

## *Bacteroides fragilis* Synthesizes a DNA Invertase Affecting both a Local and a Distant Region<sup>∇</sup>

Hazeline Roche-Hakansson, Maria Chatzidaki-Livanis, Michael J. Coyne, and Laurie E. Comstock\*

Channing Laboratory, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts

Received 28 August 2006/Accepted 8 December 2006

**The activity of a fourth conserved tyrosine site-specific recombinase (Tsr) of *Bacteroides fragilis* was characterized. Its gene, *tsr19*, is adjacent to *mpi*, encoding the global DNA invertase regulating capsular polysaccharide biosynthesis. Unlike the other described Tsrs of *B. fragilis*, Tsr19 brings about inversion of two DNA regions, one local and one located distantly.**

*Bacteroides fragilis* is a member of the intestinal microbiota and can become an opportunistic pathogen if released into extraintestinal sites. We have found that DNA inversions regulate expression of key virulence factors and may also modulate expression of surface factors involved in intestinal symbiosis. The *B. fragilis* NCTC 9343 genome encodes a large number of site-specific recombinases: three members of the serine site-specific recombinase (Ssr) family and 28 members of the tyrosine site-specific recombinase (Tsr) family. We have previously shown that two of the *ssr* genes and nine of the *tsr* genes are conserved in the species (3). We have demonstrated that Mpi is a global DNA invertase of the Ssr family that is involved in inverting 13 distinct DNA regions that are flanked by inverted repeats (IRs), including the promoters of seven of

the capsular polysaccharide biosynthesis regions (3). All of the regions that Mpi inverts are located at regions remote from *mpi*. We have also shown that three of the conserved *tsr* genes (*tsr15*, *tsr25*, and *tsr26*) encode DNA invertases that act locally to invert promoter regions immediately downstream of their gene (6).

Another conserved *tsr*, *tsr19*, is present on the *B. fragilis* genome adjacent to *mpi* and is transcribed divergently (Fig. 1A). The genomes of the two sequenced *B. fragilis* strains have been annotated such that the start codon of *mpi* is separated from the start codon of the divergently transcribed *tsr19* by only 15 bp (2, 4). The close clustering of these genes does not allow for the presence of intergenic promoter regions for either *mpi* or *tsr19*. Downstream of *tsr19* is a 315-bp region that

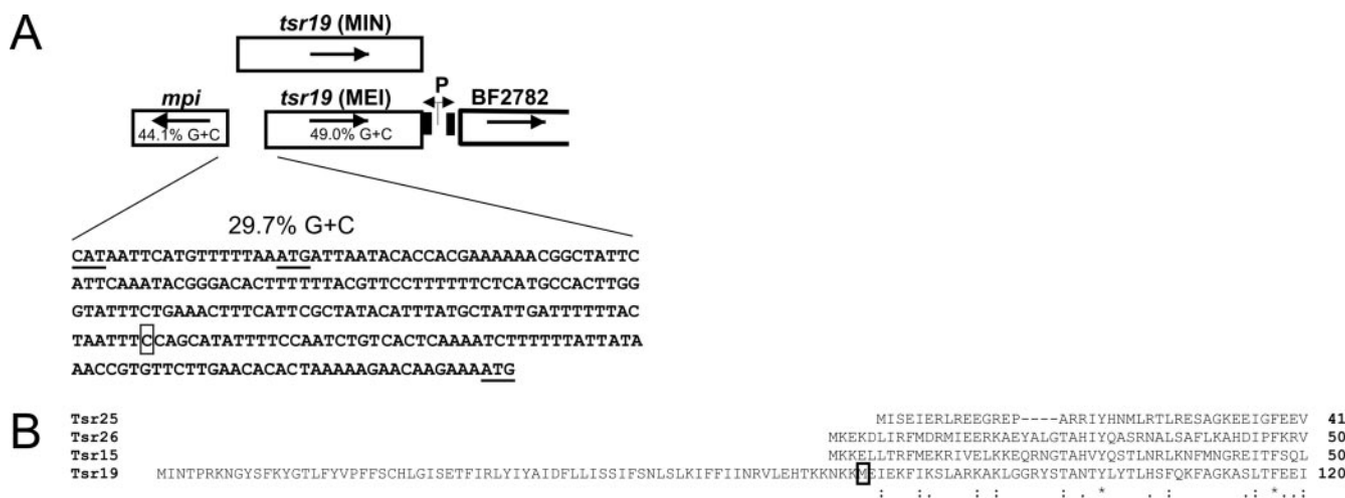


FIG. 1. Open reading frame map of the *mpi-tsr19* genetic region. A. *mpi* and *tsr19* are divergently transcribed. The predicted start of the *tsr19* gene based on the genome annotation (beginning with MIN) and an alternative predicted start site (beginning with MEI) are shown. Inverted repeat regions downstream of *tsr19* are indicated as black boxes with an intervening consensus promoter designated as P. The G+C content of the 234-bp intergenic region resulting from the MEI start of Tsr19 as well as the G+C contents of the adjacent genes are shown. The *mpi* start codon and the annotated and putative alternate *tsr19* start codons are underlined. The first base of the insert contained in pLEC84 is boxed. B. Alignment of Tsr19 with the three other functionally characterized tyrosine family site-specific recombinases of *B. fragilis*. A methionine representing an alternative putative start site for Tsr19 is boxed.

\* Corresponding author. Mailing address: Channing Laboratory, 181 Longwood Avenue, Boston, MA 02115. Phone: (617) 525-7822. Fax: (617) 264-5193. E-mail: lcomstock@channing.harvard.edu.

<sup>∇</sup> Published ahead of print on 22 December 2006.

TABLE 1. Primers used in this study

Purpose	Primer or plasmid	Primer sequence <sup>a</sup>
Creation of <i>tsr19</i> deletion mutants	<i>tsr19</i> flank 1 forward	TACCATGGTGCTTCAGAGAAGGTGACATACAT
	<i>tsr19</i> flank 1 reverse	TCGGATCCATTTGCCATAGTTCGTGAGTCTCT
	<i>tsr19</i> flank 2 forward	CTGGATCCACTTTCTGTTCCCTGAAACCATTC
	<i>tsr19</i> flank 2 reverse	ATCCATGGTATTGGCTGTGCTGTAACG
Demonstration of inversion of <i>tsr19</i> downstream region	<i>tsr19</i> forward	TGCTTCAGAGAAGGTGACATACAT
	<i>tsr19</i> central	ATTCATTTGTTTTATCGCTG
	<i>tsr19</i> reverse	ATACAATCCTTTTAATGACT
Cloning of <i>tsr19</i> for complementation (pLEC84)	<i>tsr19</i> forward	CTCGGATCCAGCATATTTTCCAATCTGTCACTC
	<i>tsr19</i> reverse	TCTGGATCCGCGAGCGATAAAACAAATGAATCT
Demonstration of inversion of PSA and PSH promoter regions	PSA forward	TGTGTAATGATAGGAGGCTAGGG
	PSA central	ACCTTTTTTCGACCTTTTCTAAAAATC
	PSA reverse	AGAAAACCTCCTGGTCTTCTTTG
	PSH forward	CTTTGCCAGTTCCCGTATGT
	PSH central	TGATGAAATTCAGAACCGGATA
	PSH reverse	CGCTCGTCTTGACGATGTA
Construction of <i>xyIE</i> transcriptional fusion clones in pLEC23	pHR20 and pHR41	ACGGATCCATTCATTTGTTTTATCGCTG ACGGATCCGCGGAATAGCGAAAAGATA
	pLEC55	TAGGATCCGTACTTACTCTCAAATAAGC TAGGATCCATAAACATAAAACAGGGGCTGG
	pMCL25	ACGGATCCATTCATTTGTTTTATCGCTG GTTGGGATCCCTTTTAGAAATGTTTGCGATAGTTG
	pMCL26	ACGGATCCGCGGAATAGCGAAAAGATA GTTGGGATCCCTTTTAGAAATGTTTGCGATAGTTG
	pMCL21	CGTTGGATCCGATAAAATCAGTCCTTTTGTTGC TCAGGATCCGCTAATGGTTTAATGTTTCATAGTTACTT
	pMCL22	ACGTGGATCCAATTTTAATGAATATATGGCTATGCA TCAGGATCCGCTAATGGTTTAATGTTTCATAGTTACTT
	pMCL23	CGTTGGATCCGATAAAATCAGTCCTTTTGTTGC GTGAGGATCCAATGTTAAGTATTTGTTGTTTCTTTAGTG
	pMCL24	ACGTGGATCCAATTTTAATGAATATATGGCTATGCA GTGAGGATCCAATGTTAAGTATTTGTTGTTTCTTTAGTG
	Demonstration of inversion of distant inverted repeat region	Forward
Central		GTTGCTGCAAATGTATGGATTCT
Reverse		TCAATGACACTTGTTCATGTTTG
PCR amplification of <i>groEL</i> probe for Northern blot analysis	<i>groEL</i> probe forward	GTGCAAGCCCTATGGATATTAAC
	<i>groEL</i> probe reverse	AATACCGGTAGTCTCGTCATCATT

<sup>a</sup> Restriction sites are underlined.

is flanked by inverted repeats and undergoes inversion (4). It has been suggested that a promoter within this invertible region may be involved in transcription of *mpi* (4). We sought to determine whether Tsr19 is a DNA invertase and, if so, whether it inverts this downstream IR region. We also sought to determine whether there is a functional promoter in this invertible region and, if so, whether it is necessary for the transcription of *mpi*.

Alignment of the three characterized Tsr family DNA invertases of *B. fragilis* with Tsr19 revealed that Tsr19 has a large N-terminal region of approximately 70 amino acids that the other three products lack (Fig. 1B). Furthermore, the second methionine of Tsr19 aligns within 4 amino acids of the initial methionines of the other three Tsrs. If it is assumed that the ATG corresponding to this second methionine of Tsr19 is the actual start codon, the percent G+C content of the resulting

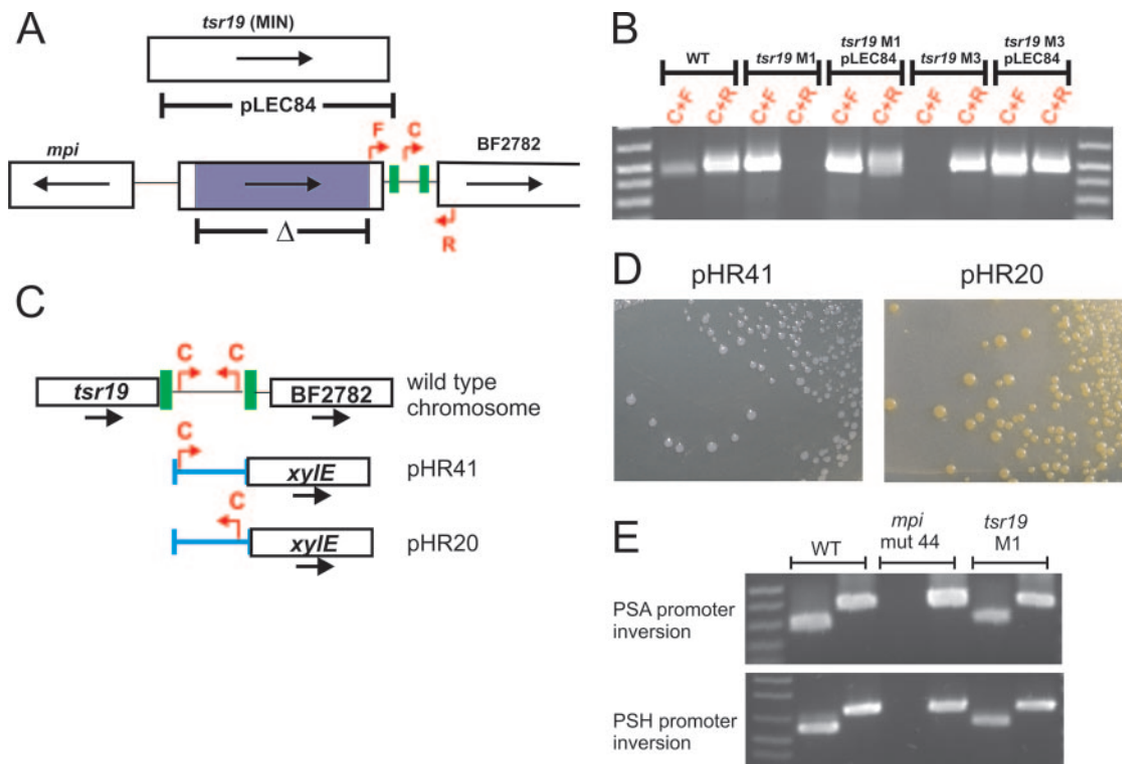


FIG. 2. Tsr19 is necessary for inversion of the downstream promoter region. A. Open reading frame map of the *mpi-tsr19* region showing the region that was deleted in the *tsr19* mutant (purple) and the extent of DNA used to complement this mutant (pLEC84). The red arrows show the primers used to assay for inversion. Inverted repeats are shown as green boxes. B. Ethidium bromide-stained agarose gel of the inversion event from the wild type (WT), *tsr19* mutants, and complemented mutants. Primers are those shown in panel A. C. Genetic regions (blue) cloned upstream of *xylE* in relation to the orientation of primer C, which undergoes inversion in wild-type bacteria and therefore is present in each orientation from a population of wild type bacteria. pHR20 and pHR41 contain identical DNA cloned in opposite orientations upstream of *xylE*. D. Analysis of functional promoter activity from *B. fragilis* 9343 containing pHR20 or pHR41 in *trans*. The transconjugants were sprayed with catechol, which is the substrate for XylE and convert the colonies to yellow if a functional promoter has been cloned. E. Ethidium bromide-stained agarose gels showing the inversion of the PSA and PSH promoter regions from the wild type, *mpi* deletion mutant 44, and  $\Delta$ *tsr19* M3.

234-bp intergenic region would be 29.7%, very dissimilar to the G+C contents of the surrounding genes (44.1% and 49.0% for *mpi* and *tsr19*, respectively). The low G+C content of this region suggests that it may be an intergenic region rather than part of the coding region of *tsr19*.

Based on the finding that Tsr15, Tsr25, and Tsr26 act on inverted repeat regions in their immediate downstream region, we predicted that Tsr19 has DNA invertase activity and mediates the inversion of its downstream IR region. Deletion mutants of *tsr19* where a 733-bp internal region was removed by allelic replacement were created (primers used in this study are listed in Table 1). The resulting mutants were tested for the inversion of the *tsr19* downstream region by using forward and central or central and reverse primer pairs. As shown in Fig. 2B, compared to that in the wild type, the DNA region between the IRs does not invert in the *tsr19* mutants. Mutants where this region was locked in each orientation were obtained (mutants M1 and M3), demonstrating that Tsr19 is necessary for the inversion of this region in both directions. For complementation studies, *tsr19* was cloned into the BamHI site of the *B. fragilis* expression vector pFD340 (5), creating pLEC84. Based on the possibility that *tsr19* may be shorter than predicted, pLEC84 was created so that the first 141 bp of the large gene was omitted (Fig. 2A), and this shorter clone was

tested for invertase activity. As shown in Fig. 2B, pLEC84 is able to mediate the inversion of this region in both  $\Delta$ *tsr19* M1 and  $\Delta$ *tsr19* M3. These data demonstrate that Tsr19 is a DNA invertase involved in the inversion of its downstream IR region and that the N terminus of the annotated gene is not necessary for its activity.

Like the other three characterized Tsr-regulated IR regions of *B. fragilis*, the region between the IRs of the *tsr19* downstream region has a perfect match with the described *B. fragilis* consensus promoter sequence (1). As it was predicted that *mpi* may be transcribed from this invertible promoter, it was necessary to determine if this region in fact contained an active promoter. A 284-bp DNA region between the IRs was PCR amplified and cloned in each orientation into the BamHI site of the *xylE* reporter plasmid pLEC23 (3) (Fig. 2C). The resulting clones, pHR20 and pHR41, were conjugally transferred into *B. fragilis*, and the presence of a promoter was qualitatively assessed by spraying the plates with the substrate catechol and by performing quantitative XylE assays as described previously (3). As shown in Fig. 2D, when this putative promoter is oriented to drive transcription of *xylE* (pHR20), 356  $\pm$  23 milliunits XylE/mg protein is produced, versus 0 milliunits XylE/mg protein when the promoter is oriented in the opposite direction in clone pHR41 (mean  $\pm$  standard deviation for two

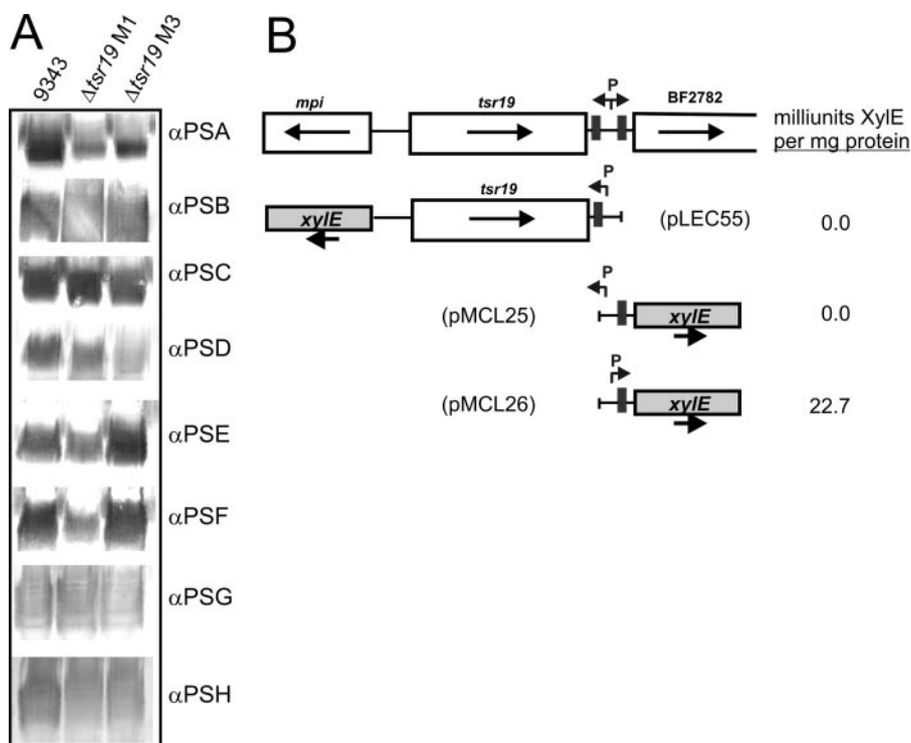


FIG. 3. The Tsr19-regulated invertible promoter does not affect capsular polysaccharide expression. A. Western immunoblot analysis of whole-cell lysates of the wild type,  $\Delta$ *tsr19* M1, and  $\Delta$ *tsr19* M3 probed with antisera to each of the eight capsular polysaccharides. B. Analysis of the transcription of *mpi* and BF2782 by the Tsr19-regulated invertible promoter, using *xylE* transcriptional fusion clones. pLEC55 detects transcription into *mpi* when the invertible promoter is oriented toward *mpi*, and pMCL25 and pMCL26 detect transcription into BF2782 when the invertible promoter is cloned in either the OFF or ON orientation, respectively. The resulting milliunits of XylE activity per milligram protein are reported for each of these constructs when in *trans* in *B. fragilis* 9343.

assays). Therefore, in the  $\Delta$ *tsr19* M3 mutant, the promoter is oriented so that it could constitutively transcribe *mpi* if this was indeed the promoter for *mpi*, whereas in the *tsr19* M1 mutant, the promoter is locked in the opposite orientation and thus could not drive transcription of *mpi*.

If the invertible promoter downstream of *tsr19* is necessary for the transcription of *mpi*,  $\Delta$ *tsr19* M1 should not express Mpi. To determine whether Mpi is synthesized in  $\Delta$ *tsr19* M1, we screened two polysaccharide promoter regions (PSA and PSH) known to invert in an Mpi-dependent manner (3). As a control, we used an *mpi* deletion mutant (mutant 44) which is unable to invert its polysaccharide promoters and has the PSA promoter locked ON and the PSH promoter locked OFF (3). As shown in Fig. 2E, inversion of these two polysaccharide promoter regions in  $\Delta$ *tsr19* M1 was similar to that in the wild type. Therefore, the promoter between the IRs of the *tsr19* downstream region is not necessary for expression of Mpi.

We next performed Western immunoblot analysis to determine if expression of any of the eight characterized capsular polysaccharides was affected by the deletion of *tsr19*. As shown in Fig. 3A, each of the eight capsular polysaccharides was synthesized in both  $\Delta$ *tsr19* M1 and  $\Delta$ *tsr19* M3, demonstrating that the invertible promoter downstream of *tsr19* does not affect capsular polysaccharide biosynthesis. To determine if this invertible promoter region may instead drive transcription of the downstream gene BF2782, *xylE* transcriptional fusion clones were constructed as shown in Fig. 3B. For both con-

structs, the upstream IR was omitted so that the promoter was locked either ON (pMCL26) or OFF (pMCL25) by PCR amplifying the region to be cloned from  $\Delta$ *tsr19* M1 or  $\Delta$ *tsr19* M3, respectively (Fig. 3B). No XylE activity was detected from pMCL25, in which the invertible promoter is oriented away from BF2782; however, when the promoter is oriented toward the gene (pMCL26), 22.7 milliunits XylE/mg protein is produced. Therefore, under the conditions of this assay, transcription of BF2782 is dependent on the invertible promoter in its upstream region.

Inverted repeats similar to those downstream of *tsr19* were detected in one other location of the *B. fragilis* genome at a distance of approximately 550 kb from *tsr19* (4). These IRs are separated by 261 bp of DNA that undergoes inversion (2, 4). To determine if Tsr19 is responsible for this inversion, PCR was performed using DNAs from the *tsr19* mutants and complemented mutants with forward and central or central and reverse primer pairs (Table 1; Fig. 4A). As shown in Fig. 4B, this distal region is unable to invert in *tsr19* mutants, and inversion is restored when *tsr19* is added in *trans* (pLEC84). In addition, in  $\Delta$ *tsr19* M1, this region is locked in the orientation opposite to that of  $\Delta$ *tsr19* M3.

This second Tsr19-regulated invertible region contains a perfect *B. fragilis* consensus promoter sequence which may be involved in the transcription of either of two divergently transcribed operons. The *groESL* operon is located 163 bp from the invertible region, and a divergently transcribed gene (BF3224)

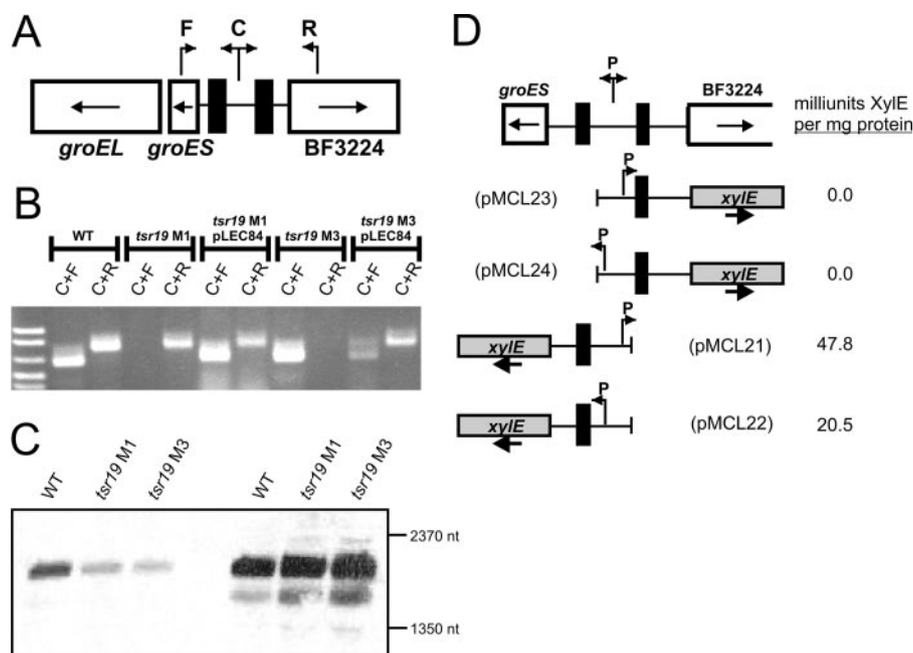


FIG. 4. Tsr19 is necessary for inversion of a distant chromosomal region. A. Schematic showing the genetic regions flanking the Tsr19-like inverted repeats (filled boxes). The direction of transcription of each gene is shown with arrows inside the boxes, and primers used to demonstrate inversion of DNA between the IRs are indicated as F, C, and R. B. Ethidium bromide-stained agarose gel of the resulting PCR products from the wild type (WT), *tsr19* mutants, and complemented mutants, using the F, C, and R primers shown in panel A. C. Northern blot analysis of wild type and *tsr19* mutants grown to early log phase (optical density at 600 nm = 0.3) (first three lanes) and mid-log phase (optical density at 600 nm = 0.8) (last three lanes), using 30  $\mu$ g of RNA per well probed with an internal portion of *groEL*. D. Analysis of the transcription of *groES* and BF3224 by the adjacent Tsr19-regulated invertible promoter, using *xylE* transcriptional fusion clones. pMCL23 and pMCL24 detect transcription into BF3224 when the invertible promoter is cloned in either the ON or OFF orientation, respectively. pMCL21 and pMCL22 detect transcription into *groES* when the invertible promoter is cloned in either the OFF or ON orientation, respectively. The resulting milliunits of XylE activity per milligram of protein are reported for each of these constructs when in *trans* in *B. fragilis* 9343.

is located 223 bp from the invertible region. The genetic organization and close proximity of the *groES* start codon to the invertible promoter suggested that transcription of the *groESL* operon might be regulated or enhanced by this invertible promoter. To determine if *groESL* transcription is dependent on this invertible promoter, Northern blot analysis was employed using RNA from the wild type,  $\Delta$ *tsr19* M1, and  $\Delta$ *tsr19* M3. Using an internal portion of *groEL* as a probe, we found that there were similar amounts of *groEL* transcript produced by  $\Delta$ *tsr19* M3 (where the promoter is locked OFF for *groESL* transcription) and  $\Delta$ *tsr19* M1 (where the promoter would constitutively drive transcription) from both early- and mid-log-phase cultures (Fig. 4C). These data demonstrate that this invertible promoter is not necessary for transcription of *groEL*. The hybridizing transcript size of  $\sim$ 2,000 nucleotides is the expected size for the *groESL* operon. To determine if this invertible promoter is involved in the transcription of the divergently transcribed gene BF3224, *xylE* transcriptional fusions similar to those of the other Tsr19-regulated invertible promoter region were constructed (Fig. 4C). No XylE activity was detected when this promoter was locked in either the OFF or ON orientation for transcription into BF3224. In addition, when the invertible promoter was locked to constitutively transcribe *groES*, no augmentation of XylE activity was detected compared to when this promoter was locked OFF for *groES* transcription. These data demonstrate that under the conditions in which the bacteria were grown, this invertible pro-

moter does not contribute to the transcription either of the *groESL* operon or of BF3224. The biological significance of this invertible promoter, therefore, is yet to be determined.

This study has demonstrated that a fourth conserved Tsr encoded by *B. fragilis* has DNA invertase activity. The three conserved Tsrs previously shown to have DNA invertase activity all localized to a single branch of a *B. fragilis* 9343 Tsr cladogram based on global protein alignment (3). Tsr19 also segregated to this same branch, allowing us to predict that this branch represents Tsr proteins with DNA invertase activity. Tsrs with other functions, such as transposition and integration, may segregate to other branches of this tree. Of the nine conserved Tsrs, five localize to this DNA invertase branch, and one other localizing to a distinct branch is predicted to be XerD (2), which is involved in chromosomal segregation. Therefore, few of the *tsr* genes that encode transposases or integrases are conserved in the species.

Tsr15, Tsr25, and Tsr26 each recognize almost identical inverted repeats; however, each is very specific in the inversion of its own downstream IR region (6). In contrast, Tsr19 recognizes two different, albeit related, IRs, one in its immediate downstream region, and one located distantly on the chromosome. This is the first report of a Tsr family DNA invertase of *B. fragilis* acting on a region distant from its gene, and Tsr19 shares this property with Mpi, the global Ssr of *B. fragilis*.

Studies designed to locate the *mpi* promoter by using the *xylE* promoter probing construct did not detect any promoter

activity immediately upstream of *mpi* (data not shown) or in regions extending beyond the invertible promoter downstream of *tsr19* (pLEC55) (Fig. 3B). Northern blot analysis also failed to detect an *mpi* transcript, suggesting that the gene is poorly transcribed from a very weak promoter. Due to this low level of *mpi* transcript, we used the functional assay of polysaccharide promoter inversion to assay for Mpi activity. The fact that Mpi activity was detected when the invertible promoter was locked in the OFF orientation for *mpi* transcription demonstrates that it is not necessary for its transcription.

We have shown that the invertible promoter downstream of *tsr19* is involved in transcription of at least one of the genes in its downstream region, BF2782. The products of the three genes of this region are similar to products involved in various aspects of extracellular polysaccharide biosynthesis and secretion. As the data have shown that this invertible promoter is not involved in regulation of capsular polysaccharide biosynthesis, our continuing studies will examine the effects of this promoter on the synthesis of an as-yet-uncharacterized extracellular polysaccharide.

This work was supported by grant AI44193 from the National Institutes of Health (NIAID).

#### REFERENCES

1. Bayley, D. P., E. R. Rocha, and C. J. Smith. 2000. Analysis of *cepA* and other *Bacteroides fragilis* genes reveals a unique promoter structure. *FEMS Microbiol. Lett.* **193**:149–154.
2. Cerdeno-Tarraga, A., S. Patrick, L. Crossman, G. Blakely, V. Abratt, N. Lennard, I. Poxton, B. Duerden, B. Harris, M. Quail, A. Barron, L. Clark, C. Corton, J. Doggett, M. Holden, N. Larke, A. Line, A. Lord, H. Norbertczak, D. Ormond, C. Price, E. Rabbinowitsch, J. Woodward, B. Barrell, and J. Parkhill. 2005. Extensive DNA inversions in the *B. fragilis* genome control variable gene expression. *Science* **307**:1463–1465.
3. Coyne, M. J., K. G. Weinacht, C. M. Krinos, and L. E. Comstock. 2003. Mpi recombinase globally modulates the surface architecture of a human commensal bacterium. *Proc. Natl. Acad. Sci. USA* **100**:10446–10451.
4. Kuwahara, T., A. Yamashita, H. Hirakawa, H. Nakayama, H. Toh, N. Okada, S. Kuhara, M. Hattori, T. Hayashi, and Y. Ohnishi. 2004. Genomic analysis of *Bacteroides fragilis* reveals extensive DNA inversions regulating cell surface adaptation. *Proc. Natl. Acad. Sci. USA* **101**:14919–14924.
5. Smith, C. J., M. B. Rogers, and M. L. McKee. 1992. Heterologous gene expression in *Bacteroides fragilis*. *Plasmid* **27**:141–154.
6. Weinacht, K. G., H. Roche, C. M. Krinos, M. J. Coyne, J. Parkhill, and L. E. Comstock. 2004. Tyrosine site-specific recombinases mediate DNA inversions affecting the expression of outer surface proteins of *Bacteroides fragilis*. *Mol. Microbiol.* **53**:1319–1330.