Telomere Exchange between Linear Replicons of \textit{Borrelia burgdorferi}

Wai Mun Huang, Margaret Robertson, John Aron, and Sherwood Casjens

Department of Pathology and HSC Core Facilities; University of Utah Medical School, Salt Lake City, Utah 84132

Received 5 January 2004/Accepted 26 March 2004

Spirochetes in the genus \textit{Borrelia} carry a linear chromosome and numerous linear plasmids that have covalently closed hairpin telomeres. The overall organization of the large chromosome of \textit{Borrelia burgdorferi} appears to have been quite stable over recent evolutionary time; however, a large fraction of natural isolates carry differing lengths of DNA that extend the right end of the chromosome between about 7 and 20 kbp relative to the shortest chromosomes. We present evidence here that a rather recent nonhomologous recombination event in the \textit{B. burgdorferi} strain Sh-2-82 lineage has replaced its right chromosomal telomere with a large portion of the linear plasmid lp21, which is present in the strain B31 lineage. At least two successive rounds of addition of linear plasmid genetic material to the chromosomal right end appear to have occurred at the Sh-2-82 right telomere, suggesting that this is an evolutionary mechanism by which plasmid genetic material can become part of the chromosome. The unusual nonhomologous nature of this rearrangement suggests that, barring horizontal transfer, it can be used as a unique genetic marker for this lineage of \textit{B. burgdorferi} chromosomes.

The Lyme disease spirochetes, \textit{Borrelia burgdorferi}, \textit{Borrelia afzelii}, and \textit{Borrelia garinii}, and most likely all members of the \textit{Borrelia} genus, have an unusual genome that is made up of an approximately 900-kbp linear chromosome and numerous smaller linear and circular extrachromosomal DNA elements. The nucleotide sequence of the genome from one \textit{B. burgdorferi} isolate, B31 MI, has been determined (8, 14). Analysis of this strain has shown that the chromosome carries nearly all of its housekeeping genes. Its 21 extrachromosomal elements include 12 linear plasmids 5 to 54 kbp in length and nine circular plasmids 9 to 32 kbp in length that carry mostly genes of unknown function that are unique to the genus \textit{Borrelia} (8). Examination of numerous independent natural \textit{B. burgdorferi} isolates has shown that sequences similar to the plasmids in isolate B31 are usually found on plasmids of similar size in other isolates (1, 9, 10, 13, 15, 17–19, 21, 23–25, 27, 31, 32, 34), and a substantial fraction of these plasmids appear to be present in all isolates examined to date (16, 23).

Physical maps of the chromosomes from 25 geographically diverse \textit{Borrelia} isolates that either cause Lyme disease or are close relatives of ones that do have been constructed (5, 11, 22; R. van Vugt and S. Casjens, unpublished data). These maps are all extremely similar, indicating that the chromosomes carry little gross structural variation across the more than 10 species that make up this cluster of species. This lack of organizational variation in the chromosome is not universal in bacteria (3); for example, the genomes of several gram-negative and -positive bacteria have been found to vary up to tens of percentages in gene content and/or have substantial rearrangements among different isolates (12, 20, 33). On the other hand, there appear to have been numerous and substantial recent rearrangements in the linear plasmid portion of the \textit{B. burgdorferi} B31 MI genome. Analysis of this genome sequence has shown that 10 of the 12 B31 linear plasmids carry a large amount of DNA that appears to be in a state of mutational decay and not to encode functional proteins. This most likely is a result of the numerous recent duplicative DNA rearrangements (8), since such duplications may release one of the copies from selection, allowing it to mutationally decay. This decay is now observed as a large number of “pseudogenes” that contain many translational frame-disrupting mutations.

Although the bulk of the chromosome appears to be very stable, we have previously noted that in \textit{B. burgdorferi} sensu stricto the extreme right end of the chromosome (as defined by Fraser et al. [14]) is variable in length, in that additional sequences extend the right telomeric region in some isolates. BB0843 is the rightmost gene in the 903-kbp “constant portion” of the chromosome; it is only a few hundred base pairs from the right telomere in \textit{Borrelia} isolates with the minimum-size chromosome, such as N40, R-IP3, W919-23, and HB19 (7, 14). The sequenced chromosome of strain B31 MI has 7.2 kbp of DNA beyond BB0843 at its right telomere. This DNA is almost entirely made up of sequences that have paralogs on the B31 MI plasmids and, except for two apparently intact genes, appears to be largely in a state of severe mutational decay (8). Different \textit{B. burgdorferi} isolates carry different lengths of DNA to the right of gene BB0843; in 31 \textit{B. burgdorferi} isolates that we have examined, 21 carry such extensions (7). None have right-end extensions longer than the 19-kbp extension of Sh-2-82. In this study we examine the nature of the right-end chromosomal extension in \textit{B. burgdorferi} isolate Sh-2-82.

\textbf{MATERIALS AND METHODS}

\textbf{Bacterial strains.} \textit{B. burgdorferi} strain Sh-2-82 was isolated from an \textit{Ixodes scapularis} tick on Shelter Island, N.Y. (26). Passage 6, passage 166, and passage 320 cultures of strain Sh-2-82 were the kind gifts of Tom Schwan, Patti Rosa, and Janis Weis, respectively. Strain 297 passage 5 was the kind gift of Justin Radolf. Strains JD1, 21305, 22921, 29968, 30757, and 28534 are described in the work of Casjens et al. (7); although we do not know exactly how each source laboratory performed these passages, each one typically represents six to nine generations.
B. burgdorferi strains were propagated, and whole-cell DNA was prepared in agarose blocks as previously described (6). Contour-clamped homogeneous electric field (CHEF) electrophoresis, DNA transfer to nylon membranes, and Southern hybridizations were performed as previously described (6).

DNA manipulation and nucleotide sequence determination. The nucleotide sequence of strain Sh-2-82 chromosomal right-end DNA was determined using a primer walking strategy and whole-genome DNA as template. Sequencing reactions with oligonucleotide primers 25 to 28 nucleotides long and Big Dye dideoxy terminator mix (PE Applied Biosystems, Foster City, Calif.) were performed in a thermal cycler as follows: 69 or 99 cycles at 95°C for 0.2 min, 1°C/s to 55°C, 55°C for 0.2 min, 1°C/s to 60°C, and 60°C for 4 min. Automated sequencers (PE Applied Biosystems) were used according to the manufacturer’s recommendations. After nearly complete sequences of the two regions described in the text were determined, all ambiguities were resolved by sequencing PCR DNA fragments amplified from Sh-2-82 (passage 320) DNA. All of the sequence reported here was thus determined from both strands, except for the 63-bp repeat regions which could be approached only from one direction on whole-genomic DNA (see elsewhere).

Nucleotide sequence accession numbers. These sequences have been deposited in GenBank with accession numbers AY309080 and AY309081 for the left and right unique regions, respectively.

RESULTS

Right chromosomal telomere of B. burgdorferi isolate Sh-2-82. We found by Southern analysis that a DNA probe (probe 1; Table 1 and Fig. 1) from the left part of the 7.2-kbp B31 extension hybridizes with the “extra” DNA at the right telomere of the strain Sh-2-82 chromosome (data not shown), suggesting at least some similarity between the right telomeric regions of these two isolates. To characterize the long right telomeric extension of the Sh-2-82 chromosome in more detail, a primer walking strategy with whole-cell DNA as template was used to determine the nucleotide sequence rightward from the conserved BB0843 gene (see Materials and Methods). Several oligonucleotide sequences were chosen from within the conserved strain B31 gene BB0843 that primed a rightward sequencing reaction on Sh-2-82 template DNA. When a good-quality sequence was obtained with one of them, a second primer was chosen from within the sequence thus determined to sequence further to the right and so on. In this way, nucleotide sequence was determined for 4,173 bp of strain Sh-2-82 DNA rightward from within gene BB0843. The left 2,692 bp of this sequence are nearly identical to similarly located B31 sequence; Sh-2-82 has a 264-bp deletion (between bp 1879 and 1880) relative to B31, but it is otherwise 99.6% identical to the

![FIG. 1. Sequence relationships among B. burgdorferi B31 MI linear plasmid lp21 and chromosomal right telomeric regions. Shading of the chromosomal and plasmid DNAs represented by rounded bars indicates the following: solid black, the constant portion of the chromosome (which extends from the left chromosomal end through gene BB0843); dark gray, the right-end extension on the B31 chromosome and very similar sequence on the Sh-2-82 chromosome; white, the unique (non-63-bp repeat) regions of B31 plasmid lp21 and the very similar sequence on the Sh-2-82 chromosome; hatched, the region of 63-bp tandem repeats on lp21 and similar sequence on the Sh-2-82 chromosome. Gray areas between DNAs highlight regions of similarity between adjacent DNAs; the darker gray (and percent values there) indicate similarities to sequence determined in this study. Black arrows indicate putative intact genes (pseudogenes are not shown). Numbered black bars show the locations of DNA probes used in this study (also Table 1). The Sh-2-82 nucleotide sequences determined here are indicated by open bars immediately above the kilobase pair scale, whose zero point is the right end of the constant portion of the chromosome.](http://jb.asm.org/ on October 23, 2017 by guest)
parallel region of the B31 chromosome (Fig. 1). This region in Sh-2-82 contains an apparently intact gene BB0844 homolog that should encode a protein of unknown function that is 99.4% identical to its B31 ortholog. The right, 1,481 bp of these 4,173 bp have no similarity to the B31 chromosome, but the sequence is nearly identical to B31 linear plasmid lp21 (similarity extends rightward from bp 2438 on the published lp21 sequence [8]). Between bp 2693 and 3870 the Sh-2-82 sequence is identical to that of the B31 lp21 plasmid. Immediately to the right of bp 3870 the sequence contains about 5.5 tandem copies (there are many more repeats beyond these copies; see below) of 63-bp direct repeats that are very similar to the B31 lp21 63-bp repeat tract which lies at an identical location there. This long repeat tract blocks further rightward sequence determination by this strategy because primers in this region would not have unique binding sites on Sh-2-82 DNA.

Since the right portion of the above Sh-2-82 chromosomal sequence is nearly identical to lp21, we attempted to determine whether sequences similar to those on lp21 to the right of its 63-bp repeat tract (the lp21 “right unique region” [Fig. 1]) were also present near the Sh-2-82 right telomere. Opposing oligonucleotides that amplify a 1.2-kbp section of the right unique region of B31 lp21 plasmid (oligonucleotides A and B; Table 1; Fig. 1) were used in a PCR and found to amplify identically sized fragments from whole-cell B31 and Sh-2-82 template DNAs. This Sh-2-82 templated PCR product, called probe 2 (Table 1), was then used in a Southern analysis of Sh-2-82 DNA to determine its location in that strain’s genome. It hybridized only with the 920-kbp chromosome in whole Sh-2-82 DNA (Fig. 2) and only with the chromosome’s rightmost BssHIII and SgrAI fragments (37 and 34 kb, respectively; data not shown, but see Fig. 4 below), showing that this amplicon lies within 34 kbp of the chromosomal right end. Restriction mapping of the rightmost chromosomal BglII, BsrGI, and NcoI fragments with the same DNA probe (data not shown), in combination with the sequences determined above and below, proved unequivocally that probe 2 hybridizes with lp21-like sequences that are present near the Sh-2-82 right telomere and distal to the 63-bp repeat tract.

Oligonucleotides A and B were therefore used to prime a second sequencing primer walk on whole-cell Sh-2-82 DNA. The resulting sequence contig had 63-bp repeats at its left end (Fig. 1), and as the walk neared the right telomere, the sequence quality deteriorated, presumably due to competition between primer annealing and snap-back of the telomeric hairpin (see below). A DNA probe derived by PCR amplification near the right end of this sequence contig (probe 3) also hybridized only with the rightmost Sh-2-82 chromosomal BssHIII fragment in a Southern analysis (data not shown). In order (i) to determine the location of the right chromosomal telomere more precisely and (ii) to confirm that the sequence determined was actually at this location, probe 3 was used in Southern analyses of singly and doubly restricted whole-genome Sh-2-82 DNA to construct a restriction map of this region of the chromosome that unambiguously located six restriction enzyme cleavage sites (data not shown)—two XbaI sites and one EcoO109I, BstXI, HindIII, and EcoRI site each (Fig. 1). Each of these sites was correctly located in the primer walk sequence, except the rightmost XbaI site, which is very near the telomere and outside the primer walk region. Each of the latter four enzymes’ right-end restriction fragments extended about 600 bp beyond the right end of the primer walk contig. Rapid reannealing (snap-back) experiments analogous to those done previously for DNA at the left end of the Sh-2-82 chromosome (7) showed that the two strands of the rightmost EcoO109I, BstXI, EcoRI, and HindIII fragments rapidly anneal (data not shown). We conclude that these fragments are in fact right-terminal chromosomal fragments and that these fragments, like other linear replicon terminal fragments in Borrelia, are tipped by a covalently closed DNA hairpin.

To determine the sequence of the tip of the Sh-2-82 right chromosomal telomere, we gel purified the size fraction 3.8 to 4.1 kbp of HindIII-restricted whole-cell Sh-2-82 DNA, which includes the rightmost HindIII fragment; nicked this DNA’s terminal hairpin by SI nuclease treatment; ligated the resulting DNA to a blunt-ended, double-stranded synthetic oligonucleotide; and used PCR to amplify between a primer that anneals within the primer walk sequence contig and a primer that anneals to the terminal, ligated synthetic sequence as previously described (7). The sequence of this PCR product was determined directly using both the amplification and internal primers, and in addition, the PCR product was cloned into the plasmid vector pCR4-TOPO (Invitrogen, Carlsbad, Calif.) and the sequences of the DNA inserts in several representative plasmids were determined. The resulting sequence overlapped the primer walk right unique region sequence in the manner expected and contains a correctly located telomere-proximal XbaI site (above) 220 bp from the right end; it extends the right unique sequence to the right by about 600 bp as predicted by the restriction map and contains at its extreme right terminus the 23-bp sequence ‘TTTTATCTAAAAAAAAACTAAA TT-3’, which is similar to the sequences at the tips of other known Borrelia telomeres (7, 14, 15, 35). We also used this methodology to determine the previously unknown telomeric...
sequence at the right end of strain B31 MI linear plasmid lp21. The reported sequence of plasmid lp21 (GenBank accession no. AE001582 [8]) was found to be lacking the rightmost terminal 25 bp of the plasmid; this 25-bp sequence, 5'-GCTTTA TACTAAAAAAAACTAATTT-3', is identical to the sequence that we determined for the right tip of the Sh-2-82 chromosome. One or a few nucleotides could be missing from the telomeres of these sequences due to possible removal by S1 nuclease. Merging of the right primer walk and the sequence of the terminal PCR amplicon resulted in 4,664 bp that contain the entire right unique region (Fig. 1). The leftmost 501 bp of this sequence contig is composed of approximately eight tandem copies of the same inexact 63-bp repeat present at the right end of the left unique region and is very similar to the parallel portion of lp21. The unique 4,163 bp to the right of bp 501 is 99.9% identical to the parallel region of lp21. In addition to three single nucleotide differences between Sh-2-82 and B31 in this region, there is an inversion of 5 bp (ACTTG centered on bp 2097). These 5 bp precisely separate a perfect 17-bp inverted repeat, which could have mediated the inversion. None of these differences disrupts the reading frame in which it occurs.

We also characterized in more detail the putative 63-bp repeat tract that lies between the two regions that were sequenced above. Southern analysis using a strain B31 probe from the 63-bp repeat region of lp21 (probe 4; Table 1 and Fig. 1) showed that the only Sh-2-82 sequences capable of hybridizing with this probe lie on the chromosome and within the 37-kbp rightmost BssHII chromosomal restriction fragment (Fig. 3). Restriction enzymes MseI (cuts at TTAA) and Asel (ATTAAT) cut the 72% A+T Borrelia DNA extremely frequently, the former giving rise to fragments that are nearly all less than 500 bp in length. There are no MseI or Asel sites in the B31 lp21 repeat tract, so in B31 they give rise to unusually large 63-bp-repeat-containing 11 ± 1.0-kbp and 13 ± 1.0-kbp DNA fragments, respectively (8). The Sh-2-82 passage 320 and B31 MI repeat tract-containing MseI and Asel fragments were indistinguishable in size (data not shown), suggesting that these enzymes also do not cut the Sh-2-82 repeat tract and that the tract length is about the same in B31 linear plasmid lp21 and the Sh-2-82 chromosome. The sequence determinations of the right and left unique regions combined with the length of the 63-bp-repeat tract thus show that there is about 19 kbp of extra sequence at the Sh-2-82 right chromosomal telomere, which agrees well with our previous estimate from the length of right terminal restriction fragment sizes (7).

The experiments described above show that DNA probes derived from the right unique region, the 63-bp-repeat tract of B31 lp21, and the Sh-2-82 right unique region all hybridize exclusively to the 37-kbp rightmost chromosomal BssHII fragment. Figures 2 and 3 show that this chromosomal hybridization is present at passages 7, 166, and 320 in culture (there are three to five generations per passage), although there is also an approximately 2-kbp-shorter form present (as the majority) in passage 7; it is not present by passage 166. Thus, although a number of linear plasmids have been lost with passage in culture (reference 26 and our unpublished analysis), we have no evidence of any substantial changes in the Sh-2-82 right telomeric region during this period; in particular we note that the length of the 11-kbp form of the chromosomal 63-bp-repeat tract appears not to have changed significantly in about 1,500 generations (313 passages). The 13 63-bp repeats in the sequence reported here are all represented exactly among the 34 types of slightly different repeats present in B31 lp21, but the repeat types are not present in the same order, suggesting that gene conversion may have been active in this region since the two sequences diverged.

Other B. burgdorferi isolates with right telomeric chromosomal extensions. We previously reported that several other B. burgdorferi isolates, 21305, 22921, 29968, and JD1, have approximately 19-kbp right-end chromosomal telomeric extensions (all quite similar in length to Sh-2-82) and 28534 has an approximately 16-kbp extension (7). In addition, strain 297 (28) has a right-end chromosomal extension similar in length to that of Sh-2-82 (J. Aron and S. Casjens, unpublished data). Each of these right-end extensions was found to hybridize to our probe 1 (Table 1). In order to determine whether the extensions in these strains might also contain lp21-like sequences, a Southern analysis was performed using probe 5 (Table 1) from the right unique region of B31 linear plasmid lp21 (this probe is 99.9% identical to, and hybridizes equally well with, the homologous Sh-2-82 right unique region). Figure 4 shows that 28534 and 29968 chromosomes do carry probe 5 sequences near their right telomeres, since the probe hybridizes to the right-end BssHII fragments in these strains (it also hybridizes to this fragment from strain 297; data not shown). MseI and Asel cleavage experiments (as in Fig. 3) estimated the length of the 63-bp-repeat tract to be 9 kbp in 28534 and 11 kbp in 29968 (data not shown). Both of these strains also carry 24- to 25-kbp linear plasmids that hybridize (but not as strongly) with this probe (Table 2). Circular DNAs are not resolved into tight bands by the CHEF electrode pulse program used, so the bands observed almost certainly represent linear DNAs. Passage 7 and 166 Sh-2-82 also carry apparently...
linear plasmids that react very weakly to this probe but which are lost by passage 320 (Table 2). The apparent change in size of these plasmids with passage is difficult to assess since (i) less total DNA was present in the passage 7 lanes; (ii) only a subset of the cells might carry the larger hybridizing plasmids at passage 7, which could have expanded by passage 166; or (iii) there may have been DNA rearrangements among plasmids during growth in culture.

Probe 5 does not react well with the chromosome in strains JD1, 21305, and 22921 and hybridizes much better with 23- to 27-kbp linear plasmids in these strains (Fig. 4). These strains also carry 63-bp repeat tracts on similar-size plasmids based on parallel Southern analyses with probe 4, and they carry 9-kbp, 9- and 13-kbp, and 9- and 11-kbp reactive MseI fragments, respectively (data not shown). In a panel of 13 additional isolates with shorter or no right-end extensions, only isolate 30757 (7-kbp right extension) was found to carry the 63-bp-repeat-hybridizing sequences, and these were on an approximately 24-kbp linear plasmid (23).

To test whether the 297, 28534, and 29968 chromosomal right-end extensions are the result of a recombination event that was identical to that of Sh-2-82, we PCR amplified (using oligonucleotides E and F, Table 1) and sequenced a 1,024-bp region that includes the Sh-2-82 recombination event from each strain (from 2515 through 3538 on the Sh-2-82 right unique region sequence). In all three cases the sequence of the amplified product was identical to the parallel Sh-2-82 sequence. Thus, since their recombination joints are identical, the right telomere replacement by lp21 in these three strains almost certainly happened only once in a common ancestor. It is interesting that Stevenson and Miller (30) recently found that Sh-2-82 and 297 also share extensive sequence identity on their cp26 and cp32 circular plasmids, supporting the notion that these are very closely related isolates. Isolates Sh-2-82, 28534, and 29968 are from ticks captured in New York, Maryland, and Connecticut, respectively, and 297 is a human isolate from Connecticut, indicating that geographic movement of the affected chromosome can happen before random mutagenic changes occur in what is thought to be nonfunctional DNA (e.g., in the gene BBU04 pseudogene homologous region). Thus, four of the seven known B. burgdorferi isolates with >15-kbp right-end extensions carry the same extensive homology to B31 linear plasmid lp21 at the right end of their chromosomes. Strains Sh-2-82, 297, 28534, and 29968 have lp21-like extensions at their right chromosomal telomeres, with the 63-bp repeat tract of 28534 being 2 kbp shorter than those of Sh-2-82 and 29968 (strain 297 was not tested). Since they carry probe 1- but not probe 4- or 5-hybridizing DNA near their right chromosomal telomeres, strains JD1, 21305, and 22921 appear to have some other DNA replacing and extending the tip of an ancestral B31-like chromosome; it seems likely that this DNA will be derived from some other Borrelia linear plasmid.

**DISCUSSION**

The structure of the right end of the Sh-2-82 chromosome is most easily explained by a simple, single recombination event between the right telomeric region of a B31-like chromosome and a B31 lp21-like linear plasmid, so that the rightmost 16 kbp of the plasmid replaced the distal 4 kbp of that chromosome (Fig. 1). In the Sh-2-82 chromosome right-end extension the proximal 2,692 bp are 99.6% identical to the B31 chromosome, and the distal lp21-like region unique (non-63-bp repeat) sequence is 99.9% identical to linear plasmid lp21. These extremely high similarities allow deduction of the nature of the recombination event in the Sh-2-82 progenitor. Figure 5 shows that there is an abrupt switch in Sh-2-82 from similarity to the
TABLE 2. Southern hybridization with right-end DNA probes

<table>
<thead>
<tr>
<th>Culture</th>
<th>Replicon</th>
<th>Probe&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1</td>
</tr>
</tbody>
</table>

| Sh-2-82 (p7) | Chromosome lp18 | +  | +  | +  | +  | +  |
| Sh-2-82 (p166) | Chromosome lp26 | ND | +  | +  | +  | +  |
| Sh-2-82 (p320) | Chromosome lp28 | ND | +  | +  | +  | +  |
| JD1 (p16) | Chromosome lp26 | +  | +  | ND | –  | –  |
| 297 (p6) | Chromosome lp29 | +  | +  | ND | ND | ND | +  |
| 21305 (p < 20) | Chromosome lp25 | +  | +  | ND | –  | –  | –  |
| 22921 (p8) | Chromosome lp27 | +  | +  | ND | –  | –  | –  |
| 28534 (p4) | Chromosome lp26 | +  | +  | ND | +  | +  | +  |
| 29968 (p < 20) | Chromosome lp23 | +  | +  | ND | +  | +  | +  |
| 30757 (p5) | Chromosome lp24 | –  | ND | –  | +  | +  | +  |

<sup>a</sup> Probes are described in Table 1 and Fig. 1. Table symbols are as follows: –, no hybridization; +, moderate hybridization; ++, strong hybridization; ND, not done. Parentheses in the table data indicate very weak hybridization that most likely represents cross-hybridization to nonorthologous, paralogous sequences.

B31 chromosome to similarity to the B31 lp21 linear plasmid. There are only 2 bp of sequence identity at the point where this recombination must have taken place, making this an essentially nonhomologous event. There are three additional locations in B. burgdorferi linear replicon sequences in which a similar deduction can be made regarding past rearrangements: (i) the 265-bp deletion in the non-lp21-like Sh-2-82 rightward extension compared to the homologous B31 sequence (Fig. 1 and 5); (ii) linear plasmid Ip56 in B31, which appears to have been generated by the integration of a 31-kbp circle (homologous to the cp32 plasmids) into an approximately 24-kbp linear plasmid (8); and (iii) a 900-bp inversion near the left end of B31 Ip56 relative to paralogous sequences on B31 plasmids lp31 and Ip36 (8). In these cases there are 0, 2, and 2 bp of identity (the latter occurred within a 20-bp region of complex imperfect similarity), respectively, at the points where the recombination events must have occurred. In addition, two apparent deletions of circular cp32 plasmids have been reported: an approximately 14-kbp deletion to form cp18 in strain N40, where the putative crossover took place at one side of a 6-of-8-bp match in its closest relative, plasmid cp32-1 (29), and an approximately 10-kbp deletion to form cp18-2 in strain 297, whose crossover point has no base pairs of identity in the two possible parental sequences in plasmid cp32-7 (2). Nonhomologous rearrangements appear to have occurred relatively frequently on these DNAs. However, given the huge number of possible events, it is unlikely that identical nonhomologous recombination events of independent origin will be found, so these unique rearrangements (e.g., the chromosome-lp21 novel joint described here in Sh-2-82, 297, 28534, and 29968) should be useful as genetic markers in the characterization of B. burgdorferi populations.

The telomeric sequences at the extreme right ends of the Sh-2-82 chromosome and the B31 lp21 plasmid are identical, but they have interesting differences from the previously characterized Borrelia termini. All previously known Borrelia telomeres have an absolutely conserved TAGTAYANA sequence (5’ to 3’ in the upper strand when the telomere is on the left; Fig. 6) that is 14 bp from the end and a highly conserved TATAAT sequence that is either 1 or 4 bp from the terminus (4). The Sh-2-82 and lp21 right-end terminal sequences determined here have a TAGTAYANA sequence that is 14 bp from the apparent end; however, they have no convincing TATAAT sequence. Compared to the other telomere sequences, AAA TAG or TAGTTT occupies the type 1 or type 2 “TATAAT positions,” respectively, and these telomeres do not fit either type (Fig. 6). Tourand et al. (32a) have used mutant target sites to show that the sequence in at least parts of the TATAAT and TAGTA regions of a type 1 telomere is indeed important for
telomere formation by protelomerase in vitro. However, the observations made here point out that the target specificity of the *Borrelia* telomere resolution machinery is not yet fully understood, especially in the “TATAAT portion” of the target sequence, and future new telomere sequences can be expected to shed additional light on terminal sequence constraints and protelomerase recognition.

The data presented here make a particularly clear case for a past recombinational exchange event between a *Borrelia* linear plasmid and a chromosomal telomere. This, coupled with the finding that the distal, non-Sh-2-82-like portion of the B31 lp21 plasmid are shown in the middle, with the covalently closed hairpin telomere on the left for ease of comparison with previous publications (3, 6, 12, 27). Gray boxes highlight the TATAAT and TAGTAYANA conserved regions (see text).

Casjens (3) and Tourand et al. (32a) have pointed out that the previously characterized *Borrelia* telomeres appear to fall into two categories, type 1 and type 2, in which an apparently conserved TATAAT sequence is present 1 or 4 bp, respectively, from the terminus. Among the nine previously sequenced telomeres, five are type 1 and four are type 2; the consensus sequences of the two types are shown at the top and bottom of the figure (R = A or G; Y = C or T; W = A or T; K = G or T; M = A or C; N indicates that three different base pairs are present among the known members of that type).

The directionality of the postulated right-end telomere exchange event is most likely the replacement of the chromosomal telomere by the lp21 sequences (as opposed to generation of lp21 by “excision” from the end of an Sh-2-82-like parental chromosome), since the recombination event truncated an apparently intact lp21 gene, BBU04, which has paralogs on several other plasmids; it is very unlikely that the intact lp21 gene BBU04 would have been generated by a nonhomologous excision event. Our previous studies suggested that the rightmost 7.2 kbp of the linear chromosome of strain B31 are all derived from plasmids in a similar but more complex manner, so that the Sh-2-82 right end is the result of at least two successive rounds of telomerere replacement. *B. burgdorferi* isolates B31 and Sh-2-82 were both isolated from *I. scapularis* ticks on Shelter Island, N.Y., but they do not appear to be especially closely related, since the constant portions of their chromosomes have several restriction site polymorphisms among the relatively small number of such sites examined (5); we do note, however, that these two isolates carry an apparently identical cp32 plasmid (30). The lp21 recombination event in the Sh-2-82 chromosome must have been rather recent, since its sequences remain more than 99% identical to the B31 lp21 sequences.

The B31 right-end extension contains two apparently intact genes, BB0844 and BB0852, both of which have paralogs on the B31 linear plasmids. These are surrounded by at least nine severely damaged plasmid-like genes (8), as if plasmid sequences had been joined to the right end of the chromosome, after which most of the plasmid genes were allowed to decay but the two currently intact genes were perhaps selected to remain functional. According to this model, the B31 chromosome had sequences added some time ago and mutational decay processes have partially removed those genes that are of no use there, while the lp21 addition to the Sh-2-82 chromosome (in which three possibly functional genes and six apparent pseudogenes of linear plasmid lp21 replaced one possibly functional gene and several pseudogenes of a B31-like chromosome) happened only rather recently, and further decay has barely begun on the newly added region. This appears to be an evolutionary mechanism which is able to sequentially move genetic material from linear plasmids onto the end of the linear *Borrelia* chromosome.

**ACKNOWLEDGMENTS**

We thank Tom Schwan, Patti Rosa, Janis Weis, Justin Radolf, and Tom Anderson for *Borrelia* strains.

This work was supported by NIH grant AI49003 to S.C.

**REFERENCES**


