

Genome Analysis and Strain Comparison of Correia Repeats and Correia Repeat-Enclosed Elements in Pathogenic *Neisseria*

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Whole genome sequences of *Neisseria meningitidis* strains Z2491 and MC58 and *Neisseria gonorrhoeae* FA1090 were analyzed for Correia repeats (CR) and CR-enclosed elements (CREE). A total of 533, 516, and 256 copies of CR and 270, 261, and 102 copies of CREE were found in these three genomes, respectively. The lengths of CREE range from 28 to 348 bp, and the lengths of multicopy CREE appear mainly in the ranges of 154 to 156 bp and 105 to 107 bp. The distribution of CREE lengths is similar between the two *N. meningitidis* genomes, with a greater number of 154- to 156-bp CREE (163 and 152 copies in *N. meningitidis* strain Z2491 and *N. meningitidis* strain MC58, respectively) than 105- to 107-bp CREE (72 and 77 copies). In the *N. gonorrhoeae* strain FA1090 genome there are relatively more 105- to 107-bp CREE (51 copies) than 154- to 156-bp CREE (36 copies). The genomic distribution of 107-bp CREE also shows similarity between the two *N. meningitidis* strains (15 copies share the same loci) and differences between *N. meningitidis* strains and *N. gonorrhoeae* FA1090 (only one copy is located in the same locus). Detailed sequence analysis showed that both the terminal inverted repeats and the core regions of CREE are composed of distinct basic sequence blocks. Direct TA dinucleotide repeats exist at the termini of all CREE. A survey of DNA sequence upstream of the sialyltransferase gene, *Ist*, in several *Neisseria* isolates showed that 5 *N. meningitidis* strains contain a 107-bp CREE in this region but 25 *N. gonorrhoeae* strains show an exact absence of a 105-bp sequence block (i.e., the 107-bp CREE without a 5' TA dinucleotide) in the same region. Whole-genome sequence analysis confirmed that this 105-bp indel exists in many homologous 107-bp CREE loci. Thus, we postulate that all CREE are made of target TA with indels of various lengths. Analysis of 107-bp CREE revealed that they exist predominantly in intergenic regions and are often near virulence, metabolic, and transporter genes. The abundance of CREE in *Neisseria* genomes suggests that they may have played a role in genome organization, function, and evolution. Their differential distribution in different pathogenic *Neisseria* strains may contribute to the distinct behaviors of each *Neisseria* species.

Since the report of repetitive extragenic palindromic (REP) sequences in *Escherichia coli* and *Salmonella enterica* serovar Typhimurium (25), short dispersed repetitive elements have been increasingly identified in prokaryotes (35). Genome analyses have confirmed the extensive repetition of REP sequences and REP elements in *E. coli* (4). Short dispersed repetitive elements have also been identified in other prokaryotes for which the complete genome sequences have been analyzed (for a review, see reference 44).

Correia et al. identified a 26-bp sequence as a repetitive element in the pathogenic *Neisseria* spp. (8). Subsequent studies showed that Correia repeats (CR) often constitute parts of longer repetitive sequence elements (9, 29). By using two-dimensional S1 nuclease heteroduplex mapping, Correia et al. estimated that in *Neisseria gonorrhoeae* there are ca. 20 copies of 152-bp elements whose ends are composed of inverted repeats of the 26-bp CR sequence (9). Gotschlich et al. identified a 105-bp sequence element that also contains the CR sequences as terminal inverted repeats, and these authors esti-

ated that the 105-bp element is present at least 20 times in the genome of *N. gonorrhoeae* R10 (23). The 154-bp Correia element(s) (CE) was considered to be a transcriptional terminator in the division cell wall (*dcw*) cluster of *N. gonorrhoeae* CH811 and a search of the sequence databases revealed another 19 copies of similar elements adjacent to different neisserial genes (14). In genome sequences of *Neisseria meningitidis*, 163 copies of the 26-bp inverted repeat were found in *N. meningitidis* MC58 (49) and a total of 286 CE (sequences bounded by 26-bp inverted repeats) were found in *N. meningitidis* Z2491 (41). More recently, Mazzone et al. reported a total of 270, 259, and 110 copies of *nemis* (*Neisseria* miniature insertion sequences) in whole genomes of *N. meningitidis* Z2491 and MC58 and *N. gonorrhoeae* FA1090, respectively (37).

In the present study, we first analyzed the sequence conservation in all CR found in three completed neisserial genomes: *N. meningitidis* Z2491 (41), *N. meningitidis* MC58 (49), and *N. gonorrhoeae* FA1090 (<http://www.genome.ou.edu/gono.html>). We then used the most conserved region of the CR to identify those sequence elements that were enclosed by an inverted pair of the CR in these three complete genome sequences. We analyzed the detailed sequence features of CR-enclosed elements (CREE) and determined DNA sequences upstream of the sialyltransferase gene, *Ist*, in several *Neisseria* strains to

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identify the size of indels in the loci of 107-bp CREE. The common sequence features identified among all CREE and the potential mechanisms for the formation of CREE may assist in an understanding of their origin, propagation, and function within the genome. The distinction between *N. meningitidis* and *N. gonorrhoeae* may supplement our understanding of the differential pathogenesis of these two clearly related but distinct pathogens.

MATERIALS AND METHODS

Bacterial strains and growth conditions. *N. gonorrhoeae* F62 was obtained from P. Frederick Sparling of the University of North Carolina at Chapel Hill; *N. meningitidis* MC58 from E. Richard Moxon, University of Oxford, United Kingdom; *N. meningitidis* DNM51 from David Dyer, University of Oklahoma Health Sciences Center; and *N. meningitidis* NMB13, DNM3, and FAM18 from David S. Stephens (Emory University, Atlanta, Ga.). Clinical isolates of *N. gonorrhoeae* were obtained from MCP Hahnemann University Hospital in Philadelphia (labeled as the "A" series of strains) and from patients attending a sexually transmitted disease clinic in Baltimore (labeled as the "B" series of strains and provided to us by John Zenilman of Johns Hopkins University). To minimize genetic change, stocks of all strains and clinical isolates were stored at -70°C and experiments were started from frozen stocks with minimal passage (one to three times). Growth of *Neisseria* isolates was performed at 37°C in a 5%, humidified CO_2 incubator (Forma Scientific, Marietta, Ohio) on GC agar (Difco, Detroit, Mich.) with supplement (30). Bacteria were suspended in sterile phosphate-buffered saline with 0.1% (wt/vol) gelatin and 0.01% (wt/vol) each of CaCl_2 and MgCl_2 (38).

PCR amplification of *lst* upstream regions (5'-*lst*). To obtain template DNA for amplification of 5'-*lst*, several colonies of *Neisseria* were suspended in 20 μl of sterile H_2O and boiled for 10 min, chilled on ice for 2 min, and then incubated with RNase A (final concentration of 16 $\mu\text{g}/\text{ml}$) at 37°C for 30 min. These suspensions were centrifuged at $8,000 \times g$ for 5 min, and supernatants were collected and placed on ice for immediate use or stored at -20°C for later use.

To amplify the 5'-*lst*, we used primers synthesized according to sequence information derived from *N. meningitidis* MC58 (accession no. U60660) (18). The forward primer, FuseBam-For1 (5'-CGCTGGATCCGACATCAATATCG G), starts 124 bp from the initiation codon of putative isocitrate dehydrogenase gene *icd* on its minus strand and contains a *Bam*HI restriction site at its 5' end (underlined). The reverse primer, FuseBam-Rev1 (5'-CAAAGGATCCCTTTT CAAGCCC), starts 24 bp from the initiation codon of *lst* on its minus strand and also contains a *Bam*HI site (underlined). PCR was performed in thin-wall glass capillary tubes by using an Air Thermo-Cycler (Idaho Technology Model 1605; Idaho Falls, Idaho) with a two-step program. Step one consisted of five cycles of 94°C for 0 s, 40°C for 0 s, and 72°C for 15 s. Step two consisted of 30 cycles of 94°C for 0 s, 55°C for 0 s, and 72°C for 15 s.

DNA sequencing. DNA fragments amplified by PCR were cut from agarose gels after electrophoresis and purified by using a Wizard PCR Preps DNA purification system (Promega, Madison, Wis.). Nucleotide sequences were determined with the terminal primers described above by cycle sequencing with fluorescently labeled dideoxynucleotides (DyeDeoxy terminators; Perkin-Elmer) by using automated DNA sequencing facilities at the MCP Hahnemann School of Medicine.

Genome sequences. The genome sequence of *N. meningitidis* MC58 (49) was obtained from The Institute of Genomic Research (ftp://ftp.tigr.org/pub/data/n_meningitidis/). The genome sequence of *N. meningitidis* Z2491 (41) was obtained from the Sanger Center (http://www.sanger.ac.uk/Projects/N_meningitidis/). The genome sequence of *N. gonorrhoeae* FA1090 was from the University of Oklahoma (<http://www.genome.ou.edu/gono.html>). The last search for CR and CREE in *N. gonorrhoeae* FA1090 was based on sequence released on 15 September 2000. It contains 2,154,110 sequence characters.

BLAST analysis of DNA sequences. BLAST searches were performed by using BLASTN (version 1.4.7) in the version 9.1 Wisconsin package (Genetics Computer Group). The ungapped BLASTN program was used for searching homologues of the prototypic 26-bp CR (5'-GTACCGTTTTTGTAAATCACTA TA) and a 105-bp sequence element was identified in the DNA sequence upstream first in *N. meningitidis* MC58. The GCG output information was parsed into a Microsoft Access database by using a custom written program. The copy number, length, and genomic location of the archived CR and CREE sequence information was analyzed by using the filtering, sorting, and querying functions of the Microsoft Access program. For sequence comparison, we converted all

sequence elements to the same stranded orientation. Consensus sequences for CRs were found by using GCG's PILEUP and LINEUP programs. Consensus sequences for 105-bp sequence elements and other CREE were obtained by using the CLUSTALW multiple sequence alignment tool (<http://dot.imgen.bcm.tmc.edu:9331/multi-align/multi-align.html>) and the BoxShade program (http://www.ch.embnet.org/software/BOX_form.html). For analysis of the flanking sequences of the 105-bp elements, a custom written computer program was used to retrieve each 105-bp sequence and its 1-kb flanking region on each side. These retrieved 2,105-bp sequences were used as queries for BLASTN searches against the GenBank "nr" (nonredundant) database by using NCBI's BLAST 2.0 website (<http://www.ncbi.nlm.nih.gov/BLAST/>) with the default parameters.

FINDPATTERNS analysis of DNA sequences. Single completed genomic sequences were divided into smaller segments of ca. 300 kb to comply with the length limitation of the GCG programs. These segments were then converted into a GCG-utilizable format by using the FROMFASTA and the data set programs in the GCG package. To avoid potential failure in detecting sequences at the ends of the divided segments, the complete genome sequences were divided into both eight and seven segments, and the number of findings were examined for agreements. The presentation of repeats in the genomes are based on original sequence coordinates indicated in the complete linear sequences. Searching for CREE was performed by using the FINDPATTERNS program of the GCG package. The search pattern used was TATAGTGGATT(N)[0, 328]AATCCACTATA, where "(N)[0, 328]" means any sequence with a length between 0 and 328 bp. This maximum limit of 328 bp in the core region is due to the total length limit of 350 bp in the FINDPATTERNS program. During the execution of the program, we used mismatch values from 0 to 6. The GCG output information was parsed into a Microsoft Access database by using a custom-written program. Then the copy number, length, and genomic location of the archived sequence information were analyzed by using the filtering, sorting, and querying functions of Microsoft Access.

Gene and ORF identification. The matched segments were manually inspected for genes or open reading frames (ORFs) as identified in the genome reports for *N. meningitidis* Z2491 (41) and *N. meningitidis* MC58 (49). The distance of 105-bp elements relative to those identified genes or ORFs was determined by manual inspection of the genomic coordinates. Fine contextual analysis was then performed for *N. meningitidis* MC58 by using ACEDB (R. Durbin and J. T. Thierry-Mieg. 1991. A *C. elegans* database. Documentation, code, and data available at <http://www.acedb.org>), as used in the annotation of the genome sequence of this strain (49).

RESULTS

CR and CREE. Using the 26-bp prototype CR (8) as a query, we found in ungapped BLAST analyses 533, 510, and 256 homologous sequence segments in the genomes of *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090, respectively (Table 1). Among these matched segments, >75% are longer than 23 bp. Examination of alignments showed that all of these CR-homologous sequences have a conserved 11-bp terminal region: TATAGTGGTTT. Thus, in the present study, we refer to any sequence that contains this 11-bp sequence but is ≤ 26 bp as a CR. We refer to any sequence that contains this 11-bp sequence but is >26 bp as CEs. Since many CEs contain CR as terminal inverted repeats (TIR) (web table 1 [web tables and figures may be found at www.pages.drexel.edu/~rr37/]), we used CREE to describe CR that are confined by two inverted CR.

FINDPATTERNS analysis of the sequence pattern TATA GTGGTTT(core)AAACCACTATA yielded 270, 261, and 102 copies of CREE for *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090, respectively (Table 1). Upon manual inspection of the CREE sequences, we noticed that some longer CREE contain an unpaired internal CR. This unpaired internal CR, when paired with a terminal CR, constitutes a shorter CREE that was also retrieved by the FINDPATTERNS program and listed in a shorter length category. For clarity, we refer to CREE enclosed by one pair of terminal

TABLE 1. Numbers of CR and CREE in three neisserial genomes

Strain	No. of CR found by BLAST with matched length of the query (bp):							CREE found by FINDPATTERNS with mismatch allowed ^a :				% CR in CREE ^b
	26	25	24	23-14	13	12-11	All (11-26)	0	1	2	All (0-2)	
<i>N. meningitidis</i> Z2491	381	14	5	18	103	12	533	183 (2)	+59 (1)	+28 (7)	270 (10)	99.4
<i>N. meningitidis</i> MC58	377	7	2	25	94	11	516	176 (0)	+51 (2)	+34 (4)	261 (6)	100.3
<i>N. gonorrhoeae</i> FA1090	153	13	26	21	37	6	256	59 (0)	+17 (2)	+26 (2)	102 (4)	78.1

^a Values indicate the number of additional CREE found upon increasing the number of allowed mismatches. The value in parentheses indicates the number of simple CREE contained inside complex CREE.

^b Calculation based on two and three CR in simple and complex CREE, respectively.

CR as “simple” CREE and CREE containing an additional unpaired internal CR as “complex” CREE (Fig. 1). Since the number of nucleotides counted for a complex CREE contain the number of nucleotides of its enclosed simple CREE, the total nonredundant copies of CREE were adjusted by subtracting the number simple CREE from their parent complex CREE. The adjusted nonredundant copies of CREE are 260, 255, and 98 copies for *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090, respectively (Table 1). In comparing the numbers of nonredundant CREE with the numbers of CR, we observed that almost all of the CR are contained in CREE in the two *N. meningitidis* genomes. In contrast, 22% of the CR are not contained in CREE in the *N. gonorrhoeae* FA1090 genome (Table 1).

Length profiles of CREE. CREE found in the above FINDPATTERNS analysis were sorted according to their length and copy number within each length (Table 2). In the three genomes, there are 50 categories of CREE based on length. *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090 have 35, 26, and 19 of the 50 different CREE lengths, respectively. The total nucleotides in all CREE in *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090 account for 1.67, 1.55, and 0.59% of their total genomic nucleotides, respectively.

Of the 50 CREE lengths, 9 are shared by all three neisserial genomes (Table 2) (341, 156, 155, 154, 107, 106, 105, 73, and 71 bp). Some CREE lengths are present in both meningococcal sequences but absent in the gonococcus (213, 185, 142, 110, 97, 96, 70, and 28 bp). CREE of 104 and 69 bp are present only in *N. gonorrhoeae* FA1090 and *N. meningitidis* Z2491. The 153-bp CREE is present only in *N. gonorrhoeae* FA1090 and *N. meningitidis* MC58. Fifteen CREE lengths are unique to *N. meningitidis* Z2491 (348, 325, 266, 219, 199, 182, 171, 159, 150, 144, 111, 108, 101, 98, and 72 bp). Seven CREE lengths are unique to *N. meningitidis* MC58 (334, 329, 220, 197, 125, 115, and 92 bp). *N. gonorrhoeae* FA1090 contains seven unique CREE lengths (319, 245, 204, 169, 165, 116, and 87 bp).

Most of the CREE lengths occur one to three times in each

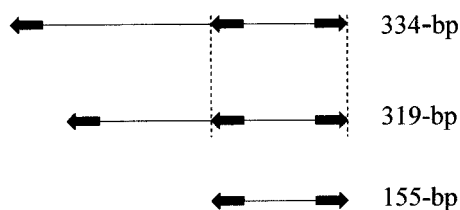


FIG. 1. Simple and complex CREE.

neisserial genome. Only the 154- to 156-bp and the 105- to 107-bp CREE have four or more copies per genome (Table 2). Interestingly, CREE of these lengths are also often found as simple CREE contained within complex CREE (Table 2). The longest simple CREE within the limitation of our FINDPATTERNS analysis is 341 bp, which was found only once in each of the three neisserial genomes. The 341-bp CREE of the two *N. meningitidis* strains are 100% homologous, whereas the 341-bp CREE of *N. gonorrhoeae* FA1090 shows little homology with the 341-bp CREE of *N. meningitidis*, having homologies of only 18 bp at the 5' end and 11 bp at the 3' end.

Sequence patterns of simple CREE. To facilitate sequence analysis, we oriented all simple CREE into the same sequence direction. We then inspected all sequences to determine whether CREE of the same length belonged to one or more subtypes. Finally, we aligned each CREE length and subtype that had two or more copies in the genome. We collected all of the consensus CREE sequences and the original sequences of the single-occurrence CREE into a new data set for further analyses.

For the convenience of sequence comparison and presentation, we separated the symmetrical terminal inverted repeat regions from the asymmetrical core regions of each CREE and ordered them in separate tables. The terminal inverted repeats of simple CREE are listed in web tables 2 to 4 for *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090, respectively, and the core regions of CREE from *N. meningitidis* strain Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090 are listed in web tables 5 to 7, respectively. Inspection of these consensus sequences further revealed the modular nature of the sequence composition in the TIR, as well as the core region of CREE (Fig. 2).

Short direct repeats flanking CREE. We compared the 5-bp terminal sequences of every CREE with their flanking 5-bp sequences. We found that TA duplication is the most common direct repeat in the flanking sequences bordering CREE, followed by 3- to 5-bp direct repeats such as TAT or ATA, TATA, TATAG, or CTATA (Table 3). Because all CREE were identified by using a pair of 11-bp inverted repeats that contain terminal TATA, the terminal regions of all CREE comprise TA direct repeats.

Genomic distribution of CREE. The distribution of all CREE in each genome of the three pathogenic *Neisseria* strains is presented in Fig. 3. From this presentation it is clear that CREE are distributed throughout the entire genome of each strain, although some local genomic regions have more of certain lengths of CREE than other regions. Whereas the overall distribution profiles are similar between the two *N.*

TABLE 2. Length profile of CREE in three neisserial genomes

Length (bp) and other parameters	<i>N. meningitidis</i> Z2491		<i>N. meningitidis</i> MC58		<i>N. gonorrhoeae</i> FA1090	
	Copy (<i>n</i>)	Internal CREE (<i>n</i>)	Copy (<i>n</i>)	Internal CREE (<i>n</i>)	Copy (<i>n</i>)	Internal CREE (<i>n</i>)
Length						
348	1	105				
341	1		1		1	
334			1	155		
329			1			
325	1	196				
319					1	155
266	1	107				
245					1	
220			1	156		
219	1	155				
213	1	155	1	155		
204					1	105
199	1	155				
197			1	155		
185	1	156	1	156		
182	1					
171	2	155				
169					1	
165					1	106
159	2	105 (1)				
157	1		3			
156	28		20		4	
155	127		120		19	
154	8		12		13	
153			2		2	
150	1					
144	1					
142	1		2			
125			2			
116					1	71
115			1	105		
111	1					
110	1		1			
108	1					
107	26		24		19	
106	30		30		17	
105	16		23		15	
104	1				1	
101	1					
98	1					
97	1		2			
96	2		2			
92			1			
87					1	
73	2		2		1	
72	1					
71	3		3		2	
70	1		2			
69	1				1	
28	1		2			
Length no.	35		26		19	
Total copies	270	10	261	6	102	4
No. of nonredundant copies ^a	260		255		98	
% Nucleotide in genome ^b	1.67		1.55		0.59	

^a Nonredundant CREE are calculated by subtracting the number of internal CREE from the total number of CREE directly counted by the FINDPATTERNS analysis.

^b The percentage of nucleotide in the nonredundant CREE in the total genome sequence.

meningitidis genomes, the CREE distribution in *N. gonorrhoeae* FA1090 is quite different from both *N. meningitidis* strains.

CREE (107 bp) as 105-bp indels upstream of the sialyltransferase gene, *lst*. During our study of the contribution of varia-

tions in sequence upstream of the *lst* (5'-*lst*) to the differences in the amount of sialyltransferase activity expressed by *N. meningitidis* MC58 and *N. gonorrhoeae* F62, we noticed that a 105-bp sequence element is absent in the 5'-*lst* of *N. gonor-*

Left-TIR		Core						Right-TIR	
Terminal	Sub-terminal	Neisseria	C1	C2	C3	C4	Sub-terminal	Terminal	
TATAGTGGTTAAA	TTTAAACAGTAC	Nm Z2491	16/14/5/0	3	50/45/39/37/0	34/33/31/25/0	GTCCTGATTTAAA	TTTAAACACTATA	
	AAAACAGTAC	Nm MC58	9/7/0	3	50/49/46/37/5/0	34/33/31/25/0	GTCCTGATTTT		
	AAAAA	Ng Fa1090	16	3	50/0	34/33/0	TTTTT		
	0						0		

FIG. 2. Common sequence blocks in CREE. The consensus sequences and common length of sequence fragments are presented in each sequence block. Slashes are used to separate alternative lengths of sequence fragments in each sequence block in the core region.

*rho*ae F62 but present in the 5'-*lst* of *N. meningitidis* MC58 (Fig. 4). This 105-bp sequence element is a 107-bp CREE when combined with the TA dinucleotide at its 5' flanking sequence. This result indicates that this 107-bp CREE reflects a 105-bp indel, a region of DNA that is present on the chromosome of an organism (insertion) but absent from closely related organisms (deletion), hence the word "indel." Further investigation of more *N. gonorrhoeae* and *N. meningitidis* strains showed that this 105-bp indel (107-bp CREE) is present in the 5'-*lst* of all of the five *N. meningitidis* strains studied but in none of the 25 *N. gonorrhoeae* strains studied (web figure 1), suggesting the differential distribution of this 105-bp indel is a common difference between *N. meningitidis* and *N. gonorrhoeae*.

Identification of unique loci for 105-bp indels in three neisserial genomes. The 105-bp indel upstream of *lst* in *N. meningitidis* MC58 was used as a query for ungapped BLASTN analysis against the databases of the three neisserial genomes. We identified 26, 23, and 17 copies of 105-bp indels in the genomes of *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090, respectively (web tables 8 to 10). These numbers are in close agreement with those found in the FINDPATTERNS analysis (Table 2), except that the latter analysis identified one more 107-bp CREE in *N. meningitidis* MC58 and two more 107-bp CREE in *N. gonorrhoeae* FA1090. To reveal the genetic locations of the 105-bp indels in the three neisserial genomes, we used the 2,105-bp sequence segments representing each of the 105-bp indel plus 1 kb of flanking sequence on either side as queries and performed self- and reciprocal-BLASTN analyses (i.e., 2,105-bp sequences from one strain were used for BLASTN analysis against the genome sequence from which it was derived, as well as against the genome sequences of the other two strains). From these analyses, we identified 51 unique loci for a total of 66 copies of 105-bp indels in the three *Neisseria* genomes (Table 4). Among these 51 unique loci, only one is shared by all three pathogenic neisseriae. There are 14 more loci shared between the two *N. meningitidis* strains. The other loci are unique to each one of the three neisserial strains. Four 105-bp indel-associated loci of *N. meningitidis* Z2491 are unique, whereas three 105-bp indel-

associated loci of *N. meningitidis* MC58 are unique. Two 105-bp indel-associated loci each from *N. meningitidis* Z2491 and *N. meningitidis* MC58 genomes do not show a significant match to the genomic of *N. gonorrhoeae* FA1090. Four 105-bp indel-associated loci of *N. gonorrhoeae* FA1090 are unique.

In most homologous loci, there is often an exact 105-bp indel that is either present or absent. However, some of the 105-bp indel-empty sites are filled with other lengths of DNA sequences (Table 4). These other indels range in length from 103 to 222 bp.

In general, 105-bp indels are scattered in each genome (Fig. 5A). However, the two *N. meningitidis* strains showed some similarity in the spacing pattern of some 105-bp elements. This similarity became clearer when the 14 matched 105-bp loci (Table 4) were plotted in pairs (Fig. 5B). Among these 14 pairs of matched 105-bp loci, 11 pairs are located in similar chromosomal locations. The other three pairs are in different chromosomal locations but the relative positions among these three loci are similar in the two *N. meningitidis* strains. When the DNA sequence segment containing these three 105-bp element loci was inverted, these three loci matched well between the two *N. meningitidis* genomes (data not shown). This result confirms that there is a major sequence inversion between the two *N. meningitidis* genomes (49).

TA direct repeats as a common feature in all loci of 105-bp indels. Comparison of the 5'-*lst* regions from the various *N. meningitidis* and *N. gonorrhoeae* strains showed that a TA dinucleotide exists in target sequences where the 105-bp indel inserts (Fig. 4 and web figure 1). Further examination of all 105-bp indels in the genomes of the three *Neisseria* strains confirmed that a TA dinucleotide always exists in places where a 105-bp indel may be inserted (web table 11).

Genes and ORFs associated with or adjacent to 105-bp indels. The homologous sequences matched by the BLAST analysis with the above 2,105-bp sequence queries were examined for genes and ORFs according to the published annotations for the two *N. meningitidis* genomes. Most 105-bp indels are intergenic (web tables 12 to 14), whereas some are located within ORFs; however, these ORFs are almost exclusively identified as transposases or insertion sequences except for two

TABLE 3. Short direct repeats flanking CREE in the three neisserial genomes

Neisserial strain	No. of short direct repeats upstream				CREE copy ^a (TATAG***CTATA)	No. of short direct repeats downstream			
	TA	TAT	TATA	TATAG		TA	ATA	TATA	CTATA
<i>N. meningitidis</i> Z2491	12	2	1	0	270	10	6	3	1
<i>N. meningitidis</i> MC58	16	2	1	1	261	17	4	2	0
<i>N. gonorrhoeae</i> FA1090	8	3	1	0	102	3	2	2	0

^a The copy number of CREE includes simple CREE contained inside complex CREE in order to show the flanking sequence features of all CREE.

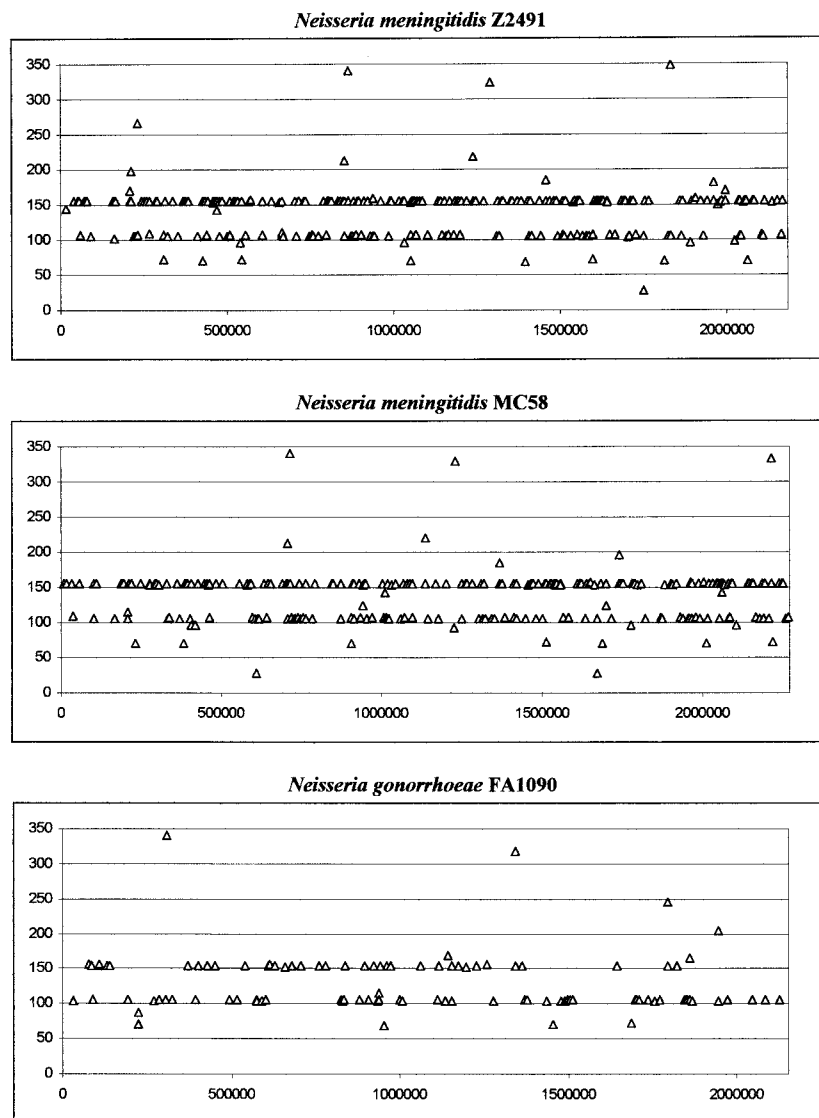


FIG. 3. Genomic distribution of CREE.

of the 17 copies of 105-bp indels in *N. gonorrhoeae* FA1090, which are inserted into ORFs other than those coding for transposases.

Importantly, some 105-bp indels are located close (<200 bp) to *Neisseria* virulence factors. For example, among the identifiable genes, 105-bp indels are associated with immunoglobulin A-specific serine endopeptidase (*iga2*), sialyltransferase (*lst*), outer membrane protein *opcA*, and class I outer membrane protein *porA* in the two *N. meningitidis* strains (web tables 12 and 13). The 105-bp indels are also associated with some metabolic genes such as glyceraldehyde-3-phosphate dehydrogenase (*gapA* and *gapC*) and acetate kinase (*ackA*) and also transporter genes for an ABC transporter ATP-binding protein. The 105-bp indels are adjacent to different sets of genes and ORFs in *N. gonorrhoeae* FA1090 (web table 14).

Detailed analysis of the *N. meningitidis* MC58 sequence using the ACEDB graphical interface revealed additional contextual information. In four instances (numbers 13, 16, 17 and

23) the 105-bp indels are located between convergent 3' ends of genes, and a further six elements (numbers 3, 4, 5, 6, 11 and 19) are located remotely from likely promoter locations so that they are unlikely to affect expression. One (number 15) is inserted into a dead gene. Four elements (numbers 10, 12, 21, and 22) are present in the same location within copies of the *IS1106* transposase. In seven instances (numbers 1, 2, 7, 8, 14, 18, and 20), the 105-bp indels are located such that they would be expected to form part of the promoter region of the associated gene, typically with the terminal TA forming part of a putative -10 element. The genes in which the 105-bp indels are likely to influence expression are (in the corresponding order): *pilF* (NMB0329), hypothetical proteins (NMB01387 and NMB0882), *lst* (NMB0922), a flavin reductase homologue (NMB1359), a hypothetical protein (NMB1782), and a peptidase homologue (NMB1877). The 105-bp indel associated with hypothetical protein NMB0882 is actually located between two divergent promoters but it is not located such that it would be

Nm MC58 AGGATTATTCTCTGTAGGTTGGTTTTTCTTTTGGAAACACATTGCGGG 50
 Ng F62 AGGATTATTCTCTGTAGGTTGGTTTTTCTTTTGGAAACACATTGCGGG
 *
 Nm MC58 GGAATGTGCGTGGCTATTATGGCATAATTTGGCGGCTTTGTTTCGCGCTTT 100
 Ng F62 GGAATGTGCGGCTATTATGGCATAATTTGGCGGCTTTGTTTCGCGCTTT
 *
 Nm MC58 GTTCGATCTTGGCGTGTGTTG[-----30bp-----] 150
 Ng F62 GTTCGATCTTGGCGTGTGTTGACACGCGCGGGGAAAGGAAAGGGGAAAT
 *
 [-----30bp-----]
 Nm MC58 AACCGCGCGCGCTGAAAGGAAGGGGAAATGGTTTTCCCGCGTTTGGCGG 200
 Ng F62 AACCGCGCGCGCGGAAAGGAAGGGGAAATGGTTTTCCCGCGTTTGGCGG
 * A *
 Nm MC58 CGG--TCGGAGGTGCTGTGCCTGATGTGCGGCGGCATATTTTCGGTGA 250
 Ng F62 CGGCGTCGCGCGCTTGTGCCTGATGTGCGGCGGCACATTTTCGGTAAAA
 ** * * * * *
 Nm MC58 TTGATTTTATAGTGGTTAAATTTAAACCAGTACAGCGTTGCCCTCGCCTT 300
 Ng F62 TTGATTTTAT[-----]-----
 Nm MC58 GTCGTA CTGCTGTACTGTCTGCGGCTTCGTCGCTTGTCTCGATTTAAA 350
 Ng F62 -----105bp-----
 -10
 Nm MC58 TTAAACCACATATAATTCGGTAACTGTCGGAATATCTGCTAAAATTC 400
 Ng F62 -----]ATATTCGCAACTGTCGGAATATCTGCTAAAATTC
 *
 [----13bp---]
 Nm MC58 GCATTTTCGTCGCGGG[----13bp---]ACACTCGGGCGTATGTTC 450
 Ng F62 GCATTTTCGTCGCGGGTTCCGACCGGGCACTCGGGCGTATGTTC
 **
 Nm MC58 ATTTGTCGGAATGGAGTTT-AGGGATATG 1st--->
 Ng F62 ATTTGTCGGAATGGAGTTT-AGGGATATG 1st--->
 SD IC

FIG. 4. A 105-bp indel (107-bp CREE) upstream of the sialyltransferase gene, *lst*. DNA fragments absent or duplicated are indicated by “[--number of base pairs--]” within or above the sequence, respectively. The single base (A) in the imperfect repeat is written below the aligned sequences. Single nucleotide differences are indicated by an asterisk. The Shine-Dalgarno (SD) sequence, the -10 box, and the initiation codon (IC) for *lst* are in boldface. Nm, *N. meningitidis*; Ng, *N. gonorrhoeae*.

expected to influence the adjacent *cysT* gene (NMB0881). Thus, 7 of 23 (ca. 30%) 105-bp indels in *N. meningitidis* MC58 may affect expression of associated genes.

DISCUSSION

We noticed discrepancies in the literature regarding CR and CE. For example, the 26-bp repetitive sequence originally reported by Correia et al. (8) was also named “*N. gonorrhoeae* inverted repeats” (49). CE (9) are sometimes termed “Correia full” (150 to 159 bp) or “Correia internal deletion” (~104 bp) (41) and sometimes “unit-length (154 to 158 bp) *nemis*” (for *Neisseria* miniature insertion sequences) and “internally rearranged (104 to 108 bp) *nemis*” (37). Estimates of the abundance of CR and CE also vary depending on the methodologies and query sequences used (37, 41, 49).

In the present study, we first identified the most conserved region of CR as an 11-bp terminal sequence. Then we used this 11-bp sequence to formulate a sequence pattern that allows retrieval of any sequence bracketed by an inverted pair of this 11-bp sequence (within a specified overall length range and mismatch in the 11-bp sequence region). The FINDPATTERNS analysis performed in this way is much easier than BLASTN analysis for identifying CREE. This FINDPAT-

TABLE 4. BLAST analysis of 105 bp loci among the three neisserial genomes^a

Locus	<i>N. meningitidis</i> Z2491		<i>N. meningitidis</i> MC58		<i>N. gonorrhoeae</i> FA1090	
	105No	Indel	105No	Indel	105No	Indel
1	1	105i	Not 105	103i	Not 105	104i
2	2	105i	23	105i	Not 105	105d
3	3	105i	No match	NA	Not 105	105d
4	4	105i	21	105i	9	105i
5	5	105i	No match	NA	Not 105	105d
6	6	105i	20	NA	Not 105	105d
7	7	105i	No match	NA	No match	NA
8	8	105i	No match	NA	Not 105	105d
9	9	105i	Not 105		Not 105	160i
10	10	105i	3	105i	Not 105	92d
11	11	105i	5	105i	Not 105	105d
12	12	105i	6	105i	Not 105	105d
13	13	105i	7	105i	Not 105	105d
14	14	105i	8	105i	Not 105	31d
15	15	105i	9	105i	Not 105	105d
16	16	105i	11	105i	Not 105	105d
17	17	105i	Not 105	174i	Not 105	176i
18	18	105i	Q?		Not 105	105d
19	19	105i	Not 105	105d	Not 105	104i
20	20	105i	15		Not 105	105d
21	21	105i	12	NA	Not match	NA
22	22	105i	Not 105	ND	Not 105	105d
23	23	105i	16		Not 105	157i
24	24	105i	17	105d	Not 105	Q?
25	25	105i	Not 105	155i	Not 105	105d
26	26	105i	Not 105	239d	Not 105	103i
27	Not 105	106i	1	105i	Not 105	105d
28	No match		2	105i	Not 105	ND
29	Not 105	105d	4	105i	Not 105	114d
30	No match	NA	10	105i	Not 105	Q?
31	Not 105	105d	13	105i	No match	NA
32	Not 105	110d	14	105i	Not 105	114i
33	Not 105	Q?	18	105i	Not 105	ND
34	No match	NA	19	105i	Not 105	ND
35	Not 105	Q?	22	105i	Not 105	ND
36	Not 105	105d	Not 105	ND	1	105i
37	Not 105	ND	Not 105	ND	2	105i
38	Not 105	105d	Not 105	105d	3	105i
39	Not 105	Q?	Not 105	Q?	4	105i
40	Not 105	105d	Not 105	154i	5	105i
41	No match	NA	No match	NA	6	105i
42	No match	NA	No match	NA	7	105i
43	Not 105	105d	Not 105	105d	8	105i
44	Not 105	105d	Not 105	189i	10	105i
45	Not 105	103i	Not 105	105d	11	105i
46	Not 105	153i	Not 105	153i	12	105i
47	Not 105	ND	Not 105	ND	13	105i
48	Not 105	132i	Not 105	132i	14	105i
49	No match	NA	No match	NA	15	105i
50	No match	NA	No match	NA	16	105i
51	Not 105	222i	Not 105	222i	17	105i

^a “Locus” refers to each unique 105-bp locus as represented by unique flanking sequence. “105No” reflects each 105 copy in each genome as originally assigned in web Tables 8 to 10. “Not 105” means the sequence match between two or three genomes does not include the 105 bp element. “No match” means there is no significant match (>80% nucleotide identity in over 351 bp). In the “Indel” columns, the “d” and “i” following a number indicate a “deletion” or an “insertion,” respectively. Here “deletion” and “insertion” merely means the presence or absence of a specific length of sequence element. “Q?” means the indel identification was not clear from the sequence alignment. “ND” means the indel identification was not done. NA, not applicable.

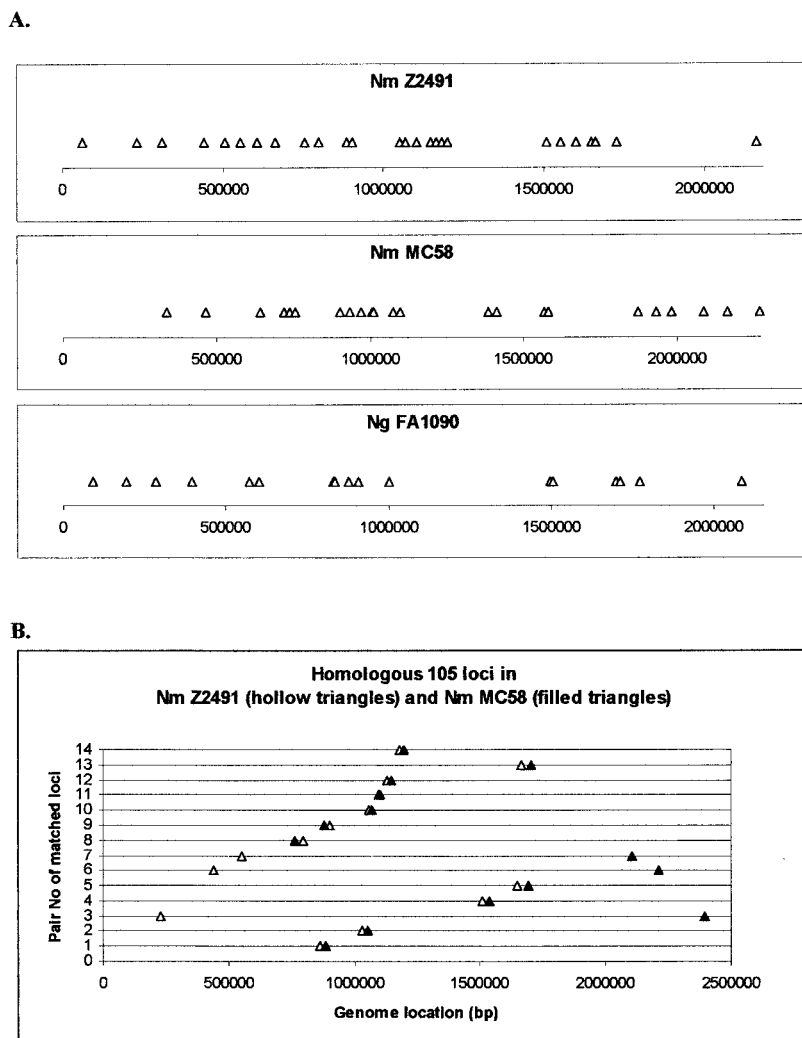


FIG. 5. Genomic locations of 105-bp indel and comparison between two *N. meningitidis* genomes. (A) 105-bp indels in each of the three neisserial genomes. (B) Fourteen homologous 105-bp indel loci in two *N. meningitidis* genomes when the two genomic sequences were aligned at the location of the 11th 105-bp indel.

TERNS analysis also gives more consistent and comparable data than the equivalent BLASTN analysis since it uses the same sequence pattern for identifying related sequences of varied lengths compared to using different queries in the BLASTN analysis. In fact, this FINDPATTERNS analysis is more powerful than the equivalent BLASTN analysis because it allows detection of greater diversity in the core regions and includes a wider range of lengths, i.e., more overall hits of more complex CREE. The accuracy of the FINDPATTERNS-based analysis was tested in a direct comparison with the BLASTN-based analysis of 107-bp CREE (105-bp indel). FINDPATTERNS analysis detected 26, 24, and 19 copies of 107-bp CREE, whereas BLASTN analysis retrieved only 26, 23, and 17 copies of 105-bp indels in *N. meningitidis* Z2491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090, respectively.

In comparing the number of CR and the number of CREE, it is apparent that CREE are indeed the most common form of CE. This is especially true for the two *N. meningitidis* genomes in which almost all identified CR are found in CREE, usually

as pairs of TIR. We made several interesting new observations in the present study. First, some longer CREE contain CREE of shorter lengths. Second, although the length profile of CREE shows several discrete clusters, there is often a continuous spreading of lengths within each cluster. Third, both the terminal regions of CREE and the core regions of CREE consist of distinctive sequence blocks. Fourth, the shortest CREE can be as short as 28-bp and comprise just the inverted CR.

Our analysis revealed that variations of CREE length reflect various combinations of several common sequence blocks (Fig. 2; see also web tables 2 to 7) and that CREE of the same length do not necessarily share the same sequence. In fact, many CREE lengths contain subtypes of sequences. These lengths include 159-, 156-, 155-, 106-, 96-, and 73-bp CREE in *N. meningitidis* Z2491 (web tables 2 and 5); 156-, 154-, 106-, and 105-bp CREE in *N. meningitidis* MC58 (web tables 3 and 6); and 155- and 105-bp CREE in *N. gonorrhoeae* FA1090 (web tables 4 and 7). The differences in the subtype sequences within

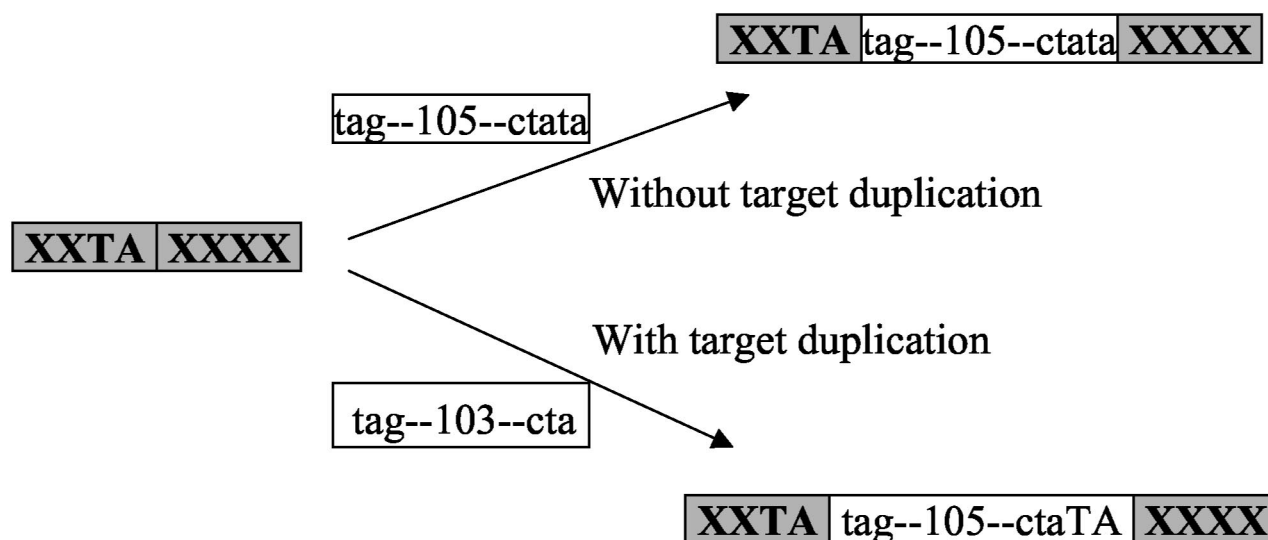


FIG. 6. Hypothetical mechanism of the formation of a 105-bp indel. Shaded boxes represent “target” sequences with uppercase sequence letters. Hollow boxes represent “insertion” sequences with lowercase sequence letters. X, unspecified target sequence.

these CREE are often due to the different combinations of the terminal inverted repeat sequence blocks and the core sequence blocks. However, the differences are sometimes also due to the presence of completely different core sequences. For example, the 341-bp CREE of *N. gonorrhoeae* FA1090 is different from the 341-bp CREE of the two *N. meningitidis* strains.

The CREE described in the present study do not contain any ORFs and cannot have any transposase activity. Thus, they are not insertion sequences by conventional definition (36). In this sense, we feel that it may be misleading to call CE (most of them are shown in the present study as CREE) as “*nemis* (*Neisseria* miniature insertion sequences)” (37). Instead, CREE may be better treated as indels, i.e., regions of DNA that are present on the chromosome of an organism but absent from closely related organisms (7). The experimental determination of DNA sequences upstream of sialyltransferase gene *lst* in multiple strains of *N. meningitidis* and *N. gonorrhoeae* revealed that a 107-bp CREE is actually a combination of a 105-bp indel with a TA dinucleotide in the flanking target sequence upstream of the 105-bp indel. Since all CREE have TA dinucleotides at their termini, it is reasonable to assume that all CREE are also made of target TA dinucleotides and indels of different lengths. This is supported by the observations of Abadi et al. that a 157-bp indel is located between the divergently transcribed *mtrR* and *mtrC* genes in *N. meningitidis* but not in the same region in *N. gonorrhoeae* (1). Upstream of this 157-bp indel in the *N. meningitidis* sequence is a TA dinucleotide. Thus, a 159-bp CREE is formed when this 157-bp indel combined with this flanking TA.

Currently, little is known about the origin or propagation of CREE. The discrete clustering of CREE around several lengths and the modular nature of CREE sequences suggest that the sequence diversity of CREE did not arise through sequential single base additions or deletions. Rather, different CREE might form via insertion or deletion of smaller sequence blocks plus some single-base-pair mutations. The ques-

tion remains whether all CREE are mobile or whether some CREE are simply “genetic fossils” of past DNA mobilization events. It also remains to be determined how frequent and by what mechanism this mobilization occurs. For example, the 105-bp indel found upstream of the *lst* gene of *N. meningitidis* strains might be formed through an insertion of a 103-bp or a 105-bp sequence element (Fig. 6). If the inserted sequence originally exists as a 103-bp sequence element, then a duplication event of the target TA sequence must occur to add another TA to the 3' end of the 103-bp element and form a 105-bp indel. The fact that all CREE contain a TA direct repeat at both ends supports the possibility of TA duplication. The observation of TA dinucleotides as the most common direct repeat found in the target sequence flanking CREE further suggests that repeated TA duplication may occur as multiple insertion events in the same target sites and that repeated insertion and target duplication events account for the spreading or diffusion of CREE length within each length cluster. Although target duplication during the insertion of noncoding short sequence elements has not been directly demonstrated in prokaryotes, such events have been described for short interspersed elements in eukaryotes (33). A recent study showed that a 107-bp interspersed repeat in *Streptococcus pneumoniae* could be mobilized via *trans*-mobilization by using the transposase of *IS630-Spn1* and, interestingly, a TA dinucleotide exists upstream of this 107-bp repeat (39). Thus, it is possible for the noncoding CREE to be passively mobilized and short sequences such as a TA dinucleotide in the insertion target to be duplicated in the insertion process.

The phenomenon of shorter repetitive sequences being enclosed within longer repetitive sequences is widely recognized. The REP/palindromic units (19, 25, 47) enclose some core sequences to form REP elements such as the IRU or ERIC (28, 45) and bacterial interspersed mosaic elements (BIME) (22). In *E. coli* K-12 there are 581 REP sequences and 314 REP elements (4). Therefore, most REP sequences are components of longer repeats. REP sequences may share similar

functions such as binding chromoid-associated protein (20), DNA gyrase (DNA topoisomerase) (51) and DNA polymerase I (21) and affecting mRNA stability (26). One subclass of BIME, called RIB (reiterative *ihf* BIME) (5) or RIP (repetitive IHF-binding palindromic elements) (40) is capable of binding integration host factor (IHF).

The relationship between CR and CREE resembles that between REP sequences and REP elements. The conservation of terminal regions in CR, and thus all CREE may reflect a highly conserved function for this DNA sequence fragment such as serving as a binding site for transposase. The 28-bp shortest CREE may represent the simplest indel structure on which a transposase can work. The diversification of core region may form the structural basis for the diverse function for various CREE. In this regard, there have been increasing reports of associating CREE with distinctive functions. For example, an earlier report demonstrated that a 106-bp CREE has promoter activity for the *uvrB* gene in *N. gonorrhoeae* (3). A study found that a 154-bp CREE appears to act as a transcriptional terminator (14). A recent study demonstrated that nemis (corresponding to some CREE described here) are cotranscribed with nearby cellular genes and subsequently processed at either one or both TIR (37). Since many CREE are located upstream of the coding regions of genes (web tables 1 and 12 to 14), the possibility for their involvement in a much wider range of gene regulation exists. For example, a 107-bp CREE is located near the *regF-regG* gene cluster (10), a 154-bp CREE is located downstream of the *mtrE* gene in *N. gonorrhoeae* (11), a 156-bp CREE exists in *N. gonorrhoeae* between divergently transcribed *frpB* and *groES* (48), a 152-bp CREE is located between *carA* and *carB* in both pathogenic and commensal *Neisseria* strains (34), and a 159-bp CREE is located between divergently transcribed *mtrR* and *mtrC* genes in *N. meningitidis* but not in the same region in *N. gonorrhoeae* (1). A total of 20 copies of CE were found in intergenic regions adjacent to different neisserial genes, including some virulence genes (14). However, the functional significance of these CREE remains to be experimentally determined.

Pathogenic *Neisseria* strains contain very "plastic" genomes (2, 6, 17) and are naturally competent for transformation (46), and they show evidence of extensive horizontal gene transfer (15, 32). The presence of a large number of interspersed DNA repeats in pathogenic genomes could affect the functional and evolutionary behaviors of these pathogens. The abundance of CREE (260, 255, and 98 nonredundant copies) and the percentage of nucleotides contained in these CREE (1.67, 1.55, and 0.59%) in the genomes of *N. meningitidis* Z1491, *N. meningitidis* MC58, and *N. gonorrhoeae* FA1090, respectively, are higher than that described for comparable intergenic repeats in other prokaryotic species. For example, the best-studied REP sequences and REP elements account for ca. 0.54% of the *E. coli* K-12 genome (4). The impact of these dispersed CREE in pathogenic neisserial genomes may be greater than currently realized because of their abundance and proximity to several virulence genes.

Pathogenic *Neisseria* strains present an interesting example of morphologically and biochemically similar organisms that cause very distinctive diseases, including life-threatening disseminated septicemia and meningitis caused by *N. meningitidis* and localized urogenital tract disease caused by *N. gonor-*

rhoeae. Analyses of the physical chromosomal maps of *N. meningitidis* (13, 16) and *N. gonorrhoeae* (12) show a high degree of conservation in overall gene organization between *N. meningitidis* Z2491 and *N. gonorrhoeae* FA1090 (13) and between *N. meningitidis* B1940 and *N. gonorrhoeae* FA1090 and MS11 (16). Previous DNA hybridization studies estimated that *N. meningitidis* and *N. gonorrhoeae* are 90% homologous in genes that are common to both species (24, 27). Whole genome sequence comparison showed that 91.2% of the 2,158 ORFs of *N. meningitidis* MC58 are similar to the ORFs of *N. meningitidis* Z2491 (49). There is no doubt that differences in the genes or ORFs (31, 42, 50) are important in determining the pathogenic differences between different *Neisseria* strains. However, the question is whether the distinctive pathogenic behaviors of different pathogens can be completely explained by these differences in the genes or ORFs. In this regard, the characterization of significant extragenic differences in one major family of neisserial repeat, CREE, among three strains of pathogenic *Neisseria* strains may offer some valuable additional insights.

A better understanding of CR and CREE may help the study of the evolutionary history and the phylogenetic classification of *Neisseria*. It is known that DNA loss via indel mutation is a determining factor for genome size reduction in eukaryotes (43). The relative genome sizes of three pathogenic *Neisseria* strains are 100% (*N. meningitidis* MC58), 96.1% (*N. meningitidis* Z2491), and 94.8% (*N. gonorrhoeae* FA1090). The contribution of CREE to the size variations of these three genomes is 0.08% between the two *N. meningitidis* genomes and ca. 1% between *N. gonorrhoeae* and *N. meningitidis* genomes. Thus, deletion or insertion of CREE alone can be a significant factor in altering the genome size of pathogenic *Neisseria* strains. CREE may serve as hot spots for genomic recombination and rearrangements, which may involve even larger segments of DNA and affect many different sets of genes, making CREE an important extragenic DNA component in the study of genome function.

ACKNOWLEDGMENTS

N. meningitidis Z2491 sequence data were determined by the *N. meningitidis* strain Z2491 Sequencing Group at the Sanger Centre (http://www.sanger.ac.uk/Projects/N_meningitidis/). *N. meningitidis* MC58 sequence data were determined by the *N. meningitidis* MC58 Sequencing Group at the Institute of Genome Research (ftp://ftp.tigr.org/pub/data/n_meningitidis/). We acknowledge the Gonococcal Genome Sequencing Project (supported by USPHS/NIH grant AI38399) and B. A. Roe, L. Song, S. P. Lin, X. Yuan, S. Clifton, Tom Ducey, Lisa Lewis, and D. W. Dyer at the University of Oklahoma. *N. gonorrhoeae* FA1090 sequence data can also be obtained online (<ftp://ftp.genome.ou.edu/pub/gono>). The GenBank accession number for the completed *N. gonorrhoeae* FA1090 genome is AE004969, and this link gives the GenBank sequence file containing the sequence as deposited in GenBank. We thank B.-J. Tang for writing computer codes for the custom programs.

This work was supported in part by grants AI33505 and AI20897 to R.F.R. from the National Institutes of Health. N.J.S. is supported by a Wellcome Trust Research Fellowship in Medical Microbiology.

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