

## Molecular Evolution of the Pathogenicity Island of Enterotoxigenic *Bacteroides fragilis* Strains

AUGUSTO A. FRANCO,<sup>1,2</sup> RODNEY K. CHENG,<sup>1,2</sup> GYUNG-TAE CHUNG,<sup>3</sup> SHAOGUANG WU,<sup>1,2</sup> HEE-BOK OH,<sup>3</sup> AND CYNTHIA L. SEARS<sup>1,2\*</sup>

Divisions of Infectious Diseases<sup>1</sup> and Gastroenterology,<sup>2</sup> Department of Medicine, Johns Hopkins University, School of Medicine, Baltimore, Maryland 21205-2195, and Laboratory of Bacterial Toxins, Department of Microbiology, National Institute of Health, Seoul, Korea<sup>3</sup>

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Enterotoxigenic *Bacteroides fragilis* (ETBF) strains, which produce a 20-kDa zinc metalloprotease toxin (BFT), have been associated with diarrheal disease in animals and young children. Studying a collection of ETBF and nontoxigenic *B. fragilis* (NTBF) strains, we found that *bft* and a second metalloprotease gene (*mpII*) are contained in an ~6-kb pathogenicity island (termed *B. fragilis* pathogenicity island or BfPAI) which is present exclusively in all 113 ETBF strains tested (pattern I). Of 191 NTBF strains, 100 (52%) lack both the BfPAI and at least a 12-kb region flanking BfPAI (pattern II), and 82 of 191 NTBF strains (43%) lack the BfPAI but contain the flanking region (pattern III). The nucleotide sequence flanking the left end of the BfPAI revealed a region with the same organization as the mobilization region of the 5-nitroimidazole resistance plasmid pIP417 and the clindamycin resistance plasmid pBFTM10, that is, two mobilization genes (*bfmA* and *bfmB*) organized in one operon and a putative origin of transfer (*oriT*) located in a small, compact region. The region flanking the right end of the BfPAI contains a gene (*bfmC*) whose predicted protein shares significant identity to the TraD mobilization proteins encoded by plasmids F and R100 from *Escherichia coli*. Nucleotide sequence analysis of one NTBF pattern III strain (strain I-1345) revealed that *bfmB* and *bfmC* are adjacent to each other and separated by a 16-bp GC-rich sequence. Comparison of this sequence with the appropriate sequence of ETBF strain 86-5443-2-2 showed that in this ETBF strain the 16-bp sequence is replaced by the BfPAI. This result defined the BfPAI as being 6,036 bp in length and its precise integration site as being between the *bfmB* and *bfmC* stop codons. The G+C content of the BfPAI (35%) and the flanking DNA (47 to 50%) differ greatly from that reported for the *B. fragilis* chromosome (42%), suggesting that the BfPAI and its flanking region are two distinct genetic elements originating from very different organisms. ETBF strains may have evolved by horizontal transfer of these two genetic elements into a pattern II NTBF strain.

Enterotoxigenic *Bacteroides fragilis* (ETBF) strains have been associated with watery diarrheal disease in livestock and young children (27, 28, 34, 35, 38, 40). The primary virulence factor proposed for these strains is a zinc-dependent metalloprotease toxin (*B. fragilis* toxin [BFT]) (23). This toxin stimulates fluid accumulation in lamb ligated ileal loops (LLIL) and alters the morphology of human intestinal epithelial cells in vitro, especially HT29/C1 cells (3, 17, 27, 30, 40, 49).

We recently reported cloning and sequencing the *B. fragilis* toxin gene (*bft*) from a cosmid library of ETBF strain 86-5443-2-2 (piglet isolate) (9) and, at the same time, Kling et al. reported sequencing the *bft* gene from ETBF strain VPI 13784 (lamb isolate) (16). The *bft* gene from both strains encodes a preprotoxin of 44 kDa that is processed to yield a biologically active toxin of ca. 20 kDa, which is secreted into the culture supernatant. By comparing the reported *bft* sequences from ETBF strains 86-5443-2-2 and VPI 13784, we developed specific oligonucleotide probes derived from an area of reduced homology in the sequence of these *bft* genes. Hybridization of these oligonucleotide probes to ETBF strains 86-5443-2-2 and VPI 13784 revealed that each gene could be independently identified (9). Furthermore, examination of a collection of ETBF strains revealed that ETBF strains contain one of these

*bft* genes but not both (9). We designated these alleles *bft-1* and *bft-2* representing the genes derived from strains VPI 13784 and 86-5443-2-2, respectively. In additional studies (50), we found that (i) BFT purified from culture supernatant of ETBF strain 86-5443-2-2 (BFT-2) had modest but consistently greater biological activity when tested on HT29/C1 cells than BFT purified from ETBF VPI 13784 (BFT-1) and that (ii) BFT-1 and BFT-2 were eluted with different concentrations of NaCl from a high-resolution anion-exchange column (MonoQ) and exhibit different electrophoretic mobilities on sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) gels. These results indicate that ETBF VPI 13784 and 86-5443-2-2 produce two distinct isotypes of BFT.

In studying a collection of ETBF and nontoxigenic *B. fragilis* (NTBF) strains, we also found that *bft* is located in an ~6-kb region present only in ETBF strains (7). Later, sequence analysis of this 6-kb region revealed that, in addition to *bft*, this region contains another metalloprotease distinct from *bft*. Together, these data indicate that *bft* is contained in a pathogenicity island (8). Moncrief et al. recently reported the sequence of this 6-kb region (termed a pathogenicity islet by the authors) from strain VPI 13784 (24). In the present study we analyzed the DNA sequence of this specific 6-kb region and its flanking region from ETBF 86-5443-2-2 and a large collection of ETBF and NTBF strains. Our results suggest that the specific 6-kb region and its flanking region form different mobile genetic elements and that ETBF strains may have evolved by acquisition of these two genetic elements.

\* Corresponding author. Mailing address: Divisions of Infectious Diseases and Gastroenterology, Johns Hopkins University School of Medicine, Ross Bldg., Rm. 933, 720 Rutland Ave., Baltimore, MD 21205. Phone: (410) 955-9680. Fax: (410) 955-9677. E-mail: csears@welchlink.welch.jhu.edu.

TABLE 1. *B. fragilis* strains used in this study

Strain	Origin	Characteristic <sup>a</sup>	Source or reference
86-5443-2-2	Pig	ETBF (intestinal)	L. L. Myers
VPI 13784	Lamb	ETBF (intestinal)	T. D. Wilkins
J38-1	Human	ETBF (intestinal)	26, 49
20793-3	Human	ETBF (intestinal)	26, 49
DS-64	Human	ETBF (intestinal)	26, 49
DS-233	Human	ETBF (intestinal)	26, 49
DS-49	Human	ETBF (intestinal)	26, 49
I-1356-1	Calf	ETBF (intestinal)	L. L. Myers
TM4000	Human	NTBF (abscess)	M. H. Malamy
077225-2	Human	NTBF (intestinal)	L. L. Myers
I-1345	Calf	NTBF (intestinal)	L. L. Myers
LM-20	Human	NTBF (blood)	26
LM-12	Human	NTBF (blood)	26
K518	Human	NTBF (abscess)	G.-T. Chung

<sup>a</sup> ETBF indicates that culture supernatants of the strains stimulate morphologic changes in the HT29/C1 cells, whereas culture supernatants of NTBF strains do not.

## MATERIALS AND METHODS

**Bacterial strains, vectors, and growth conditions.** ETBF 86-5443-2-2 (piglet isolate) was used for the identification and characterization of the *B. fragilis* pathogenicity island (BfPAI). *B. fragilis* strains used for sequencing, determination of integration site, and the mobilization of BfPAI are listed in Table 1. For colony blot hybridizations, a total of 113 ETBF and 191 NTBF strains were tested. Of 75 intestinal ETBF strains, 39 were from the United States (human), 22 were from Bangladesh (human), 4 were from Thailand (human), 7 were from the United States (calf), and 3 were from the United States (lamb). Of 38 extraintestinal ETBF strains, 4 were isolated in the United States (1 from blood, 1 from pleural fluid, and 2 from wound), and 34 were isolated in Korea (12 from blood, 6 from wound, 4 from pus, 3 from peritoneal fluid, 2 from bile, and 7 from other body fluids). Of 75 intestinal NTBF strains, 6 were from the United States (human), 25 were from Bangladesh (human), 38 were from Thailand (human), 5 were from the United States (calf), and 1 was from the United States (lamb). Of the 116 extraintestinal NTBF strains, 60 were from the United States (27 from blood, 6 from pleural fluid, 6 from abscess, 6 from bile, 11 from wound, and 4 from other body fluids), and 56 were isolated from Korea (11 from blood, 18 from wound, 11 from pus, 4 from peritoneal fluid, and 12 from other body fluids). Isolates were identified as ETBF or NTBF on the basis of the activity on HT29/C1 cells and/or the LLLI assay.

Human intestinal and extraintestinal *B. fragilis* strains from the United States were gifts from R. B. Sack, L. L. Myers, or the microbiology laboratory of the Johns Hopkins Hospital, as previously reported (26, 49). Strains from Korea, Bangladesh, and Thailand and the animal (calf and lamb) *B. fragilis* strains were gifts from G.-T. Chung, R. B. Sack, P. Escheverria, and L. L. Myers, respectively.

*Bacteroides thetaioaomicron* BT4107 and BT400, used as bacterial mating controls, were gifts from N. B. Shoemaker and A. A. Salyers. *Escherichia coli* HB101 and DH5 $\alpha$  were used as hosts for maintaining cosmid and plasmid clones, respectively. The cosmid vector pHC79 was used to construct the *B. fragilis* 86-5443-2-2 genomic library, whereas pBluescript II SK(+) (Stratagene, Inc., La Jolla, Calif.) was used for subcloning cosmid DNA. *B. fragilis* strains were propagated anaerobically on BHC medium (3.7 g of brain heart infusion base [Difco Laboratories] per liter and 0.1 mg of vitamin K per liter, 0.5 mg of hemin per liter, 50 mg of L-cysteine per liter [all from Sigma, St. Louis, Mo.]), and *E. coli* strains were grown on Luria agar or in Luria broth. The following antibiotic concentrations were used unless otherwise noted: ampicillin, 200  $\mu$ g/ml; tetracycline, 3  $\mu$ g/ml; rifampin, 20  $\mu$ g/ml; and trimethoprim, 100  $\mu$ g/ml.

**DNA isolation analysis.** Plasmid DNA was extracted by the alkali lysis method (37) or by using Qiagen columns (Qiagen, Inc.). The CsCl-ethidium bromide density gradient centrifugation technique was used to identify plasmids in ETBF strain 86-5443-2-2. Cosmids 5G11 and 11F11 containing *bft-2* were restriction mapped by Southern blot walking according to the conditions described previously (9). Purification of DNA fragments and extraction from gel slices were performed with Qiaex II gel extraction kit (Qiagen, Inc.).

**Colony blot hybridizations.** Colony blots of *B. fragilis* strains were prepared by the technique described previously (9). Briefly, *B. fragilis* organisms grown overnight on BHC agar were transferred to Whatman 541 filters. The filters were microwave processed in alkali solution (0.5 M NaOH, 1.5 M NaCl) followed by neutralization in 2 M ammonium acetate. The filters were hybridized with either fragment probes or specific oligoprobes. The fragment probes were labeled with [ $\alpha$ -<sup>32</sup>P]dATP by random priming (Random-Prime DNA-Labeling Kit; Boehringer GmbH, Mannheim, Germany), hybridized in 50% formamide-5 $\times$  SSC (1 $\times$  SSC is 0.15 M NaCl plus 0.015 M sodium citrate)-0.1% SDS-1 mM EDTA-1 $\times$  Denhardt's solution at 37°C, and washed with 5 $\times$  SSC-0.1% SDS at 65°C.

TABLE 2. Primers and oligonucleotide probes used in this study

Primer or oligoprobe	Sequence (5' to 3')
<b>Primers</b>	
P1T7	.....GCTGGTAGACTACCTGAGTAAGGAGTC
P1T7-1	.....GCTTCGGTACCCAGGTATCTCTCCATA
P1T3	.....TTCAACCTGATCGATCCGGAAGATCCG
P1T3-1	.....GGTAGTGCTTATGTCCTCGCAACCTCA
PrimA	.....GCATGGATCCATGTGCTGACGATCTTCTT
PrimB.1	.....CGCGCTCGAGTTTTGTATGCATTCACAC
Confir1	.....CAGCTTCATTAGAACATGCAGCTAATAA
Confir2	.....CTGTCCATAACTGTATTCCATTTGATAC
Confir3	.....CACCGACAGAACATATCCACTCAACTG
Confir4	.....GGTGTGCAAGAGAGGCTCAAACACAT
OliртеB1	.....GACTACGCCGAGTTAATACCG
OliртеB2	.....TCGTGCAGGCGGTACAGAAGAT
<b>Oligoprobes</b>	
MT086	.....GGTGCTAGGCATGCGGATGATC
MTVPI	.....GGCGCTGAGCATACGGATAATT
K469	.....CAGATGCAGGATGCGGCGAACT
Olibf3	.....CGAGGTGTCTGCTCTTGAATC

Synthetic oligonucleotide probes were end labeled with [ $\gamma$ -<sup>32</sup>P]dATP by using T4 polynucleotide kinase (37), hybridized in 6 $\times$  SSC-5 $\times$  Denhardt's solution-1 mM EDTA (pH 8) for 16 h at 56°C and washed twice in 1 $\times$  SSC-1% SDS at 56°C for 10 min.

**Nucleotide sequence analysis.** Recombinant plasmids and PCR products were sequenced by the fluorescent dideoxy terminator method of cycle sequencing (20) on a PE/ABd 373a automated DNA sequencer at the DNA Analysis Facility of Johns Hopkins University. Sequences were compiled with Sequencer software (Gene Codes Corp., Ann Arbor, Mich.). DNA and amino acid sequences were analyzed by using programs developed by the Genetic Computer Group at the University of Wisconsin (4), and a sequence homology search was performed by using the National Center for Biotechnology Information (NCBI) BLAST server (1).

**Expression of MPII in *E. coli*.** The T7 promoter polymerase expression system (Novagen pET-26b[+]) was used to express metalloprotease MPII in *E. coli*. Primers PrimA and PrimB.1 (sequences are described in Table 2) generated a PCR product of the DNA region encoding the proprotein and mature protein of MPII and containing a *Bam*HI and *Xho*I at the 5' and 3' ends of the product, respectively. Vector pET-26b(+) carries an N-terminal *pelB* signal sequence for potential periplasmic location. Cloning of the PCR product in the *Bam*HI and *Xho*I sites of the pET-26b(+) vector generated a fusion protein containing *pelB* signal sequence-MPII proprotein-mature protein, whose expression was under the control of the T7 promoter. As recommended by the manufacturer, the *mpII* construction was first cloned into a nonexpression host (*E. coli* HB101) lacking the T7 RNA polymerase. Once established in a nonexpression host, plasmids were transferred into the expression host (*E. coli* BL21) containing a copy of the T7 RNA polymerase gene, which permitted expression of MPII. Expression of MPII in *E. coli* was determined by Western blot with anti-ETBF 86-5443-2-2 antibodies.

**Activity of MPII on HT29/C1 cells and E-cadherin.** The cell-free supernatant and the supernatants of whole-cell lysate of strain F2 (*E. coli* expressing MPII) were tested for biological activity on HT29/C1 cells and E-cadherin. Activity on HT29/C1 cells was determined at 3 or 24 h as described previously (26, 49). Activity of MPII on E-cadherin was determined as described previously (51). Briefly, HT29/C1 cells were treated with sterile MPII-containing filtrates for 16 h at 37°C. Cleavage of E-cadherin was determined by Western blot by using antibodies to the cytoplasmic domain of E-cadherin (a gift of James Nelson, Stanford University, Palo Alto, Calif.) as the primary antibody (1:1,000 dilution) and anti-rabbit immunoglobulin G-horseradish peroxidase as the secondary antibody (1:10,000 dilution). Purified BFT from ETBF strain 86-5443-2-2 was used as a positive control in both assays. Negative controls included medium alone and crude supernatant filtrates of the NTBF strain 077225-2 and *E. coli* BL21 containing plasmid pET26.

**Bacterial matings.** Thymidine-dependent mutant strains 086Thy<sup>-3</sup>, I-1356-1Thy<sup>-1</sup>, I-1345Thy<sup>-2</sup>, and K518Thy<sup>-2</sup> derived from ETBF 86-5443-2-2 (contains *bft-2*), ETBF I-1356-1 (contains *bft-1*), NTBF I-1345 (pattern III; see Results), and NTBF K518 (pattern II; see Results), respectively, were used as donor strains. All donor strains were resistant to tetracycline (Tc<sup>r</sup>) and sensitive to rifampin (Rif<sup>r</sup>). The presence of the tetracycline-inducible transposon (Tc<sup>r</sup> element) in each Tc<sup>r</sup> strain was confirmed by PCR by using a set of primers derived from the regulatory gene *rtxB* (OliртеB1 and OliртеB2 [sequences in Table 2]) (46). NTBF TM4000 (Tc<sup>r</sup> Rif<sup>r</sup>) was used as the recipient strain. *B. thetaioaomicron* BT4107 (a thymidine-dependent donor containing the Tc<sup>r</sup>Em<sup>r</sup>-DOT element) and *B. thetaioaomicron* BT4001 (a rifampin-resistant recipient)

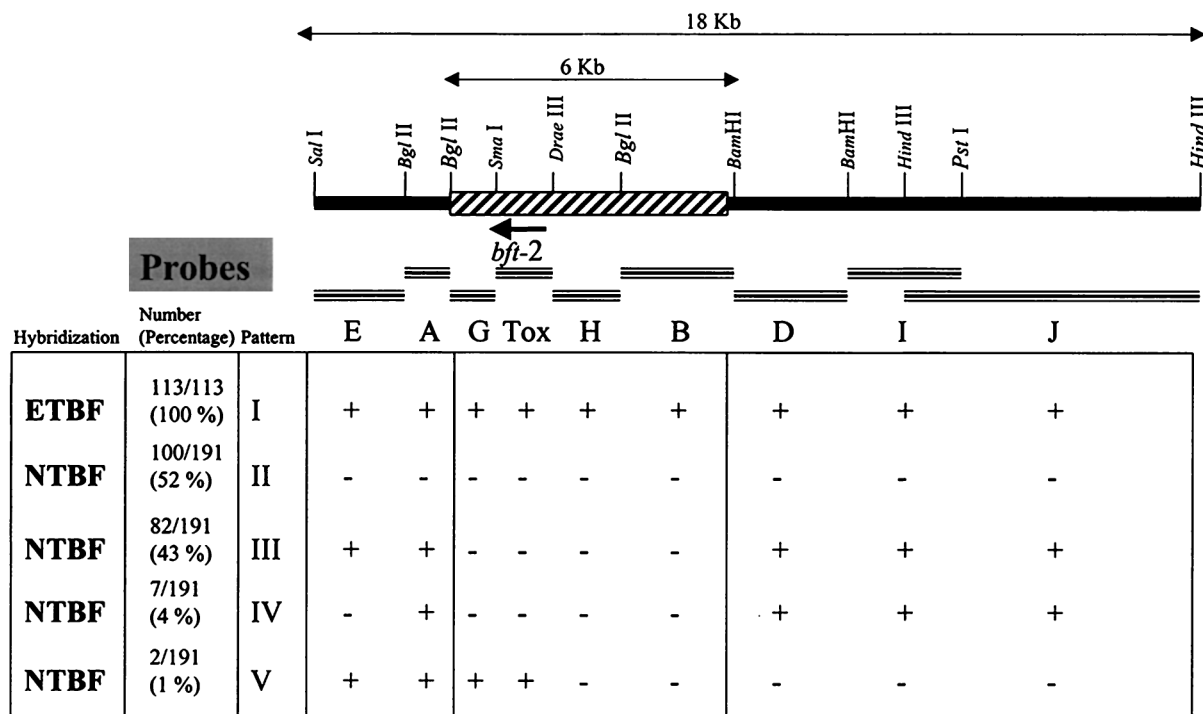


FIG. 1. Partial restriction map of the region flanking *bft* in ETBF 86-5443-2-2. Arrow shows *bft* and the direction of its transcription. Probes used in the colony blot hybridizations are shown in the black bars. The results of hybridization of these probes to *B. fragilis* strains (Table 2) are indicated beneath each probe. The 6-kb region spanning probes G, Tox, H, and B (the striped bar) is the specific locus present only in ETBF strains (BfPAI). Only the restriction sites used for the construction of the probes are shown in the figure.

strains were used as bacterial mating controls. Conjugation matings were done by using the filter mating procedure as described by Shoemaker et al. (42). Briefly, exponential cultures of donor and recipient bacteria were centrifuged, washed, and resuspended in fresh medium. Donor and recipient cells were mixed in a final ratio of 2:1, and 20 µl of this mix was applied to nitrocellulose filters placed on BHC medium containing thymidine (100 µg/ml). The filter matings were incubated overnight at 35°C, and transconjugants were selected on BHC medium containing rifampin (20 µg/ml) and tetracycline (3 µg/ml) and lacking thymidine to avoid the growth of donor Rif<sup>r</sup> spontaneous mutants. Transconjugants were confirmed to contain the Tc<sup>r</sup> element by PCR by using primers OIriteB1 and OIriteB2 (Table 2). Donor strains were grown in medium containing 1 µg of tetracycline per ml before mating to induce factors in the Tc<sup>r</sup> element. The frequency of transfer of the Tc<sup>r</sup> element was determined by dividing the number of transconjugants by the number of recipients.

**PCR conditions.** PCR primers used in this study are listed in Table 2. PCRs were performed with *Taq* polymerase (1.5 U) in 50 µl containing either chromosomal DNA (20 to 50 ng) or plasmid (5 to 10 ng), primers (25 pmol), deoxynucleoside triphosphates (200 µM), and MgCl<sub>2</sub> (1.5 M). Amplification used a hot start (94°C for 5 min), followed by denaturation at 94°C for 1 min, annealing at 66°C for 2 min, and extension at 72°C for 1 min. The amplification cycle was repeated either 29 times for determination of the integration site of BfPAI or 19 times for generating *mpII* for the cloning experiments and was followed by a final extension at 72°C for 5 min. The amplified products were directly sequenced or cloned into the TA cloning vector (Invitrogen, San Diego, Calif.).

**Data analysis.** Data were analyzed by the chi-square formula; a *P* value of <0.05 was considered statistically significant.

**Nucleotide sequence accession numbers.** We originally submitted the *mpII* gene sequence from ETBF strain 86-5443-2-2 to GenBank as *bft3* (GenBank accession number AF056297). However, we adopt here the nomenclature *mpII* in accordance with that used by Moncrief et al. (24). The *bfmA*, *bfmB*, *bfmC*, and open reading frame (ORF) ORF-A genes were assigned accession numbers AF118241, AF118242, AF118243, and AF118244, respectively.

**RESULTS**

**The region flanking the *bft* gene is present only in ETBF strains.** To determine whether the DNA region flanking *bft-1*/*bft-2* was specific for ETBF, two overlapping cosmids (11F11 and 10G11) containing *bft-2* (from ETBF strain 86-5443-2-2)

were mapped with several restriction enzymes. Eight adjacent restriction fragments were used to probe colony blots of 113 ETBF and 191 NTBF strains. Five patterns of hybridization were identified (Fig. 1). All of the probes spanning an 18-kb sequence hybridized to the 113 ETBF strains (pattern I). Of 191 NTBF strains, 100 (52%) lacked this 18-kb chromosomal region (pattern II), and 82 (43%) lacked only an ~6-kb region (containing probes G, Tox, H, and B) flanking *bft-2* (pattern III). Of the 191 NTBF strains, 7 (3.6%) lacked the region spanning probe E in addition to the 6-kb region flanking *bft* (pattern IV). Two NTBF strains containing the *bft* gene (i.e., hybridized to the Tox probe) did not hybridize to probes J, I, D, B, and H in the region spanning ca. 11.5 kb upstream of *bft-2* (pattern V). These two strains hybridized with an oligo-probe specific for the DNA region downstream of the metalloprotease motif of the *bft-2* allele (oligoprobe MT086), but neither of them hybridized to an oligoprobe (oligoprobe K469) derived from a common region of *bft-1* and *bft-2* that encodes the N-terminal region of the mature protein, suggesting that these two NTBF strains contain a deletion in the DNA encod-

TABLE 3. Frequency of patterns of hybridization in *B. fragilis* strains isolated from intestinal and extraintestinal sources

Pattern	No. of strains/total (%)	
	Intestinal	Extraintestinal
I	75/113 (66)	38/113 (33.6)
II	35/100 (35)	65/100 (65)
III	40/82 (48.7)	42/82 (51)
IV	0/7	7/7 (100)
V	0/2	2/2 (100)

TABLE 4. Frequency of patterns of hybridization in intestinal *B. fragilis* strains isolated from The United States, Bangladesh, and Thailand and in extraintestinal *B. fragilis* strains isolated from The United States and Korea

Source and pattern	No. of strains/total (%), type
<b>Intestinal (United States)</b>	
I .....	49/49 (100), ETBF
III .....	12/12 (100), NTBF
<b>Extraintestinal (United States)</b>	
I .....	4/4 (100), ETBF
II .....	36/60 (60), NTBF
III .....	15/60 (25), NTBF
IV .....	7/60 (11), NTBF
V .....	2/60 (3), NTBF
<b>Intestinal (Bangladesh)</b>	
I .....	22/22 (100), ETBF
II .....	11/25 (44), NTBF
III .....	14/25 (56), NTBF
<b>Intestinal (Thailand)</b>	
I .....	4/4 (100), ETBF
II .....	24/38 (63), NTBF
III .....	14/38 (37), NTBF
<b>Extraintestinal (Korea)</b>	
I .....	34/34 (100), ETBF
II .....	29/56 (52), NTBF
III .....	27/56 (48), NTBF

ing the amino terminus of the mature BFT protein, as well as the region upstream of the gene.

Overall, pattern I ETBF strains were isolated at a significantly higher frequency from intestinal (75 of 150 [50%]) than extraintestinal (38 of 154 [25%]) samples ( $P < 0.001$ ) (Table 3). All *B. fragilis* strains with hybridization patterns IV and V were isolated from extraintestinal sources (Table 3). In the United States and Thailand, only 6% (4 of 64) of extraintestinal *B. fragilis* strains or 10% (4 of 42) of intestinal *B. fragilis* strains, respectively, are ETBF strains (pattern I). Interestingly, in these locations the NTBF strains with pattern II were identified with higher frequency than the NTBF strains with

pattern III (Table 4). In contrast, in intestinal *B. fragilis* strains from the United States and Bangladesh, as well as extraintestinal *B. fragilis* strains from Korea, places where ETBF strains have been detected with higher frequency (3a, 34, 35), either NTBF strains with pattern II were not detected (intestinal, United States) or a similar proportion of NTBF strains with patterns II and III were identified (Table 4). Because no significant differences in the proportion of pattern II versus pattern III NTBF strains were found when areas with lower (extraintestinal, United States; intestinal, Thailand) and higher (extraintestinal, Korea; intestinal, Bangladesh) numbers of ETBF strains were compared, the groups were combined and analyzed. This comparison revealed that pattern II NTBF strains were identified in a significantly higher proportion from areas with lower recovery of ETBF isolates ( $P < 0.02$ ).

**Nucleotide sequence of the specific 6-kb region.** The colony blot hybridization results indicate that an ~6-kb region flanking *bft* is present exclusively in all ETBF strains ( $n = 113$ ). To examine this region for the presence of additional potential virulence genes, in addition to the *bft-2* sequence already reported (9), we determined the sequence of 3,045 bp downstream of *bft-2* and 5,799 bp upstream of *bft-2* (total sequenced region, 10,038 bp) (Fig. 2a). Analysis of the specific 6-kb region revealed that, in addition to *bft-2*, this region contains two other ORFs (ORF-A and ORF-B), which are 375 and 1,180 bp in length, respectively. ORF-A is transcribed in the opposite direction from *bft-2*, and its stop codon is located 270 bp downstream of stop codon of *bft-2* (Fig. 2a). The predicted amino acid sequence of the protein encoded by ORF-A showed no significant similarity to any sequence in the database. ORF-B also transcribes in the opposite direction of *bft-2*, with the start codon located 1.6 kb upstream of the start codon of the *bft-2* gene (Fig. 2a). The predicted protein encoded by ORF-B is 44.36 kDa with a predicted lipoprotein signal peptide (18 amino acids) and a zinc-binding metalloprotease motif similar to BFT-1 and BFT-2. However, this predicted protein is only 29 and 28% identical to BFT-1 and BFT-2, respectively, and contains an ATP/GTP-binding site motif A (48) not found in BFT-1 or BFT-2. The predicted protein encoded by ORF-B was termed metalloprotease II (MP-II) in accordance with the results of Moncrief et al. (24).

To correlate the presence of *bft-1/bft-2* and *mpII*, a specific oligoprobe derived from *mpII* sequence (Olibft3) was de-

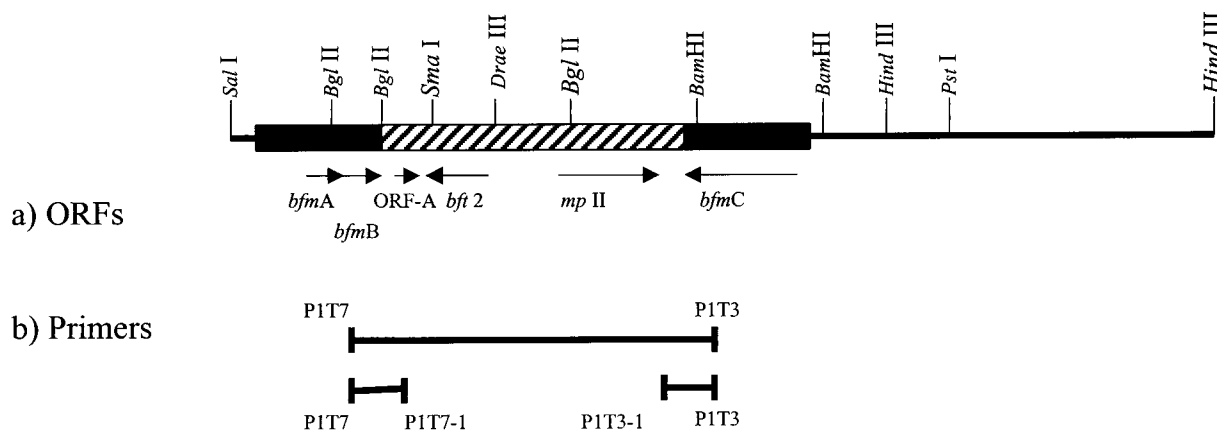


FIG. 2. Partial restriction map of the region flanking *bft* in ETBF 86-5443-2-2. The thicker bar (both solid and striped) shows the region sequenced, and the striped bar shows the BfPAI. (a) ORFs detected in the BfPAI and flanking regions. Arrows show the locations of the ORFs and the direction of their transcription. (b) Relative positions of the primers and the expected PCR products to determine the integration site of the BfPAI. Only the restriction sites important for the construction of the probes (Fig. 1) are shown in the figure.

signed. In colony blot hybridization, Olibft3 hybridized with all 113 ETBF strains containing either *bft-1* or *bft-2*, but none of 191 NTBF strains. These results strongly suggest that *bft-1/bft-2* and *mpII* colocalize in ETBF strains.

The G+C content of the specific 6-kb region (35%) differs substantially from that reported for the *B. fragilis* chromosome (42%) (45). This result and the presence of *bft-2* and *mpII* in the 6-kb locus, which is unique to ETBF strains, suggest that this element is a pathogenicity island or islet (herein designated BfPAI for *B. fragilis* pathogenicity island).

**Nucleotide sequence of the DNA region flanking the BfPAI.** Analysis of the sequence of the left end flanking the BfPAI revealed two ORFs of 540 and 948 bp (Fig. 2a), which encode predicted proteins of 316 and 180 amino acids residues with calculated molecular masses of 21.37 and 37.24 kDa, respectively. A search in the GenBank and EMBL databases for related sequences revealed that these two ORFs, as well as the region immediately upstream of the ORFs, have the same organization as the mobilization region of the 5-nitroimidazole and clindamycin resistance plasmids of *B. vulgatus* (pIP417) and *B. fragilis* (pBFTM10), respectively (12, 47). The mobilization regions of both antibiotic resistance plasmids consist of two genes, named *btgA* and *btgB* in the clindamycin resistance plasmid, and *mobA* and *mobB* in the 5-nitroimidazole resistance plasmid. *BtgA/MobA* and *BtgB/MobB* proteins share 49 and 70% identity, respectively. Both genes are organized in an operon, and immediately upstream of the operon there is a putative origin of transfer (*oriT*). By using the NCBI BLAST server (1), the highest homology score for the predicted 316-amino-acid protein was found with the mobilization protein MobB [P(N)4.5e-15 by BLAST algorithm], with 41% of identity between amino acids 42 and 84, 46% identity between amino acids 176 and 203, 33% identity between amino acids 214 and 234, 46% identity between amino acids 17 and 31, and 38% identity between amino acids 139 and 151. The predicted 180-amino-acid protein shared 25 and 40% identity and similarity, respectively, with a limited region (102 amino acids) of MobA (with both the Gapped BLAST and PSI-BLAST searches). Due to the sequence similarity of the predicted proteins encoded by the ORF of 948 bp (316 amino acids) to MobB and the ORF of 540 bp (180 amino acids) to MobA and the analogous gene arrangement, we designated these ORFs *bfmB* (*B. fragilis* mobilization gene B) and *bfmA* (*B. fragilis* mobilization gene A), respectively.

Figure 3A shows the comparison of this region upstream of the BfPAI and the mobilization region of plasmid pBFTM10. Similar to *btgB/mobB* and *btgA/mobA*, the ATG initiation codon of *bfmB* overlaps the final codon of *bfmA*, suggesting that both genes are transcribed together, and the region immediately upstream of these genes revealed several features consistent with the origin of transfer (*oriT*) of conjugative or mobilizable plasmids (Fig. 3B). This region is ca. 200 bp long and has a higher A+T content than flanking regions (72 versus 56%). The region contains three inverted repeat (IR) sequences (IR1, IR2, and IR3) with lengths of 7, 13, and 9 bp, respectively (Fig. 3B). None of these three IR sequences have similarity with those found in the putative *oriT* from plasmid pIP417 or pBFTM10. The IR1 and IR3 elements have perfect right (R)- and left (L)-arm inverted nucleotide sequences. Only 11 of the 13 bp of IR2R and IR2L are identical. IR1(R)-IR1(L) and IR3(R)-IR3(L) are separated by 3 and 8 bp, respectively. Approximately 165 bp upstream of the start codon of *bfmA* a putative nick site was identified according to the consensus sequence established by Pansegrau and Lanka (31) from the IncP plasmid (RK2/RP4) (Fig. 3C). There was no

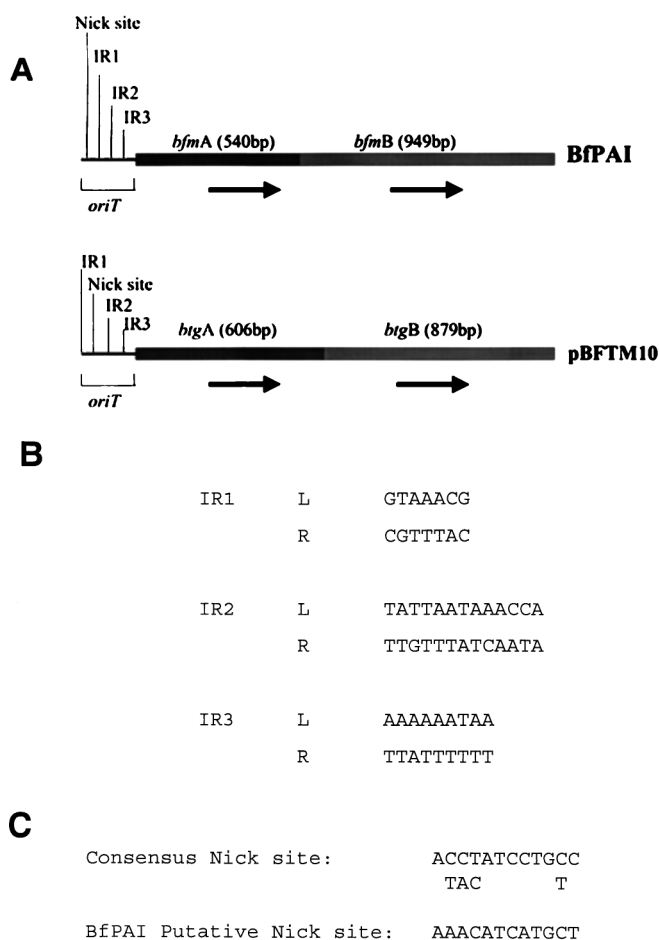


FIG. 3. Putative mobilization region of the BfPAI. (a) Comparison between the putative mobilization region of BfPAI and the mobilization region of the clindamycin resistance plasmid pBFTM10. Arrows show the direction of gene transcription. (b) Nucleotide sequence (5' to 3') of the three IRs. (c) Nick site present in the putative *oriT* gene of the BfPAI.

sequence similarity with the proposed nick region of plasmids pIP417 and pBFTM10.

The right end flanking the BfPAI contained an ORF (designated *bfmC*) of 1,602 bp which transcribes in the same direction as *bft-2* but in the opposite direction of *bfmA* and *bfmB* (Fig. 2a). The *bfmC* gene yields a predicted protein product (BfmC) of 534 residues with a molecular mass of 61.0 kDa. By using the BLAST search program, BfmC shares the highest homology score with a conjugal transfer protein TrsK of conjugative bacteriocin-producing plasmid pMRC01 from *Lactococcus lactis* [P(N)5e-09] (5) sharing 22% identity and 41% similarity over 395 amino acids. BfmC also shares significant similarity with mobilization proteins TraG of *Helicobacter pylori* [P(N)1.5e-07], TrsK of plasmid pGO1 from *Staphylococcus aureus* [P(N)3e-04], TraD of plasmids F and R100 from *E. coli* [P(N)0.003], and TrbC of *Salmonella typhimurium* [P(N)0.092] (14, 25, 29, 52). BfmC contains a perfect ATP/GTP-binding motif A (48) at the amino-terminal region; the same motif appears to be present in TrsK encoded by plasmids pMRC01, as well as TraD encoded by plasmids F and R100. Alignment of these proteins in the region surrounding this motif is shown in Fig. 4. The organization of the left and right ends flanking the BfPAI suggests that the BfPAI may have integrated into a plasmid. However, several different types of plasmid prepara-

BfmC	N F L V Y <u>G G A G S G K T K S I</u>
TrsK (pMRCO1)	N v M v i G G S G S y K T Q S V
TrsK (pGO1)	N i f v v G G p G S y K S a g y
TraD (R100)	N F c L h G t v G A G K S E v I
TraD (F)	N F c L h G t v G A G K S E v I

FIG. 4. Alignment of the region surrounding the ATP/GTP-binding site motif A in BfmC, TrsK (pMRCO1), TrsK (pGO1), TraD (R100), and TraD (F). Amino acids similar to those of the BfmC sequence are in capital letters and boldface. The ATP/GTP-binding site motif A in BfmC is underlined.

tion procedures performed on ETBF strain 86-5443-2-2 failed to reproducibly reveal detectable plasmid DNA (data not shown).

Analysis of the G+C content of the left-end (47%) and right-end (50%) sequences flanking the BfPAI revealed that the G+C content of these regions is significantly different from that of the BfPAI (Fig. 5). Interestingly, the G+C content of the regions immediately upstream of the *bfmA/bfmB* operon, *bft*, *mpII*, and *bfmC* all decrease sharply (asterisks in Fig. 5). Similar results were found by Smith in the DNA sequences of transposon Tn4555 and plasmid pBI143 (45a). This may be due to the fact that promoter regions often have higher A+T content to facilitate separation of DNA for the initiation of transcription. The differing G+C contents of the BfPAI, the flanking region, and the *B. fragilis* chromosome suggest that the BfPAI is contained in yet another foreign genetic element.

**Determination of the integration site of the BfPAI.** Based on the results of colony blot hybridization (Fig. 1) and the above sequence analysis, we hypothesized that ETBF strains could be generated by the integration of the BfPAI in certain NTBF strains (hybridization pattern III). Since the exact left and right borders of the BfPAI were contained in the DNA region spanning probes A and D, respectively (Fig. 1), two primers (P1T7 and P1T3 sequences in Table 2) for these regions were designed (Fig. 2b) and used in PCR experiments to amplify DNA from one strain of each pattern of hybridization. As expected, PCR yielded a product (ca. 1.6 kb) only in the NTBF strain with patterns III (strain I-1345) and IV (strain LM20) but not in any strain with the other patterns of hybridization (Fig. 6A). NTBF strains of patterns II and V did not hybridize with probes A and/or D, which contain primers P1T7 and/or P1T3,

respectively (Fig. 1), and in ETBF strains (pattern I) primers P1T7 and P1T3 flank a ca. 7-kb region (Fig. 2b), a fragment too large to be amplified by *Taq* DNA polymerase under the conditions tested. To determine whether the integration of ETBF is site specific, we examined five additional NTBF strains with hybridization pattern III. All strains yielded products of 1.6 kb similar to strain I-1345 (data not shown). In addition, we designed two new primers (P1T7-1 and P1T3-1) containing sequences for the region in the BfPAI spanning probes G and B, respectively (Fig. 2b). When P1T7 and P1T7-1 were used as primers, PCR experiments with six ETBF strains yielded the same predicted product (1.3 kb) (Fig. 6B). Similarly, the six ETBF strains yielded the predicted 1.1-kb product when P1T3 and P1T3-1 were used as primers (Fig. 6C). All of these results indicate that the integration of the BfPAI is in the same small region and in the same orientation in the cases examined and suggest that the integration of the BfPAI is site specific and unidirectional.

To determine the actual integration site and the exact borders of the BfPAI, we sequenced the 1.6-kb PCR product obtained from NTBF strain I-1345 by using primers P1T7 and P1T3. Comparison of this sequence and the appropriate sequence of ETBF strain 86-5443-2-2 defined the integration site and the putative left and right borders of the BfPAI (Fig. 7). Integration of BfPAI in ETBF strains is predicted to lie between the stop codons of *bfmB* and *bfmC*. ETBF strains lack a span of 16 bp at the integration site which is present in pattern III NTBF strain I-1345. According to this alignment, the BfPAI is calculated to be 6,036 bp in length. Analysis of the borders of the BfPAI revealed nearly perfect 12-bp direct repeats present 45 and 40 bp from the left and right ends of the BfPAI, respectively (data not shown).

**Sequence comparison of 86-5443-2-2 and VPI 13784 BfPAIs.** Moncrief et al. (24) recently reported the sequence of the pathogenicity island (islet) from an ETBF strain containing *bft* subtype 1 (strain VPI 13784). Comparison of the BfPAI sequence from strain 86-5443-2-2 and the reported sequence from ETBF strain VPI 13784 reveals that the two sequences are not identical. A summary of this comparison is shown in Table 5. Overall, both sequences share 97% identity. However, the BfPAI region containing the *mpII* gene is more conserved than the region containing *bft* (99 versus 95% identity, respec-

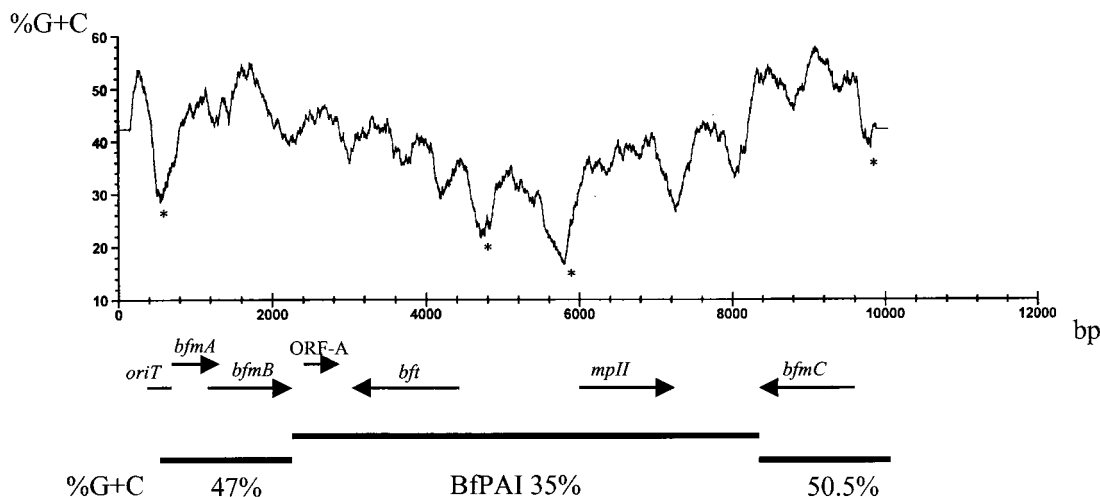


FIG. 5. G+C content across the BfPAI and its flanking regions. Arrows show the location of genes and the direction of their transcription. Asterisks show the sharp declines in G+C content immediately upstream of the designated genes.

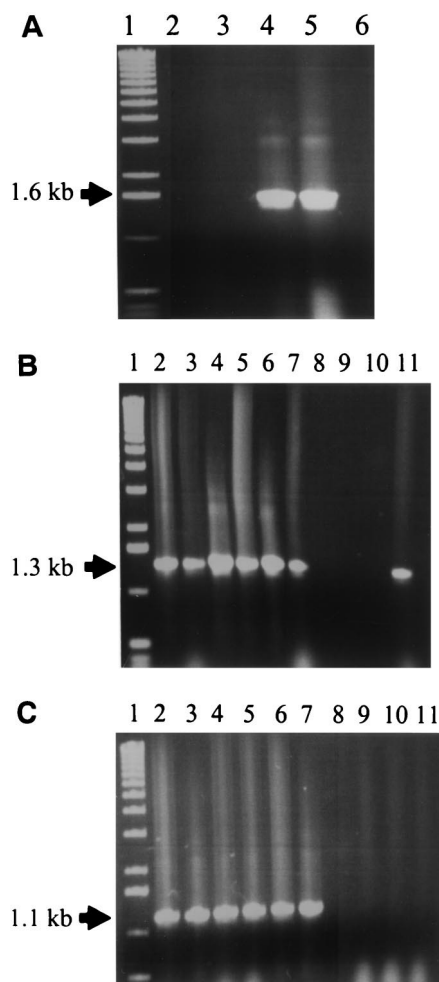


FIG. 6. PCR analysis of the left and right junctions of BfPAI. (A) Screening for the integration site by using primers P1T7 and P1T3 (see Fig. 2b). PCR yielded a product (1.6 kb) only in NTBF strains with pattern III (strain I-1345) and IV (strain LM20). Lanes: 1, 1-kb ladder molecular weight marker; 2, ETBF 86-5443-2-2 (pattern I, positive control); 3, NTBF 077225-2 (pattern II); 4, NTBF I-1345 (pattern III); 5, NTBF LM-20 (pattern IV); 6, NTBF LM-12 (pattern V). (B) Screening for the left junction by using primers P1T7 and P1T7-1 (see Fig. 2b). PCR yielded a similar-sized fragment (1.3 kb) in six ETBF strains (pattern I) and NTBF LM12 (pattern V). Lanes: 1, 1-kb ladder molecular weight marker; 2, ETBF 86-5443-2-2 (pattern I); 3, ETBF VPI 13784 (pattern I); 4, ETBF J-38-1 (pattern I); 5, ETBF DS-64 (pattern I); 6, ETBF DS-233 (pattern I); 7, ETBF DS-49 (pattern I); 8, NTBF 077225-2 (pattern II); 9, NTBF I-1345 (pattern III); 10, NTBF LM-20 (pattern IV); 11, NTBF LM-12 (pattern V). (C) Screening for the right junction by using primers P1T3-1 and P1T3 (see Fig. 2b). PCR yielded a similar-sized fragment (1.1 kb) in all six ETBF strains (pattern I). Lanes are as described for panel B.

tively). Similarly, the predicted proteins encoded by *mpII* from ETBF 86-5443-2-2 and VPI 13784 are 99% identical; whereas BFT-1 from VPI 13784 and BFT-2 from 86-5443-2-2 are 92% identical (9).

Near the left end of the *bft* gene from strain VPI 13784, Kling et al. identified an ORF (ORF1) of 261 nucleotides that encodes a predicted protein with homology to a snake cytotoxin (16). This ORF1 was not identified in the BfPAI from strain 86-5443-2-2. Deletion of nucleotide 16 and the addition of one nucleotide between bases 47 and 48 and bases 142 and 143 of ORF1 changes the frame three times in strain 86-5443-2-2. To determine whether ORF1 was specific for ETBF strains containing *bft-1*, we amplified by PCR and sequenced

the region spanning ORF1 from a strain containing *bft-1* (strain J38-1) and a strain containing *bft-2* (strain 20793-3). We found that the nucleotide sequence of strains J38-1 and 20793-3 were identical to the appropriate sequences of strains VPI 13784 and 86-5443-2-2, respectively. Conversely, we also found that ORF-A detected in BfPAI from strain 86-5443-2-2 (Fig. 2a) was not present in strain VPI 13784. The deletion of nucleotide 60 and the addition of one nucleotide between bases 172 and 173 of ORF-A change the frame twice in strain VPI 13784. Sequence analysis revealed that ORF-A was not present in either strain J38-1 or 20793-3. However, the region spanning this ORF in strain 20793-3 shared 99% similarity with that from 86-5443-2-2 but only 95% similarity with that of VPI 13784, whereas the same region from strain J38-1 shares 99% similarity with that of strain VPI 13784 and 95% similarity with that of strain 86-5443-2-2. The role of the proteins encoded by ORF1 and ORF-A in the virulence of strains VPI 13784 and 86-5443-2-2, respectively, if any, remains to be determined.

The near-direct repeat sequences present at the borders of both BfPAIs are identical. We found that the insertion site of the BfPAI in strain 86-5443-2-2 is similar to that of strain VPI 13784 and that the precise location of the BfPAI is between the stop codons of *bfmB* and *bfmC* (Fig. 7). Our results indicate that the first nucleotide (G) of the 17 bp found in the NTBF strains (but not in the ETBF strains) reported by Moncrief et al. (24) is part of the *bfmC* stop codon, suggesting that 16 bp may be the target sequence for the integration site of the BfPAI and that the size of the VPI 13784 BfPAI is 6,032 bp. However, we cannot rule out the possibility that the last base of the *bfmC* codon is part of the target sequence for the integration of the BfPAI.

**Mobilization of the BfPAI.** Analysis of the nucleotide sequence of the DNA region flanking the BfPAI suggested that *bft-1/bft-2* and *mpII* are in a genetic element that may be mobilized from ETBF to NTBF strains. Transfer of *Bacteroides* plasmids pIP417 and pBFTM10 is linked to the presence of the *Bacteroides* tetracycline resistance transposons (termed  $Tc^r$  elements) (12, 36, 47).  $Tc^r$  elements are large chromosomal self-transmissible elements (ca. 65 to 150 kb) that also facilitate the mobilization of other elements, such as coresident plasmids, other transposons, and discrete unlinked 10- to 12-kb segments of chromosomal DNA nonreplicating *Bacteroides* units (NBUs) (36). Interestingly, the transfer efficiency of  $Tc^r$  elements is enhanced by pretreatment of donor cells with concentrations up to 1  $\mu$ g of tetracycline per ml. Under this condition,  $Tc^r$  elements may be transferred in a frequency of  $10^{-3}$  to  $10^{-7}$ , and ca. 10% of transconjugants will contain  $Tc^r$  elements which cotransferred with plasmids, other transposons, or NBU elements (36). To determine whether  $Tc^r$  elements can mobilize the transfer of BfPAI, we performed mating experiments between ETBF strain 086Thy<sup>-3</sup> (a thymidine-dependent mutant strain derived from ETBF 86-5443-2-2 which contains an inducible  $Tc^r$  transposon [see Materials and Methods]) and NTBF strain TM4000 (tetracycline sensitive). However, under different conditions of induction, we failed to detect tetracycline-resistant transconjugants, indicating that the  $Tc^r$  element of ETBF 086Thy<sup>-3</sup> was not mobilized. When we used the mating positive controls, *B. thetaiotaomicron* strains BT4107 (donor) and BT4001 (recipient), transfer of  $Tc^r$  element was observed at a frequency of  $1.3 \times 10^{-5}$  to  $2.4 \times 10^{-6}$ . These results suggested that the  $Tc^r$  element is not conjugative in strain 086Thy<sup>-3</sup>, or, alternatively, that BFT and/or MP11 may inhibit the mobilization of the  $Tc^r$  element. To test these possibilities, we selected ETBF strain I-1356-1Thy-1 (containing *bft* subtype 1) as donor strains, as well as NTBF strains K518Thy<sup>-2</sup> (pattern II) and I-1345Thy<sup>-2</sup> (pattern III). No

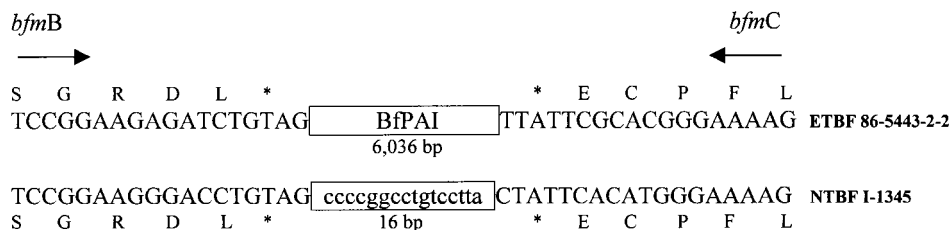


FIG. 7. Comparison of the nucleotide-amino acid sequences of the region flanking the BfPAI of ETBF strain 86-5443-2-2 with the equivalent region of NTBF strain I-1345. Nucleotide sequences belonging to *bfmB* and *bfmC* are given in capital letters. Arrows show direction of gene transcription, and stop codons are indicated by asterisks. The 16-bp sequence lost in ETBF is given in lowercase letters.

transfer of Tc<sup>r</sup> element was observed under any of the conditions of induction used when ETBF I-1356-1Thy-1 or NTBF K518Thy<sup>-2</sup> were used as the donor strains. However, the Tc<sup>r</sup> element was transferred at a frequency of  $1.5 \times 10^{-5}$  to  $1.0 \times 10^{-6}$  when NTBF I-1345Thy<sup>-2</sup> was used as a donor strain. To determine whether the region flanking BfPAI was cotransferred with the Tc<sup>r</sup> element in strain I-1345Thy<sup>-2</sup>, we performed colony blot of 600 Tc<sup>r</sup> transconjugants by using the region spanning *bfmB* as a probe. No transfer of the region flanking BfPAI was detected in any of the 600 tetracycline-resistant transconjugants probed.

**Evaluation for the biological activity of MPII.** BFT (BFT-1 or BFT-2) alters the morphology and physiology of human intestinal epithelial cells (HT29/C1, T84) (49); recently, we demonstrated that BFT cleaves the zonula adherens protein, E-cadherin (51). To determine whether MPII had a biological activity similar to that of BFT-1 and BFT-2, MPII was overexpressed in *E. coli* by using the T7 polymerase system (see Materials and Methods), and its biological activity was tested on HT29/C1 cells. By Western blot analysis, proteins with a molecular mass consistent with the proprotein-mature protein (44 kDa) and mature protein (20 kDa) were detected in the supernatants of whole-cell lysates, but no protein was detected in the cell-free supernatants (data not shown), suggesting that MPII is synthesized as a precursor protein similar to BFT (9, 16) and then processed to a mature 20-kDa protein. However, only a small amount of the putative pre-pro-mature protein was processed to a 20-kDa protein, indicating that MPII is not fully processed or secreted in *E. coli*. The supernatant of whole-cell lysates of MPII clones neither had biological activity on HT29/C1 cells nor cleaved E-cadherin (data not shown).

## DISCUSSION

The data contained in this report strongly suggest that ETBF strains arose via the chromosomal acquisition of a 6-kb region containing the *bft* gene from a foreign source. By studying a collection of *B. fragilis* strains isolated from different countries and from different sources (intestinal and extraintestinal), we showed that this 6-kb region of DNA is found exclusively in ETBF strains. Of note, at least an additional 12 kb of flanking DNA were also found in about half of the NTBF strains, suggesting a second recombination event potentially modulating *B. fragilis* virulence. We have further found that the DNA flanking the ETBF-specific 6-kb region contains mobilization genes and have confirmed that the 6-kb region contains both *bft* and a second metalloprotease gene. We propose that, consistent with the report of Moncrief et al. (24), the specific 6-kb locus be called a BfPAI.

Recently, it has been found that genes involved in virulence, the presence of which distinguish pathogenic and nonpathogenic strains of a species, are often clustered in pathogenicity islands (10, 19). For example, enteropathogenic *E. coli* (EPEC), enterohemorrhagic *E. coli*, and uropathogenic *E. coli* all contain pathogenicity islands inserted at the identical site in the nonpathogenic *E. coli* chromosome (2, 21). In addition, pathogenicity islands have been identified in *Yersinia pestis* (6), *Salmonella typhimurium* (22, 41), *Clostridium difficile* (11), *Vibrio cholerae* (15), and other bacterial pathogens. The BfPAI is similar to other pathogenicity islands in that (i) it contains virulence genes (*bft* and possibly *mpII*), (ii) it has direct repeat sequences at the ends of the locus and is flanked by putative

TABLE 5. Sequence comparison of the BfPAI from strains 86-5443-2-2 and VPI 13784

Nucleotide sequence or coding region	Value
<b>Nucleotide sequence</b>	
Length <sup>a</sup> .....	6,036 bp (86-5443-2-2), 6,032 bp (VPI 13784)
Overall % identity.....	97
Overall number of gaps.....	33
% Identity of region surrounding <i>bft</i> (bp 1 to 3,232).....	95
No. of gaps in region surrounding <i>bft</i> (bp 1 to 3,232).....	33
% Identity of region surrounding <i>mpII</i> (bp 3,233 to 6,036).....	99
No. of gaps in region surrounding <i>mpII</i> (bp 3,233 to 6,036).....	0
% Identity of partial direct repeats at the ends.....	100
<b>Coding region</b>	
% Identity of BFT.....	92
% Identity of MPII.....	99
ORF-A.....	Not present in VPI 13784
ORF1.....	Not present in 86-5443-2-2

<sup>a</sup> Defined by using 16 bp as the target sequence for integration of the BfPAI (see Fig. 7).



mobilization genes, and (iii) it has a G+C content different from the rest of the chromosome.

Our previous study (9) showed that there are two isotypes of *bft* (*bft-1* [VPI-*bft*] and *bft-2* [086-*bft*]) whose predicted encoded proteins share 92% identity (95% identity in nucleotide sequence). In studying a collection of ETBF strains we found that all ETBF strains contain *bft* but the two alleles are found in different amounts (9). The results found in this study suggest that the divergence found between *bft-1* and *bft-2* is perceived in the BfPAI sequence. *B. fragilis* strains are composed of different genetic groups (33), and no predominant clones have been found between the ETBF and NTBF strains (43). The identification of two groups of ETBF strains based on the sequence of the BfPAI supports two paths of evolution for these organisms. Perna et al. (32), suggested that the variability found between the locus of enterocyte effacement from an enterohemorrhagic *E. coli* O157:H7 strain and that of an EPEC strain might originate by natural selection after specific host adaptation or evasion of the host immune response. We have not found any correlation between *bft* subtype and source of the strains (intestinal or extraintestinal and human or animal) (9, 9a). Interestingly, we found that the region of the BfPAI surrounding *mpII* is more conserved than the region surrounding *bft*. The conservation of the MPII sequence in strains 86-5443-2-2 and VPI 13784 suggests that this protein is intolerant of even conservative amino acid changes and that MPII may have an important role in ETBF strains and/or in ETBF-induced disease. However, no biological activity has yet been identified for this protein.

The BfPAI is integrated specifically between potential mobilization genes (Fig. 2a) having significant homology with proteins encoded by conjugal plasmids. It has been reported that *Bacteroides* mobilizable plasmids and transposons utilize a common mobilization strategy. That is, the genes for DNA-processing enzymes and the *cis*-acting *oriT* are located on an adjacent DNA sequence in a small, compact mobilization region (45). The region flanking the left end of the BfPAI has an organization similar to these mobilization regions: two genes organized in an operon (*bfmA/bfmB*) and an *oriT* located adjacent to these genes (Fig. 3A). Phylogenetic analysis of the protein coding sequence of the mobilization region of *Bacteroides* mobilizable plasmids and transposons has shown that they can be clustered together in three major groups based on protein sequence homology (44). We found that the putative mobilization region of BfPAI has the same organization as the mobilization regions of *Bacteroides* antibiotic resistance plasmids pBFTM10 and pIP417. These two plasmids are clustered in the second major homology group of *Bacteroides* mobilization proteins, a plasmid system that requires two gene products for transfer (*btgA* and *btgB* in pBFTM10 and *mobA* and *mobB* in pIP417) (12, 47). Parsimony analysis of the alignment of BfmB with the mobilization proteins of plasmids pIP417 and pBFTM10, as well as other *Bacteroides* mobilizable elements, showed that the mobilization proteins of BfPAI are clustered in the same group as plasmids pIP417 and pBFTM10 (data not shown). It has been proposed that BtgA/MobA and BtgB/MobB bind to the *oriT* region and catalyze the nicking reaction that initiates the transfer process (45, 47). BfmB shares significant identity with BtgB/MobB; however, BfmA shares weak identity with MobA, and the sequence of the putative *oriT* of BfPAI does not share sequence similarity with those of plasmids pIP417, pBFTM10, or other *Bacteroides* mobilizable elements, suggesting that the mobilization region of the BfPAI is unique.

Despite the similarity of the organization of the putative BfPAI mobilization region to that of the mobilization regions

of *Bacteroides* mobilizable plasmids and transposons, it was not mobilized by Tc<sup>r</sup> elements under the conditions studied. This result indicates that the BfPAI mobilization region is not activated by the Tc<sup>r</sup> element regulatory proteins RteA and RteB as occurs with other *Bacteroides* mobilizable elements (46). The right end flanking the BfPAI contains an ORF, whose encoded protein (BfmC) shares significant amino acid similarity with TraD-related proteins. It has been demonstrated that TraD is essential for DNA transfer (13), and it has been proposed that TraD has some role in coupling transfer of the unwound T strand to the transfer pore (18). The similarity of BfmC to TraD proteins suggests that BfPAI might be contained in a self-transmissible element which contains its own transfer regulatory proteins. Alternatively, it has been suggested that certain pathogenicity islands were originally plasmids that were integrated into chromosomal sites and subsequently lost their replication functions (2, 11).

The presence of the BfPAI in a region that appears to be of plasmid origin suggests that ETBF strains might have originated by horizontal transfer of the BfPAI and its flanking element. The different G+C content of the BfPAI (35%) and the flanking elements (47 to 50%) versus that for the *B. fragilis* chromosome (42%) indicates that these two genetic elements come from very different sources. We found that 52% of NTBF strains, in addition to lacking the BfPAI, do not contain the mobilization genes flanking the BfPAI (pattern II), whereas 43% of the NTBF strains lack the BfPAI but present these mobilization genes (pattern III). One hypothesis to explain the origin of these NTBF strains as well as ETBF strains (pattern I) is that the 6-kb region (BfPAI) and the flanking region may be different mobile elements that can be spread independently, namely, pattern II NTBF strains may acquire by horizontal transfer the element containing the mobilization genes, yielding pattern III NTBF strains. Subsequently, these pattern III strains may acquire the BfPAI, yielding ETBF. The 16-bp GC-rich sequence present in NTBF strains with pattern III (Fig. 7) may be the target sequence for integration of the BfPAI into the flanking element. Consistent with this hypothesis, higher numbers of NTBF strains containing the mobilization genes (pattern III) have been isolated in regions where ETBF strains are more common than in places where ETBF were isolated at a low frequency (Table 4). Alternatively, the flanking element may have previously acquired the BfPAI, forming a single mobile element. In this case, ETBF strains might originate by horizontal transfer of this element into NTBF with pattern II, and pattern III NTBF strains might originate by spontaneous deletion of the 6-kb BfPAI by recombination within the 12-bp direct repeats at the ends of the BfPAI.

We found that a small percentage of strains lack portions of the BfPAI as well as its flanking regions (patterns IV and V). All of these NTBF strains were isolated from extraintestinal sources, suggesting that these strains might result from spontaneous deletions after adaptation to the extraintestinal environment. Schubert et al. (39) found that some *E. coli* strains can present the "high-pathogenicity island" of *Yersinia* species but that in 3% of these *E. coli* strains the pathogenicity island manifests deletions of different sizes. These authors suggested that *E. coli* strains acquired the *Yersinia* pathogenicity island by horizontal transfer and then suffered spontaneous deletions after adaptation to a new environment.

In summary, the present study indicates that a BfPAI is a constant feature of the recently recognized ETBF strains. However, divergence within this BfPAI suggests two paths of evolution for ETBF organisms. The BfPAI and its flanking region (with homology to mobilizable plasmids) appear to be

two distinct foreign genetic elements, suggesting that ETBF evolved by horizontal transfer of these two elements into NTBF strains. Our future studies will focus on determining the size and chromosomal location of the locus flanking the BfPAI, as well as on determining whether these loci are self-transmissible elements.

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