent in Lyme arthritis (9), which, however, was not confirmed by two other studies (10,11).

The nuclear genomes of all *B.burgdorferi sensu lato* species consist of one linear chromosome and varying amounts of several linear and circular plasmids (12). The chromosomes are highly similar (13), but plasmids show a wealth of diversity and can be lost during culture of the bacteria (14,15). The genomic sequence of the *B.burgdorferi* B31 chromosome was determined in 1997 (16). The sequence analysis of *B.burgdorferi sensu stricto* showed that the chromosome has a length of 0.94 Mb harbouring all genes for basic cellular functions. The plasmids often share sequence motifs and segments, and seem to contain a large number of fragmented genes, since several sequence motifs occur in predicted non–coding as well as coding regions. This fact and the lack of orthologs in other species made it difficult to define genes (17). Although the plasmid fraction of the genome seems to be responsible for host range selection and pathogenicity (6,18), large parts of it are likely to be dispensable for viability in culture and are highly variable, even in a single species (19).

The *Borrelia* species belong to the spirochetes, a group of bacteria that have long, helically coiled cells. To date, the genomes of the following spirochete species are known in addition to *Borrelia burgdorferi*: *Treponema denticola*, *Treponema pallidum* and *Leptospira interrogans* (16,20–22). Their genome sizes range from 1.6 to 4.3 Mb, with *B.burgdorferi* possessing the smallest genome. The *B.burgdorferi* sequence was recently used for microarray experiments to identify similar sequences in *B.hermsii* (23). This analysis revealed that at least 81% of the chromosome and 41% of the plasmid sequences of *B.burgdorferi* are present in this distantly related species. Yet, hybridization experiments can only detect highly similar sequences shared between organisms. On the other hand, a comparative analysis of sequences is able to detect not only low similarities that would not give hybridization signals, but provides also access to sequence differences of the organisms compared. Closely related species reveal species–specific differences and evolutionary selection pressures on genes. At the same time, a comparative sequence analysis provides the means for a better annotation as was shown with several *Saccharomyces* species (24). The comparison of more distantly related species helps to describe core sets of proteins within a specific evolutionary branch (25). The goal of the present comparative study was (i) to define the orthologous gene set, which presumably is the basic set of borreliosis causing *Borreliae*; (ii) to identify the stable and variable parts in

the genomes of *Borrelia*; (iii) to determine, which genome part is dispensable without losing pathogenicity; (iv) to improve the the *B.burgdorferi* annotation by the *B.garinii* orthology information. Furthermore, gene groups exposed to different evolutionary changes can be defined. Since positive selection is an indicator for environmental adaptations of the pathogen, genes evolving in such a manner are highly suspicious to be involved in pathogenicity. Many of the *B.burgdorferi* genes are so far described as only hypothetical. Thus, a comparative analysis would also help to define true orthologous pairs and exclude false positive predictions.

Therefore, we decided to sequence and analyse another species from the *B.burgdorferi* sensu lato complex, *B.garinii*. Previous studies had revealed that selected genes of both species are highly similar with *>90% sequence identity on DNA level. Moreover, there were hints that the chromosomes of the two species are collinear* (26). This relatively high similarity enabled us to apply a low redundant sequencing strategy. Thus, we were able to generate a complete analysis of the chromosome and two conserved plasmids of *B.garinii*. The diverse plasmid fraction of the genome could also be defined and analysed although no clear-cut assignment to *B.burgdorferi* plasmids was possible due to redundancies and rearrangements.

MATERIALS AND METHODS

The *B.garinii* strain PBi (OspA serotype 4), a cerebrospinal fluid (CSF) isolate from a German patient with neuroborreliosis, was used for sequence analysis (27). OspA-serotype 4 strains are enriched in CSF, but they have been isolated only exceptionally from ticks (27). A low passage of strain PBi (12th subculture) was cultured in MVP-medium as described (28). Passage 12 is still infectious for gerbils, infectivity was lost between passage 30 and 60. DNA was extracted using the Genomic DNA Bufferset (Quiagen GmbH, Hilden) and Genomic tip 500/6 and 100/6 (Quiagen, GmbH, Hilden). A genomic library with a target insert size of 1.5 kb using total DNA was constructed as described previously (29). *>>From this library, 5740 clones were sequenced from both ends resulting in an estimated coverage of the whole genome of three times. According to the similarity of the obtained sequences to B.burgdorferi counterparts they were binned into a plasmid and a chromosome group using BLAST* (30). The chromosomal sequences were assembled using the assembler GAP4 (31) utilizing the finished chromosome of *B.burgdorferi* (16,17) as a backbone. Using the backbone sequence as a ruler and orientation measure, we defined primer pairs for PCR reactions to close the gaps. For convenient primer design, we have written a Perl script for the automatic definition of primers in a Staden package project with the program 'primer3' (http://www-genome.wi.mit.edu/genome_software/other/primer3.html) as kernel (R.Lehmann, unpublished data). Nucleotide differences between the *B.garinii* and the *B.burgdorferi* chromosome were calculated automatically if the Phred score was 20 or better at the differing base. Bases below this score were inspected by eye to ensure proper

difference calculation.

Initial gene finding was performed using GeneMarkS (<http://opal.biology.gatech.edu/GeneMark>) (32). Orthology to *B.burgdorferi* counterparts was determined by aligning the best bidirectional hit (BBH) to each predicted protein. Only proteins located at the same position within the different genomes and with <10% length difference were considered as orthologs. Orphan genes of both organisms contained no functional domains according to an InterPro analysis (<http://www.ebi.ac.uk/interpro/>). Transmembrane domains were predicted using TMHMM (<http://www.cbs.dtu.dk/services/TMHMM/>). The GenBank gene descriptions of the *B.burgdorferi* genome (NC_001318 [GenBank], NC_000948 [GenBank] to NC_000957 [GenBank], NC_001849 [GenBank] to NC_001857 [GenBank], NC_001903 [GenBank], NC_001904 [GenBank], NC_004971 [GenBank]) were used for the annotation of the corresponding *B.garinii* coding sequences.

To perform an independent cross-check of the reliability of this annotation approach, all potential protein coding sequences (CDS; potential start codon to stop codon without length threshold) of *B.garinii* were used for BLASTP searches against the GenBank database. Whenever a gene is referred to as 'hypothetical', no match in any database could be found. A 'conserved hypothetical gene' is a gene, which can be detected with sufficient similarity ($p < 10^{-10}$) in other genera.

The alignment between the *B.garinii* and *B.burgdorferi* collinear chromosomes and plasmids was generated using the program stretcher, which is part of the EMBOSS package (<http://www.emboss.org>).

The *B.garinii* sequences were deposited in GenBank with the accession numbers CP000013 [GenBank], CP000014 [GenBank], CP000015 [GenBank] and AY722917 [GenBank] to AY722953 [GenBank]. The *B.garinii* genome data as well as the results of the comparative analysis of *B.garinii* with *B.burgdorferi* are also available from the Spirochetes Genome Browser at <http://sgb.imb-jena.de/>. The browser is based on our genome annotation and analysis system GenColors (manuscript in preparation).

RESULTS

The DNA for the libraries was obtained from the total DNA content of *B.garinii* strain PBi. Thus, besides the chromosome, the plasmids of this strain should also be represented at least in part in the shotgun data. All sequences from the whole-genome library were binned according to their similarities to the *B.burgdorferi* genome parts (i.e. chromosome and plasmids): 37.8% of all *B.garinii* reads were derived from plasmids (Table 1). This value is comparable to that of the plasmid fraction of *B.burgdorferi* (40%).

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Table 1. Comparison of the *B.garinii* low redundancy WGS assembly to the *B.burgdorferi* sequence

Altogether the assembly comprises 1.227 Mb of *B.garinii* sequence. Three contigs completely cover the counterparts of the corresponding *B.burgdorferi* chromosome and plasmids lp54 and cp26. Additional 37 contigs >2 kb amounting to 239 kb were obtained for the remaining plasmid fraction of *B.garinii* (Figure 1). Clear-cut and error-free assignment of these plasmid contigs to defined *B.burgdorferi* plasmids was not possible, further underlining the variability of the plasmid complement in *Borrelia* species. The whole chromosome and two plasmids (lp54, A and cp26, B) are completely represented in *B.garinii*. Eight plasmids of *B.burgdorferi* are highly similar (cp32: L, M, N, O, P, R, S; lp56: Q). These seem to be also completely represented by *B.garinii* plasmid contigs although neither an exact assignment of individual contigs to a specific *B.burgdorferi* plasmid nor the calculation of their copy number is possible. The remaining plasmid contigs of *B.garinii* show also similarities to *B.burgdorferi* plasmids, but some regions are either unique to *B.burgdorferi* or have low DNA similarities. In four contigs, we observed similarities to different plasmids of *B.burgdorferi* indicating breakage/fusion points between different plasmids.

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Figure 1. View of the *B.burgdorferi* genome indicating similarities to sequences of the *B.garinii* genome. The calculation of the similarities was done using BLAST. Threshold for identity was 75% on a length of 40 bases. Portions of the *B.burgdorferi* genome with matches are depicted in red, unmatched in blue. The complex consisting of eight nearly identical plasmids is drawn in yellow. Here, no exact orthologous regions could be defined since sequences as well as CDSs are redundant. The scale for the plasmids is 10x magnified compared to the chromosome.

Chromosome

In our low redundancy project, the average coverage of the chromosome assembly is 3.38 (Table 1). The initial assembly of the *B.garinii* chromosome had 260 gaps (comprising sequencing and clone gaps). After applying gap closure procedures, we obtained one contig covering almost (99.5%) the complete linear *B.burgdorferi* chromosome. Despite the low redundancy of the sequence reads >80% of the chromosome is endowed with a error frequency of <10⁻⁴. The overall expected error rate is 0.26% (Supplementary Figure S1). The comparative analysis shows an unbroken collinearity between the *B.burgdorferi* and *B.garinii* chromosomes. The only parts of the *B.burgdorferi* chromosome with no counterpart in *B.garinii* reside in both telomeric regions with a size of 168 and 8458 bp, respectively. Since ends of linear chromosomes are not clonable without further manipulation, the missing bases on the left end are probably due to a cloning bias. It was previously shown that the right

end of the chromosome in *B.burgdorferi* exhibits length variations in defined steps in different strains (33). The shorter right end of the *B.garinii* chromosome is comparable to one form of these stepwise variable lengths of the *B.burgdorferi* chromosomes.

Substitutions, insertions and deletions

Base substitutions are a measure for evolutionary distance between organisms. The overall identity of the *B.garinii* with the *B.burgdorferi* chromosome is 92.7%. A calculation based on the shared CDS on amino acid level gives the same result indicating an equal distribution of substitutions over the chromosome irrespective of information content. The decrease of similarity between the two chromosomes below 80% in three regions around the origin of replication as can be seen in Figure 2 is apparently caused by larger insertions and deletions (indels; Figure 2, number 4–6). In total, we found 66 482 single base substitutions (Table 2). Transitions and transversions are almost equally distributed in the genic and intergenic regions.

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Figure 2. A sketch of the *B.garinii* chromosome compared to the collinear *B.burgdorferi* counterpart. Base identity of *B.garinii* versus *B.burgdorferi* is shown as green line, GC content as light blue line (both left scale) and GC skew as purple line. All values were calculated using a sliding window of 10 kb with a step width of 1 kb. Positions of all indels (deletions–negative peaks, insertions–positive peaks) in the alignment of the two analysed chromosomes are shown as black bars (right scale). Indels with a distance <10 bp to each other were defined as one single indel. All larger indels are indexed with numbers in red, exact positions are given in Supplementary Table S1. A cluster of five insertions appears in this figure as one peak (number 4), in a similar manner two deletions are located near position 540 000 (number 5 together with an insertion). The deletion of 8458 bases at the telomeric end is not shown.

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Table 2. Frequency of single base substitutions in the collinear genomic elements of *B.burgdorferi* and *B.garinii*

Besides the shorter right end of the chromosome, we found eight insertions and six deletions with a size >100 bp (Figure 2, numbers 1–8; Supplementary Table S1) in the *B.garinii* chromosome as major structural differences relative to the chromosome of *B.burgdorferi*. The largest observed insertion with a size of 1878 bp is caused by a duplication of a region containing the *bmpA* gene and part of the *bmpB* gene (see below) resulting in a tandem repeat of these genes (Figure 2, number 3). A series of five insertions is separated by short orthologous sequences of at most 470 bases (Figure 2, number 4;

Supplementary Table S1). This cluster of insertions is located in a region containing two tRNA genes (tRNA-Ile-1, tRNA-Ala-1) and expands only intergenic regions.

Indel region 5 consists of an insertion of 538 bases (Figure 2, peak 3) followed by two deletions of 211 and 498 bases, respectively, which are separated by 133 bases. In this indel region 5 resides the gene encoding inositol monophosphatase (BB0524) in *B.burgdorferi*. This gene is only partly represented (59 of 284 amino acids) in *B.garinii*. The eighth insertion (Figure 2, number 7) is located in an intergenic region. According to the observed indel regions, two 'hotspots' for rearrangements on the *Borrelia* chromosome could be defined: indel region 4 and indel region 5. Most interestingly, these two regions are located in close vicinity to the origin of replication of the chromosome at position 475 kb.

All deletions >100 bases including the missing chromosomal ends comprise 10 448 bases, all insertions 4099 bases. Thus, the *B.garinii* chromosome is by 0.7% shorter than that of *B.burgdorferi* (Table 2).

Genes

A comparative annotation of the *B.garinii* chromosome was performed using the previously published *B.burgdorferi* gene prediction and annotation (GenBank NC_001318 [GenBank] .1) (16). In parallel, we used GeneMarkS for ab initio gene predictions. This program was not able to detect 36 of the original gene predictions on the *B.burgdorferi* chromosome (Supplementary Table S3). Two of these non-verified coding sequences are fused to neighbouring coding sequences in *B.garinii* (BB0410, BB0510). The failure of GeneMarkS to identify BB0412 is possibly a false negative result, as the program predicted the ortholog in *B.garinii*. Additional ten potential genes of the remaining 814 genes annotated on the chromosome of *B.burgdorferi* are not predicted on the chromosome of *B.garinii* (Supplementary Table S4). Interestingly, these genes are annotated in *B.burgdorferi* only as predicted coding region without any other supportive evidence like similarities to other genes. Twelve predicted *B.burgdorferi* genes are fused in *B.garinii* (BB0078 + BB0079, BB0356 + BB0357, BB0410 + BB0411, BB0510 + BB0511, BB0521 + BB0522, BB0710 + BB0711). Eight annotated genes show extensive length divergence and overlap only partly their *B.garinii* counterpart, seven of which are altered due to differing open reading frames (ORFs) on an otherwise orthologous genomic sequence (BB0475, BB0524, BB0532, BB0546, BB0591, BB0749, BB0758). The *lmp1* gene (BG0212) is affected by the largest deletion in *B.garinii* (Figure 2, number 1) and thereby shortened by 648 bases at the 5' end. In summary, 786 GenBank annotated genes of *B.burgdorferi* are supported by GeneMarkS predictions in both *Borrelia* species and thus most likely represent the orthologous set of chromosomal genes. The length of 452 orthologs is unaltered, 255 genes are longer in *B.burgdorferi* and 79 genes are longer in *B.garinii*, but since these length differences are only small, the core of the deduced amino acid sequence is not affected.

BB0086 is split in *B.garinii*. Two genes are affected by a large duplication in *B.garinii* leading to a second copy of *bmpA* (BB0382) and a partial copy of *bmpB* (BB0383; Figure 2, number 3). Due to nonsense mutations, this partial copy is represented in the predicted gene set by four small ORFs.

In addition to the 807 predicted *B.garinii* genes with homologous sequences (including split, fused and other altered gene structures) in annotated CDS of *B.burgdorferi*, GeneMarkS predicts 33 further genes. These genes are comparably small (<60 amino acids) and most likely represent false positive predictions. This is further underlined by the fact that four of these potential genes lie within rRNA and tRNA gene regions and additional four predictions are apparently derived from the truncated copy of *bmpB*. The additional 39 genes on the *B.burgdorferi* chromosome, which are predicted by GeneMarkS, may also be false positive predictions.

On the DNA level, no predicted protein-coding gene of *B.garinii* is identical to its ortholog in *B.burgdorferi* whereas 20 tRNA genes (out of a total of 33 tRNA genes) are identical to their orthologous counterparts. Interestingly, the mutations seem to affect tRNA genes not randomly, since most sequence changes occur in non-unique tRNA genes (11 of 13). All four copies of tRNA-Leu, all three copies of tRNA-Ser, two of the three tRNA-Arg copies, and the second copy of tRNA-Lys and tRNA-Thr, respectively, are mutated.

Due to the high similarity of the chromosomes, the statistics of the codon usage shows only a slight difference between the two species. For example, in *B.burgdorferi* a higher preference for TTG as a start codon than in *B.garinii* predicted genes could be observed (Supplementary Table S2).

On the protein level, only 11 of all *B.garinii* genes are not altered in comparison to *B.burgdorferi*. This includes the ribosomal proteins *rpsU*, *rpsL*, *rpmG*, *rpsJ*, *rpsS*, *rpmF*, a putative subunit K of an ATPase, the flagellar motor switch protein *fliG-2*, the phosphocarrier protein *ptsH-2*, and the chemotaxis-related proteins *cheX* and *cheY-3*.

Additional 25 genes are affected by conservative exchanges with amino acids having the same chemical properties, thus increasing the number of highly conserved proteins to 38; not surprisingly 18 genes of this expanded group code for ribosomal proteins.

As an indication for positive selection, 94 (11.7%) of all orthologous genes and genes with similar sequences contain more non-synonymous than synonymous exchanges. Of these, 61 have no functional assignments. Interestingly, a higher than average proportion of the deduced amino acid sequences is predicted to contain transmembrane domains (39% compared to 26% for all proteins, Supplementary Table S8). The remaining 33 predicted genes are listed in Table 3. Many of these proteins seem to be located, according to their function, on the

surface of the cell.

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Table 3. Proteins with ascribed function with more non-synonymous (Ka) than synonymous (Ks) codons

Plasmids

Different strains of the same *Borrelia* species can carry different sets of plasmids. The differing plasmid repertoire of the cells can be partly a function of the living conditions (34). Additionally, strains can lose parts of their equipment due to a lack of selection pressure (35). Since the primary assembly of the chromosomal reads without additional gap closure sequences resulted in 91% coverage of the chromosome, we may conclude that also the plasmids are represented in the same range of coverage. Using the whole-genome shotgun data, it was possible to assemble two individual plasmids of the *B.garinii* PBI strain completely (Table 1). These two plasmids are highly similar and collinear to the linear plasmid lp54 and the circular plasmid cp26 of *B.burgdorferi* B31. The nearly two times higher coverage of one of these plasmids (lp54) compared to that of the chromosome indicates that it should be present in about two copies per cell. Compared to the chromosome, we find an equal number of base substitutions (8.9%) on the cp26 plasmid. Interestingly, with 15% substitutions are twice as frequent on plasmid lp54. Most remarkably, the transition:transversion ratio on lp54 is 3:2, whereas that of the chromosome as well as that of cp26 is approximately 4:1 (Table 2). The coding capacity of both plasmids is comparable to their counterparts in *B.burgdorferi*. Only three *B.garinii* lp54 genes predicted by GenMark are orphans, whereas the majority of predicted genes (49 of 74) have orthologs in the *B.burgdorferi* plasmid: 22 of the predicted genes match as pairs to different parts of 11 *B.burgdorferi* genes indicating nonsense mutations leading to split CDSs in *B.garinii*. On the other hand, 14 predicted *B.burgdorferi* genes have no counterparts on the *B.garinii* plasmid (Supplementary Table S5). None of these genes has an ascribed function. Interestingly, the *B.burgdorferi* lp54 gene family (BBA68 to BBA73) appears to be almost completely conserved in *B.garinii*, only BBA72 being split into two predicted genes. The analysis of the coding capacity of cp26 showed that all 26 predicted *B.garinii* genes have orthologs in *B.burgdorferi*, only three (BBB15, BBB20, BBB21) of the *B.burgdorferi* predicted genes are orphans (Supplementary Table S5). Interestingly, two cp26 encoded genes are subjected to a rapid positive selection: ospC and BBB08. OspC is well characterized as outer surface protein, whereas BBB08 so far has no assigned function. On the other hand, only 17 of the 55 lp54 encoded proteins may be subjected to a neutral evolution or purifying selection.

The assignment of clusters of orthologous groups (COG) (36) to the predicted proteins is clearly different between the chromosome and the plasmids. Whereas 81.6% of the chromosome-encoded orthologous proteins

can be assigned to a COG, only 53.9% (cp26) and 26.5% (lp54) can be categorized this way (Supplementary Table S6).

All other *B.garinii* plasmids are represented in our assembly as 37 contigs >2 kb comprising 239 kb. We here refer to these plasmid parts as variable plasmid segments (VPS). As it is known from the assembly of the *B.burgdorferi* plasmids, there are redundant segments distributed over several plasmids (37,38). The same holds true for the *B.garinii* VPS. Some redundant regions containing polymorphisms could separately be assembled. Yet, the read coverage of some contigs is higher than that of the chromosome and cp26. Thus, it is very likely that these portions of the VPS represent paralogous sequences. Therefore, they cannot be assembled properly into individual plasmids. Accordingly, due to this non-unique nature of many segments in the plasmids, a clear 1:1 assignment to defined plasmids of *B.burgdorferi* is not possible.

A GeneMarkS gene prediction revealed 338 complete and truncated potential protein-coding genes on the VPS: 284 of these predicted genes have matches to predicted genes in the *B.burgdorferi* genome on protein level. Many, mainly small genes (117) show partial matches, but 167 predicted genes exhibit similar lengths in both genomes. One of these genes is a true ortholog to BB0844, which is encoded on the chromosome in *B.burgdorferi*. All other predicted genes are related to plasmid-encoded genes.

To get an overview of our *B.garinii* VPS assembly, we performed a BLAST search on nucleotide level of all plasmid-derived contigs against a database of *B.burgdorferi* plasmids. This BLAST search revealed that 70% (167 kb) of the VPS are similar enough to *B.burgdorferi* plasmid sections to be detected (Figure 1). The remaining sequences (73 kb) have no detectable similarity to *B.burgdorferi* plasmids on the DNA level. Yet, if we search for similarities on protein level, we find matches (40–70% identity on amino acid level) for all contigs to putative *B.burgdorferi* plasmid-encoded proteins. The contig with the lowest similarity to *B.burgdorferi* sequences is contig AY722928 [GenBank], which encodes a *vls* locus involved in antigenic variation in the mammalian host (39). This locus is located on lp28-1 in *B.burgdorferi*. Thus, all VPS sequences seem to be represented in *B.burgdorferi* plasmids.

We then asked, which part of the coding capacity of the *B.burgdorferi* plasmids is present in *B.garinii*. Since small predicted genes are often false positives, for a reliable comparison of the gene sets, we took into account only *B.burgdorferi* plasmid-derived proteins >100 amino acids, and searched for their counterparts in the whole *B.garinii* shotgun data. Only one protein each from plasmids lp5 (T), lp54 (Q), lp21 (U), cp32-3 (S), lp25 (E), lp28-1 (F), lp28-2 (G), lp28-3 (H) and lp28-4 (I) had no counterpart. The failure to detect these proteins in the *B.garinii* genome could be due to missing shotgun data. Interestingly, from plasmids lp38 (J) 15 of 21 and from plasmid lp36 (K) 9 of 24 proteins were not represented in the shotgun data

(Supplementary Table S7). A more detailed inspection revealed that the predicted protein-coding regions from these two plasmids that have matches to the shotgun data belong to protein families. Members of these protein families are encoded also on different plasmids. These results taken together indicate that *B.garinii* PBi lacks the counterparts of plasmids lp38 and lp36.

Since the copy number of plasmid segments can affect the phenotype of *Borreliae* (40), we also analysed the data in this respect. According to the BLAST hits against *B.burgdorferi* proteins >100 amino acids plasmids lp28-1 (F), lp28-3 (H), lp28-4 (I), lp17 (D), lp25 (E) and lp5 (T) are present in one copy per cell. Plasmid lp28-2 is represented with two independent segments in our assembly. Thus, it should exist as two slightly different copies in *B.garinii*. For the proteins from the highly redundant cp32 and cp56 plasmids, we observed between three and four copies each. Furthermore, since parts of these segments are identical, the assembly resulted in parts of these segments in a higher coverage than average. We thus estimate that this plasmid group is at least present in five copies. The plasmid cp9 encodes similar proteins as the cp32 plasmids, albeit with much lower similarities. Thus, we are not able to determine whether a counterpart of this particular plasmid belongs to the *B.garinii* genome.

DISCUSSION

Closely related pathogenic species can cause different symptoms or even a different disease based on unique species-specific features. To reveal the molecular basis of such differences, one can examine the genomic repertoire of two closely related species. In cases where previous studies revealed not only a high similarity on DNA level but also almost complete collinearity of the chromosomes or large segments thereof, a direct comparative analysis without a completely finished genome is feasible (41,42). The limitation of this method lies in the inability to resolve highly rearranged and repetitive structures of the genome. Here, we report on an in-depth exploration of the *B.garinii* genome in comparison to that of *B.burgdorferi*. The analysis revealed a complete collinearity of the chromosome as well as of two plasmids in the compared species. The other genome parts seem to be subjected to rapid sequence changes as well as rearrangements and duplications. Thus, *Borreliae* genomes seem to consist of a remarkably invariant part mainly responsible for survival in ticks and a fluctuating part responsible for pathogenicity and disease symptoms in humans.

Borrelia core gene repertoire annotation on the invariant collinear genome fraction

An ab initio prediction of genes is flawed by the uncertainty as to what constitutes a real gene in a given organism (43). In recent years, it was successfully shown that a comparative genomics approach could improve if not largely replace an annotation from scratch (44,45). If the relationship between the two compared species is close enough, most

if not all genes should have an ortholog in the sister species. Thus, the use of orthology information is the best approach to discern between true genes and false positive predictions or species-specific adaptations. On the stable fraction of the genome (chromosome and plasmids lp54 and cp26), we could easily define, which of the predicted genes in both organisms are orthologous gene pairs. Some of the orthologous pairs found may not be true genes especially if they are short. But the high conservation would suggest at least a regulatory function of these chromosomal regions. In total, we found 861 orthologous gene pairs compared to 955 (*B.burgdorferi*) and 932 (*B.garinii*) predicted genes on these conserved genomic elements. Since the other plasmids seem to be dispensable for viability, this set of orthologous protein-coding gene pairs is very likely the basic repertoire of *Borrelia burgdorferi sensu lato* species. This is supported by the fact that for example the proteins OspA, OspB and OspC, which seem to be required for survival in the tick midguts (46), are encoded on the two conserved plasmids.

COG categories show interspecies relationships, i.e. only proteins more common than for one genus can be categorized this way (36). The fraction of categorizable proteins decreases from the chromosome to cp26 and then to lp54 considerably, whereas the fraction of positively selected genes increases. Recently, it was shown that *B.burgdorferi* is not able to live without cp26 (47). Our analysis further supports the view that cp26 is an essential genome part of *Borrelia* species. Both the presence of two members of a family of genes (*ospA* and *ospB*) known to be needed in tick midguts and the conserved collinearity of the plasmid leads us to the conclusion that lp54 very likely plays also a pivotal role in survival in ticks. Since most of the encoded proteins are subjected to positive selection, we hypothesize that lp54 is a major player in host response evasion.

Selection pressure

The analysis of the ratio of non-synonymous to synonymous substitutions shows that the overwhelming majority of functionally described proteins of the chromosome (475/513; 92.6%) are under neutral evolution or purifying selection. The remaining 33 proteins (Table 3) show positive selection to different degrees. As expected, we find many surface-associated proteins in this list, which may be involved in the escape from the host response. On the other hand, of the 295 proteins described only as predicted or hypothetical, 56 (19%) seem to be under positive selection. This is a larger fraction than that in the subset with known function. Many of the proteins with functional assignment as well as the hypothetical proteins seem to be located at the surface of the cell and presumably generate a strain variability to fool the immune response from ticks as well as from humans. Thus, this subset of hypothetical proteins would be a rewarding target for further functional studies.

Variable plasmid complement of *B.garinii*

Only an isolation and analysis of each single plasmid would allow the

resolution of the entire plasmid fraction of *B.garinii*. But since the whole-genome shotgun data should represent most parts of the plasmids (see Results), we are able to calculate similarities between contigs and plasmids to uncover the relationship of *B.garinii* to *B.burgdorferi* plasmids, and to reveal differences in plasmid content between the two species.

The noncollinear plasmids are not only rearranged but also the encoded proteins are not as conserved as in the collinear genome parts. Yet, two-thirds of the *B.garinii* VPS sequences are similar enough on DNA level to match *B.burgdorferi* counterparts. For the remaining sequences, no DNA similarities could be found. Nevertheless, our ability to detect similarities on the protein level shows that these plasmid parts are more likely subject to an accelerated evolution than unique *B.garinii* genome constituents: 167 of the 338 predicted genes have counterparts of similar length in the *B.burgdorferi* genome. Despite their high divergence, they may thus be ascribed as proteins with orthologous functions. BB0844 is encoded on the right-end extension of the *B.burgdorferi* chromosome and its ortholog on a plasmid in *B.garinii*. This may point to a mechanism, which enables variable *Borrelia* chromosome lengths by exchanging parts between chromosome and distinct plasmid segments.

Based on comparisons of the coding capacity of the VPS with the *B.burgdorferi* plasmids, we are able to define, which plasmids are presumably present in *B.garinii* and which not. With this analysis, we could show that two plasmids (lp38, J and lp36, K) are missing from the *B.garinii* genome. This is especially important since lp38 encodes a member of the osp family of proteins (ospD), a protein family widely studied for its role in pathogenicity and survival in the hosts.

It was shown that interplasmidal duplications and rearrangements are able to change the virulence phenotype of *Borrelia* species (40). Thus, it is of high value to know, which plasmids or segments are represented more than once in the genome. Generally, duplication and diversification events seem to affect the same plasmids in the two species. We also see an amplification of the cp32 plasmid sequences, although there may be a few copies less in *B.garinii* than in *B.burgdorferii*. Most other plasmid sequences are represented only as one copy in each species. Yet, in *B.garinii* plasmid lp28-2 sequences underwent also duplication and diversification. Thus, while the main difference in the protein family sets lies in the presence or absence of plasmids lp36 and lp38, additional diversity is achieved not only by mutation of single genes but also by a duplication of lp28-2, and possibly modification of the copy number of cp32 plasmid sequences.

Evolution and species definition

In previous studies (48–50), a large genetic distance between and within *Borrelia* species was observed on the basis of highly variable genes and genomic regions. Most of the genes examined are located on

the plasmids, which are far less conserved than the chromosome. In contrast, we found a strong conservation of similarity and collinearity between *B.garinii* and *B.burgdorferi* not only of the chromosome but also of two plasmids. Interestingly, the amino acid identity of chromosomally encoded proteins is not higher than the conservation of the whole chromosome on DNA level. Thus, despite positive selection observed for specific proteins (Table 3), the chromosome on the whole is subjected to a neutral evolution.

Since plasmid repertoire variability is observed also in closely related strains causing similar disease patterns, the species definition is based only on the chromosome. Proteins encoded on the chromosome may also slightly influence the disease pattern. As discussed before, the survival in ticks may be mediated by the two collinear plasmids. Thus, pathogenicity in vertebrate hosts could be mainly dependent on the VPS. Future work has to determine, which plasmids or plasmid parts are lost during loss of pathogenicity. Since both *Borrelia* species have amplified mainly the same plasmids or segments thereof, it is conceivable that it is not enough to keep one member of each plasmid-encoded protein for maintenance of pathogenicity. Rather a larger number of paralogous proteins encoded by redundant plasmids could be required to successfully infect vertebrates.

Possibly the genes under positive selection are also causative for the symptom differences of the various *Borrelia* species. Thus, both the variable plasmid part of the genome and the positively selected genes represent prime targets for further functional studies.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at NAR Online.

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