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Characterization of a Periplasmic ATP-Binding Cassette Iron Import System of *Brachyspira (Serpulina) hyodysenteriae*

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The nucleotide sequence of the pathogenic spirochete *Brachyspira hyodysenteriae* bit (for “Brachyspira iron transport”) genomic region has been determined. The bit region is likely to encode an iron ATP-binding cassette transport system with some homology to those encountered in gram-negative bacteria. Six open reading frames oriented in the same direction and physically linked have been identified. This system possesses a protein containing ATP-binding motifs (BitD), two hydrophobic cytoplasmic membrane permeases (BitE and BitF), and at least three lipoproteins (BitA, BitB, and BitC) with homology to iron periplasmic binding proteins. These periplasmic binding proteins exhibit lipoprotein features. They are labeled by [3H]palmitate when tested in recombinant *Escherichia coli*, and their signal peptides are typical for substrates of the type II secretory peptidase. The FURTA system and Congo red assay indicate that BitB and BitC are involved in iron binding. The Bit system is detected only in *B. hyodysenteriae* and is absent from *B. innocens* and *B. pilosicoli*.

In Gram-negative bacteria, active transport of molecules from the periplasm to the cytosol occurs via well-conserved systems (28). These specialized high-affinity transport systems are members of the ATP-binding cassette (ABC) superfamily of transporters, which are also responsible for the transport of a wide variety of solutes, including sugars, iron, amino acids, ions, proteins, and polysaccharides (19, 28). The common denominator of ABC transporters is the presence of high-homology stretch of about 200 amino acids in one of the polypeptides. This stretch contains a highly conserved ATP-binding motif consisting of two conserved sites (A and B) that form an ATP-binding pocket, also called the Walker motif (14). The classical ABC transporters possess a common structure consisting of four domains that can be on separate polypeptides; two are hydrophobic integral membrane proteins (permeases), and two are hydrophilic ATP-binding proteins, which can act as homodimers and couple ATP hydrolysis to import movement. In the bacterial ABC transporters-importers, the cytoplasmic permease and the nucleotide-binding protein activities exist on separate polypeptides (21, 28). The periplasmic proteins of importer systems are solute specific and determine the substrate specificity of the transport systems. The periplasmic substrate-binding protein interacts with and concentrates the specific incoming substrate and then presents it to the import complex formed by the cytoplasmic membrane permease and the nucleotide-binding protein.

Cytoplasmic permeases are energized by ATP (reviewed by Ames in 1990 [4]). Common amino acid sequences (EAA---G--------I-LP) were observed in the hydrophilic region of cytoplasmic permeases (33). These sequences, called EAA loops, have been proposed to interact directly with portions of the nucleotide binding site of the ATP-binding protein (33).

Transport of free iron from the periplasmic space into the cytoplasm is proposed to occur by a classic active-transport (import) process involving a periplasmic binding protein, a specific cytoplasmic permease, and an energy-supplying nucleotide binding protein. This active-transport system is widely distributed among gram-negative pathogenic bacteria.

Spirochetes form a distinct bacterial group within the eubacterial kingdom (61, 70). *Brachyspira hyodysenteriae* (Serpulina hyodysenteriae [48]) is a member of the spirochete family and is the agent of swine dysentery (26). *B. hyodysenteriae* is characterized by its strong beta-hemolytic activity and its enteropathogenicity in swine (34). Different virulence factors of *B. hyodysenteriae* have been cloned and identified. Inactivation of flagellar proteins (FlaA1 and FlaB1) resulted in a strong reduction in virulence and colonizing ability, indicating that motility is an essential virulence factor of *B. hyodysenteriae* (32, 55, 56). Different hemolysins have also been cloned and are considered important for the virulence of *B. hyodysenteriae* (29, 43, 67, 68). A *B. hyodysenteriae* mutant with a mutation of one of the hemolysins (thyA) showed a reduced virulence in mice in a swine dysentery model (29, 68). Furthermore, *B. hyodysenteriae* possesses a NADH oxidase (Nox) (63) that reduces chemically the oxygen in water. The NADH oxidase may play an essential role in the colonization of the mammalian intestinal tract by diverse bacterial species.

In this study, we describe the cloning and characterization of an ABC importer system of *B. hyodysenteriae* implicated in iron import (the Bit system). The Bit system possesses all the features characteristic of bacterial periplasmic importers; it bears strong primary sequence and secondary-structure similarities to the importer systems, suggesting that it has a similar mechanism of action. We suggest that in *B. hyodysenteriae* the periplasm-to-cytosol transport of iron proceeds by an analogous mechanism to those observed in gram-negative bacteria.
The Bit system has similarities to the periplasmic binding protein-dependent iron transport systems in *Actinobacillus pleuropneumoniae* (*afuABC*) (13), *Haemophilus influenzae* (*bitABC*) (58), *Neisseria meningitidis* (*fbpABC*) (1), and *Serratia marcescens* (*sfuABC*) (6).

MATERIALS AND METHODS

Bacterial strains, plasmids, and growth conditions. *B. hyodysenteriae*, *B. innocens*, and *Escherichia coli* strains, as well as plagues and plasmids used in this study, are listed in Table 1. Bacteria were grown on solid medium by using blood agar base no. 2 (Oxoid Ltd., Basingstoke, England) containing 5% bovine blood. The plates were incubated anaerobically at 37°C for 4 days in jars (Oxoid) under a GasPak Plus generator atmosphere (BBL, Becton Dickinson and Co., Cockeysville, Md.). *E. coli* LE392, used for phage multiplication, was grown in Luria-Bertani medium under aerobic conditions.

**TABLE 1. Strains, plagues, and plasmids used in this study**

<table>
<thead>
<tr>
<th>Strain, plague, or plasmid</th>
<th>Genotype</th>
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<tbody>
<tr>
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<tr>
<td>B78</td>
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<td>Reference strain serotype 8</td>
<td>(ATCC 49887)</td>
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<td></td>
</tr>
<tr>
<td>B1265</td>
<td>Type strain (ATCC 29796)</td>
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<td><strong>E. coli</strong> K-12</td>
<td>F′<em>hsdR15 (ts−mcr−)</em> supE44 supF58 lacY1 recA1 endA1 gyrA96 thi-1 hsdR17 supE44 relA1 lacI2ΔM15 Tn10 (Tetr)</td>
<td>Promega</td>
</tr>
<tr>
<td>XL1 Blue</td>
<td></td>
<td>Stratagene</td>
</tr>
<tr>
<td><strong>Phage</strong> λGEM-11</td>
<td>Bacteriophage vector, SpI selection</td>
<td>Promega</td>
</tr>
<tr>
<td><strong>Plasmids</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pBluescript II KS(+)</td>
<td>Ap′ Plasmid vector</td>
<td>Stratagene</td>
</tr>
<tr>
<td>pDJ2</td>
<td>pKS(+) bciBCDEF</td>
<td>This study</td>
</tr>
<tr>
<td>pDJ3</td>
<td>pKS(+) bciA′B′</td>
<td>This study</td>
</tr>
<tr>
<td>pDJ4</td>
<td>pKS(+) bciF</td>
<td>This study</td>
</tr>
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<tr>
<td>pDJ9</td>
<td>pKS(+) bciBC</td>
<td>This study</td>
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**FM88-90** whole cells (41). The antiserum was preabsorbed overnight at 4°C with *B. innocens* reference strain B256 and *E. coli* LE392. Bound antibodies were detected with horseradish peroxidase-conjugated goat anti-rabbit immunoglobulin G (Jackson Immunoresearch, West Grove, Pa.) and 4-chloro-1-naphthol (Sigma) as described by Sambrook et al. (57). Selected positive clones were purified by several rounds of single-plaque isolation. These purified plagues were then tested for their nonreactivity with an antiserum of *B. innocens* type strain B256 by Western blotting and sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE); the positive ones were discarded. Plague 42 has been selected for further studies.

DNA purification. Phage DNA was extracted from each of the positive plagues that reacted only with the *B. hyodysenteriae* antiserum, by scraping off agarose containing nearly confluent plaques in dilution buffer. After 3 h of elution at room temperature, the agarose and bacterial cells were pelletted by centrifugation. The DNA was then extracted from the supernatant by an extraction method involving Phase Lock Gel II (3Prime-3Prime, Boulder, Colo.). Plasmid DNA was isolated from *E. coli* by alkaline extraction (57).

Southern blotting and hybridization. DNA was transferred to Zeta-Probe blotting membranes (Bio-Rad Laboratories Ltd, Mississauga, Ontario, Canada). DNA probes were labeled with [32P]dCTP (DuPont Canada, Mississauga, Ontario, Canada) by using a random oligolabelling kit (Pharmacia-LKB, Biotechnology Inc., Dorval, Canada) as previously described (25). The blots were hybridized with DNA probes as recommended by the supplier of the Zeta-Probe blotting membranes and exposed for autoradiography.

SDS-PAGE and Western blot (immunoblot) analysis. Proteins were purified by scraping the top agarose from plates containing nearly confluent plaques into phase dilution buffer. After a 3-h elution period, the agarose was pelleted and proteins in the supernatant were recovered by precipitation with 10% trichloroacetic acid as previously described (27), washed twice in phosphate-buffered saline, suspended in Laemmli sample buffer (37), and subjected to SDS-PAGE. Protein profiles of subclones were analyzed for the expression of *B. hyodysenteriae* proteins by Western immunoblot analysis with whole bacterial cells.

DNA sequencing. Restriction fragments of DNA from recombinant plagues were ligated into phage vectors KS plasmid. Plasmids carrying the appropriate inserts were isolated from double-stranded DNA sequencing with both strands as template by the dideoxynucleotide method of Sanger et al. (59) with the AutoRead sequencing kit (Pharmacia-LKB). Oligonucleotides were synthesized on a Cyclone nucleic acid synthesizer (Millipore Corp., Bedford, Mass.); DNA sequence analysis was performed with the GeneWorks software package (IntelliGenetics, Inc., Mountain View, Calif.). DNA sequence information was analyzed with the Intelligenetics Suite package and programs from the University of Wisconsin Genetics Computer Group software sequence analysis package (18).

Analysis of hydrophathy and sequence homology of proteins. Average hydrophathy profiles of the proteins were obtained by the method of Kyte and Doolittle (36) with the GeneWorks program (IntelliGenetics), using a window length of 9. Amino acid comparisons were effected with the BESTFIT program and the Intelligenetics Suite package (IntelliGenetics, Inc., Mountain View, Calif.). DNA sequence information was analyzed with the Intelligenetics Suite package and programs from the University of Wisconsin Genetics Computer Group software sequence analysis package (18).

[1H]Palmitate labeling and immunoprecipitation of BitB and BitC lipoproteins. The transformants were labeled in vivo with [1H]palmitate as previously described (54). Briefly, transformants XL1 containing pDJ2, pDJ3, and pDJ5 were grown aerobically at 37°C in LB containing ampicillin (100 mg/liter) and by adding 0.01% to Trypticase soy agar plus 10% IsoVitalex (BBL, Becton Dickinson and Co.) for determination of the Crb (Congo red-binding) phenotype (8). The DNA was transferred to Zeta-Probe membranes (Bio-Rad Laboratories Ltd, Mississauga, Ontario, Canada). DNA probes were labeled with [32P]dCTP (DuPont Canada, Mississauga, Ontario, Canada) by using a random oligolabelling kit (Pharmacia-LKB, Biotechnology Inc., Dorval, Canada) as previously described (25). The blots were hybridized with DNA probes as recommended by the supplier of the Zeta-Probe blotting membranes and exposed for autoradiography.

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**F[3H]palmitate labeling and immunoprecipitation of BitB and BitC lipoproteins.** The transformants were labeled in vivo with [3H]palmitate as previously described (54). Briefly, transformants XL1 containing pDJ2, pDJ3, and pDJ5 were grown aerobically at 37°C in LB containing ampicillin (100 mg/liter) and by adding 0.01% to Trypticase soy agar plus 10% IsoVitalex (BBL, Becton Dickinson and Co.) for determination of the Crb (Congo red-binding) phenotype (8). The DNA was transferred to Zeta-Probe membranes (Bio-Rad Laboratories Ltd, Mississauga, Ontario, Canada). DNA probes were labeled with [32P]dCTP (DuPont Canada, Mississauga, Ontario, Canada) by using a random oligolabelling kit (Pharmacia-LKB, Biotechnology Inc., Dorval, Canada) as previously described (25). The blots were hybridized with DNA probes as recommended by the supplier of the Zeta-Probe blotting membranes and exposed for autoradiography.

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transcription of the fhuF::lacZ. The lacZ expression assay was performed with MacConkey lactose plates (Difco) supplemented with 25 to 50 mM FeCl$_3$·6H$_2$O.

**Immunofluorescence purification of antibodies.** To detect the presence of the cloned proteins in *B. hyodysenteriae*, the rabbit antiserum raised against *B. hyodysenteriae* was enriched for antibodies to the Bit proteins by the method of Oakes et al. (47). This enriched antiserum was used to localize the proteins on the bacteria by immunoelectron microscopy. Rabbit anti-*B. hyodysenteriae* polyclonal antiserum was incubated with nitrocellulose onto which clone 42 plaques had first been adsorbed and blocked with 2% skim milk–Tris-buffered saline. After a 3-h incubation period, the nitrocellulose was washed first in Tris-buffered saline with 0.05% Tween 20, then in Tris-buffered saline, and then in 0.15 M NaCl. Finally, the bound antibodies were eluted in 0.2 M glycine–0.15 M NaCl (pH 2.8). The antibody solution was then neutralized with 0.75 M Tris (pH 8.8) and used undiluted for electron microscopy.

**Electron microscopy and immunogold labeling.** The electron microscopy and immunogold labeling were done as previously described (39). Briefly, a drop of bacterial suspension was placed on a Formvar-coated grid. After blocking for 5 min with 1% egg albumin, the grids were incubated for 30 min with undiluted antibody-enriched solution (described above) and rinsed with distilled water five times. The grids were then incubated for 30 min with protein-A gold particles (10 nm; Sigma), rinsed, and negatively stained with 1% phosphotungstate for 10 s. They were examined under an electron microscope (Philips 201) at an accelerating voltage of 60 kV.

**Nucleotide sequence accession number.** The nucleotide sequence data reported in this paper appears in the GenBank Database Library under accession no. U75349.

## RESULTS

**Construction and screening of the library.** Chromosomal DNA isolated from strain FMS8-90 of *B. hyodysenteriae* serotype 8 was partially digested with *Sac*I, and the sites were partially filled with dGTP and dATP. A library was then constructed in AGEM-11 phage. We obtained approximately 3 × 10$^9$ recombinants per μg of DNA. To obtain clones expressing *B. hyodysenteriae* specific proteins, the library was screened with anti-*B. hyodysenteriae* rabbit sera adsorbed against *B. innocens* B256 reference strain and *E. coli* LE392. Five recombinant phages positive for *B. hyodysenteriae* and nonreactive with *B. innocens* were chosen. We report here the characterization of the recombinant bacteriophage 42.

**Restriction mapping and expression of recombinant proteins.** The recombinant bacteriophage 42 consisted of the vector AGEM-11 and a genomic insert of 6.6 kb. The insert was then subcloned in the phagemid pBluescriptII KS as a *Sac*I–*Sac*I fragment (Fig. 1). The plasmid was named pDJ1. Western blot analysis of pDJ1 revealed two proteins with similar molecular masses of 42 and 43 kDa that reacted with the anti-*B. hyodysenteriae* specific antibodies raised against the recombinant bacteriophage 42.

**Imunoaffinity purification of antibodies.** Plasmids pDJ5 and pDJ6, containing an insert from pDJ1 was sequenced by gene-walking PCR. The procedure is based on single-specific-primer PCR amplification of chromosomal DNA adjacent to the known sequence (46). The sequence bitA (1,008 nucleotides) encodes a putative polypeptide of 336 amino acids with a deduced molecular mass of 37,027 Da. BitA has homology to many proteins involved in iron periplasmic binding, such as AfuA (56% similarity) in *A. pleuropneumoniae* (12) and a putative periplasmic iron-binding protein (51.6%) of *Sinorhizobium meliloti* (GenBank accession no. AAC64671). BitA has homology of 49.4% to a thiamin-binding lipoprotein of a member of the *Archaea*, *Pyrococcus horikoshii* (31). Downstream of bitA, a 1,026-nucleotides ORF, bitB, encodes a putative polypeptide of 342 amino acids (BitB), with a deduced molecular mass of 37,262 Da. BitB has homology to many proteins involved in iron periplasmic binding, such as AfuA (51.3% similarity) in *A. pleuropneumoniae* (13) and HitA (45.5% similarity) from the system HitABC in *H. influenzae*. The third (1,017-nucleotide) ORF, bitC, encodes a putative polypeptide of 339 amino acids (BitC) with a deduced molecular mass of 36,885 Da. The amino acid sequences of BitA, BitB, and BitC showed homology to each other (approximately 90% similarity). The genes encoding BitB and BitC expressed a 42- and a 43-kDa protein, respectively, as revealed by Western blotting with anti-*B. hyodysenteriae* serum (Fig. 2).

The fourth ORF, bitD (of 1,122 nucleotides) encodes a putative polypeptide of 374 amino acids (BitD) with a deduced molecular mass of 42,589 Da. BitD has homology to many proteins belonging to the superfamily of the ABC transporter proteins and contained an ATP-binding domain consisting of two highly conserved amino acid sequences known as the Walker motif A (amino acids 37 to 53) and Walker motif B (amino acids 134 to 167 and 188 to 194). More specifically, the BitD protein has homology to a hypothetical ABC transporter of *H. influenzae*, Hi0126 (67.1% similarity), to PotA (63.7% similarity) of *H. influenzae*, to AttA1 (63.6% similarity) of *Agrobacterium tumefaciens*, to AfuC (62.8% similarity) of *A. pleuropneumoniae*, and to HitC (58% similarity) of *H. influenzae*.

The fifth ORF (bitE) of 831 nucleotides encodes a polypeptide of 277 amino acids (BitE) with a deduced molecular mass of 30,853 Da. The sixth ORF (bitF) of 771 nucleotides encodes a polypeptide of 257 amino acids (BitF) with a deduced molecular mass of 28,343 Da. Both BitE and BitF proteins have homology to hydrophobic permeases and show homology to each other (48.7% similarity). BitE has homology to the first half of AfuB (54.4% similarity) (*A. pleuropneumoniae*) and to the first half of HitB (58.4% similarity) (*H. influenzae*). BitF has homology to the second half of AfuB (57.8% similarity) and to the second half of HitB (48.6% similarity). Both AfuB and HitB are permeases involved in iron transport. BitE and BitF proteins also have homology to a putative sulfate permease, NifC, of *H. influenzae*, also named Hit0129 (22). BitE has homology (55.0% similarity) to the first half of NifC, and BitF has homology to the second half (54.6% similarity). BitE and BitF are very hydrophobic proteins, their mean hydrophobicity (36) being 0.89 and 0.91, respectively. Their hydrophathy profiles indicate that BitE and BitF contain six putative transmembrane-spanning segments linked by hydrophilic segments of variable length (not shown). A sequence matching the consensus permease EAA motif (EAA---G-------------I-LP) (33, 60) was found in both BitE and BitF (amino acids 177 to 198 for BitE and 157 to 178 for BitF).

A homology search using the default parameters of Blastp at the National Center for Biotechnology Information and restricted to the amino acid sequences of the spirochete protein has been applied to BitABCDEF. BitA showed homology to a
thiamine ABC transporter of *Treponema pallidum* (accession no. AAC65133), 47% similarity on a stretch of 78 amino acids. BitB showed homology to the surface protein VspA of *Borrelia turicatae* (12), 37% similarity on a stretch of 216 amino acids. BitC, like BitA, showed homology to a thiamine ABC transporter of *T. pallidum* (AAC65133), 43% similarity on a stretch of 138 amino acids. BitD showed homology to an ATP-binding protein (PotA) of the spermidine/putrescine ABC transporter of *T. pallidum* (AAC65627), 63% homology on a stretch of 296 amino acids. BitE showed homology to a permease protein (PotB) of the spermidine/putrescine ABC transporter of *Borrelia burgdorferi* (AAB91526), 45% homology on a stretch of 269 amino acids. Finally, BitF showed homology to another permease protein (PotC) of the spermidine/putrescine ABC transporter of *B. hyodysenteriae* (AAB91526), 37% similarity on a stretch of 306 amino acids.
transporter of *B. burgdorferi* (AAB91527), 48% homology on a stretch of 263 amino acids.

Preceding the putative ATG start codon of all ORFs, a ribosome-binding site characteristic of gram-negative bacteria was found. The first 20 amino acids of BitB and the first 21 amino acids of BitC, the putative proteins homologous to the periplasmic protein-binding substrate, have putative lipoprotein cleavage signal sequences (GCG software). No typical signal peptide sequence was found for BitD, the putative ATP-binding protein, or for BitE or BitF, which are homologous to hydrophobic permeases. Moreover, 235 and 210 consensus sequences (24) were found upstream of the sequence of *bitB* and *bitC*. A potential rho-independent termination site has been found just downstream the last ORF of the Bit system, *bitF*. Furthermore, the 84-bp intragenic region between the *bitC* stop codon and the first ATG codon of *bitD* includes a strong stem-loop structure with the potential to form 26 hydrogen bonds with a ΔG of −14.6 kcal/mol as determined with Oligo version 5.0 distributed by National Biosciences (Hamel, Minn.).

Thus, sequence analysis indicates that the Bit polypeptides belong to an ABC transporter system where BitD has the characteristic feature of an ABC transporter and BitE and BitF have the characteristics of permeases and homologies to permeases. Moreover, the homologies of Bit periplasmic binding proteins consistent with many periplasmic binding proteins involved in iron periplasmic binding and the overall homology of BitD, BitE, and BitF to the ABC transporter and permeases of an iron importer system suggest that the Bit system could be involved in iron import. A dendrogram was constructed by using amino acids sequences of periplasmic iron binding (PIB) proteins present in different gram-negative bacteria (Fig. 3). Two clusters were obtained. The PIB proteins sequences of Bit system were grouped with those of some members of the *Pasteurellaceae*, *H. influenzae* and *A. pleuropneumoniae*, in division I. The majority of the sequences of the PIB proteins of gram-negative bacteria were clustered in division II. Division II was formed by the sequences of the PIB proteins belonging to FbpABC, HitABC, and YfuABC systems.

**BitB and BitC are lipoproteins.** The signal sequences of the BitA, BitB, and BitC proteins are typical type II signal sequences that could be cleaved by the type II signal peptidase. Transformant cells harboring pBluescript II KS(−) (negative control), pDJ2 (BitB), and pDJ5 (BitC) were labeled in vitro with [3H]palmitate (Fig. 4). The size of the labeled lipoproteins

<table>
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<th>Iron-PBP</th>
<th>Organism</th>
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<td>Hi0131</td>
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<td>BitB</td>
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</table>
expressed by pDJ2 and pDJ5 corresponded to the size of BitB and BitC proteins when compared in Western blots where these recombinant proteins were revealed by antiserum against \textit{B. hyodysenteriae}. These lipoproteins were absent from the cells containing the plBluescript II KS without insert. Thus, BitB and BitC are periplasmic binding lipoproteins. When the prebleed antiserum was used, no palmitate-labeled lipoproteins were immunoprecipitated from the different transformants (data not shown). Furthermore, when \textit{B. hyodysenteriae} antiserum was used, no palmitate-labeled lipoproteins were immunoprecipitated from transformant cells harboring plBluescript II KS without insert (data not shown).

Characterization of the BitA/BC system by FURTA and Congo red binding. FURTA was used to determine if the \textit{bit} genes contain \textit{fur} boxes but also can reveal the presence of an iron-binding system. \textit{E. coli} H1717 contains an iron-regulated \textit{fhuF:\lacZ} fusion which is very sensitive to small changes in the iron concentration in the medium because the \textit{fhuF} promoter region has a weak binding affinity for the Fur-Fe$^{2+}$ repressor. The repressor Fur in presence of iron binds to the \textit{fhuF} promoter region, and therefore the \textit{lacZ} reporter gene is not expressed and colonies are white on MacConkey lactose plates. However, Fur will not bind to the \textit{fhuF} promoter region in the absence of iron or in the presence of Fur binding sequences carried on a multicopy plasmid that will also titrate out the Fur protein from binding sites within \textit{fhuF}. In both of these latter cases, the \textit{lacZ} reporter gene will be expressed and the colonies will be red on MacConkey lactose plates. Three putative Fur boxes were identified by homology to the putative Fur-box of \textit{E. coli} (44). Two of them are localized between or near the −10 and −35 sites of \textit{bitB}. One sequence which shares 12 of 19 positions with the consensus Fur-binding site overlaps the promoter of \textit{bitB} (nucleotides 1048 to 1066), which is 100\% identical to the promoter of \textit{fepA} of \textit{E. coli}. Similarly the \textit{fepA} −35 promoter sequence is also localized into a putative Fur box. The \textit{FepA} protein is part of the ferric enterobactin transport system in \textit{E. coli} (49). The second Fur box (nucleotides 1159 to 1177) is localized just downstream of the ribosome-binding site of \textit{bitB} (13 nucleotides identical to the 19 nucleotides of the Fur box). The third Fur box (nucleotides 3302 to 3320) has been found between BitC and BitD (14 nucleotides identical to the 19 nucleotides of the \textit{E. coli} Fur box).

To determine whether these Fur boxes were binding the Fur protein, we introduced constructs pDJ1, pDJ2, pDJ4, pDJ5, pDJ6, pDJ7, pDJ8, pDJ9, plBluescript KS, and a positive control pBCFUR into \textit{E. coli} H1717. For these tests, two different concentrations of iron (FeCl$_3$) were tested; 25 and 50 $\mu$M. No LacZ expression was observed when strain H1717 was transformed with plasmids carrying a putative Fur box as deduced by sequence homology (pDJ8, pDJ9, and pDJ3) (Fig. 1). This indicates that the sequence carried by these recombinant plasmids did not titrate out the Fur repressor from binding to \textit{fhuF}-binding sites. However, colonies of strain H1717 transformed with plasmids expressing BitB and/or BitC (pDJ2, pDJ5, and pDJ6) were red on MacConkey plates. This observation suggests that in the presence of the periplasmic binding proteins, \textit{E. coli} Fur repressor is not binding the \textit{fhuF} promoter region, because BitB and BitC are also competing for iron. Thus, BitB and BitC are periplasmic iron-binding proteins and the Bit system is involved in iron transport. Furthermore when the constructs pDJ1, pDJ2, pDJ4, pDJ5, pDJ6, pDJ7, pDJ8, pDJ9, and plBluescript KS were transformed into \textit{E. coli} XL1 Blue, only colonies transformed with plasmids expressing BitB and/or BitC (pDJ2, pDJ5, and pDJ6) were red on Congo red-containing plates (Fig. 1). This indicates that these proteins have the capacity to bind Congo red.

Localization of the Bit proteins in \textit{B. hyodysenteriae}. To locate the proteins expressed from phase 42, an immunogold-labeling experiment was performed on \textit{B. hyodysenteriae} cells grown on blood agar plates. The antiserum against \textit{B. hyodysenteriae} affinity purified from phase 42 expressing the Bit proteins by using the method of Oaks et al. (47). Rabbit anti-\textit{B. hyodysenteriae} polyclonal antiserum was incubated with nitrocellulose adsorbed with phase 42. The antiserum did not react with \textit{E. coli} cells used for the phase infection. These antibodies specific to the proteins expressed by phase 42 were used to localize the Bit proteins on the bacteria by immunogold electron microscopy. The affinity-purified antiserum was tested in electron microscopy on whole cells of \textit{B. hyodysenteriae} and \textit{B. innocens}. The antigens were localized external to the cytoplasmic membrane of \textit{B. hyodysenteriae} cells and could be detected when the outer sheet was disrupted (Fig. 5A). Intact \textit{B. hyodysenteriae} cells were poorly labeled with the anti-Bit protein antibodies (data not shown).

Genetic characterization of the Bit system. The Bit system of \textit{B. hyodysenteriae} was characterized genetically by Southern hybridization analysis of genomic DNAs of \textit{B. hyodysenteriae} and \textit{B. innocens}. The probe \textit{bit} consisted of a SacI-Clal fragment including \textit{bitA}, \textit{bitB}, and \textit{bitC} genes. The probe hybridized with BglII-digested DNAs from the \textit{B. hyodysenteriae} parental strain and from the type strain of \textit{B. hyodysenteriae} (Fig. 6). The \textit{bit} probe detected 4 BglII fragments of 1.0, 1.9, 10, and $>$12 kb in each strain of \textit{B. hyodysenteriae} tested. The 1.0-kb fragment corresponded to the BglII internal fragment of the \textit{bit} DNA (Fig. 1). Since two BglII sites are present in the \textit{bit} DNA sequenced, three fragments should hybridize with the \textit{bit} probe. The additional fragment hybridizing with the \textit{bit} probe indicates that there is an additional sequence to homologous \textit{bit} periplasmic genes in the \textit{B. hyodysenteriae} genome. Furthermore, the \textit{bit} probe did not hybridize with \textit{B. innocens}, a non-pathogenic spirochete, or with \textit{E. coli} HB101. \textit{B. hyodysenteriae} serotype 1 to 9 reference strains, as well as \textit{B. hyodysenteriae} B78$^T$, \textit{B. innocens} B25$^T$, and \textit{B. pilosicoli} P43/6/78$^T$, were tested by Southern blotting with the \textit{bit} probe (results not shown). Only \textit{B. hyodysenteriae} strains hybridized with the \textit{bit} probe. Furthermore, by immunodot blotting with anti-Bit serum purified by an immunoadfinity assay as described in Materials and Methods, only \textit{B. hyodysenteriae} strains reacted with anti-Bit serum (results not shown). Thus, these results suggest that the Bit system is specific to \textit{B. hyodysenteriae}.

**DISCUSSION**

A unique feature of the bacterial ABC importer systems is that they all have a periplasmic substrate-binding protein that interacts with the incoming substrate, binds to it, and presents...
it to the import complex formed by the permease and the ATP-binding protein in the inner membrane (28). This import complex has the ATP-binding domain and the membrane-spanning domains present on separate polypeptide (21, 28). The Bit system presents characteristics of a periplasmic iron transport system. It includes periplasmic proteins encoded by at least three genes, bitA, bitB, and bitC, that are proposed to be the iron-binding proteins. These genes belong to a gene cluster purported to encode a classic import system that includes BitA, BitB, and BitC; a nucleotide-binding protein, BitD; and two cytoplasmic membrane permeases, BitE and BitF. The characteristics and comparison of the predicted protein products of bitDEF with the products derived from the afuBC, fbpBC, hitBC, sfuBC and yfuBC genes illustrate conserved similarity of the Bit system with respect to the general class of ABC iron periplasmic importers.

In the Bit periplasmic transport system, the BitA, BitB, and BitC proteins correspond to the substrate-binding lipoproteins homologous to periplasmic iron-binding protein AfuA of the A. pleuropneumoniae afuABC operon (13), similar to HI0131, which is suggested to be an iron-binding periplasmic protein of H. influenzae (66), and to HitA of the H. influenzae HitABC periplasmic iron import operon (58). The dendrogram obtained from the homologies of the substrate-binding proteins BitB and BitC showed that the periplasmic iron-binding proteins of the Bit system were distant but clustered with the other periplasmic iron-binding proteins. We chose to derive the dendrogram from the homologies and the differences between the iron periplasmic protein of the different iron transport systems since these proteins have a strong amino acid sequence heterogeneity, varying from species to species. Interestingly, the BitB and BitC proteins from the Bit system are grouped with those of another swine pathogen, A. pleuropneumoniae, the causative agent of porcine pleuropneumonia. Spirochetes form a distinct group within the eubacterial kingdom (61). The homologies and differences observed across the Bit system argue that the afu, fbp, hit, sfu, and yfu operons and the bit system have evolved separately but seem to function similarly in the periplasm-to-cytosol transport of iron. The homologies between the BitA, BitB, and BitC amino acid sequences suggest a duplication phenomenon. The sequencing and hybridization results suggested that there are possibly more than three duplications of the periplasmic binding substrate gene in B. hyodysenteriae. This feature has not been observed in homologous iron transport systems of gram-negative bacteria. In H. influ-
enzae at least two related periplasmic iron-binding proteins physically distant on the genome have been identified (58, 66). In B. hyodysenteriae, a family of genes with 45 to 95% homology have been found in a locus called VspA-H (variable surface protein) (23). Furthermore, in spirochetes, gene duplication has been observed, particularly in Borrelia hermsii (35). The consequence of gene duplication in B. hermsii is to evade the host immune response through multiphasic antigenic variation by sequential expression of genes of variable major proteins (52). For the Bit system, the reason for this gene duplication is unclear and the minor amino acid sequence differences observed between the Bit periplasmic binding proteins might reflect a difference in the affinity for iron binding or for different iron sources.

The only proteins identified by Western blotting with B. hyodysenteriae antiserum were the two hydrophilic periplasmic binding lipoproteins BitB and BitC exposed to the periplasm. BitD, BitE, and BitF are more hydrophobic proteins and are hidden into the cytoplasmic membrane and probably not accessible to the anti-whole-cell B. hyodysenteriae antiserum. Interestingly, a strong stem-loop structure was found between the bitC and bitD. This structure was also found in the fbp, hit, and sfu operons, between the gene coding for the periplasmic substrate-binding protein and the gene coding for the ATP-binding protein (1). It has been proposed (1) that since stem-loop structures could function in mRNA stability (51), the presence of this structure might allow bitB and bitC expression at higher levels than that of bitD, bitE, and bitF. Therefore, the products of bitDEF could be less strongly expressed and not detectable. The BitABC transport system includes genes that are physically linked. The different plasmid constructions might indicate that two promoters are likely to exist in the bit gene cluster, since deletion of the 5′ region upstream of bitC, including bitB, did not affect its expression. However, vector promoters can be responsible for the expression observed.

BitD is the ABC transporter polypeptide with two highly conserved sequences, called Walker motifs A and B, which are believed to form the ATP-binding pocket involved in active transport (28).

BitE and BitF are the inner membrane permeases with membrane-spanning segments (six for each protein) that form the pathway through which substrate crosses the membrane. They bear strong homology to each other, indicating that they originated by a gene duplication, as suggested for other cytoplasmic membrane permeases, HisQ and HisM (3). BitE and BitF are supposed to form a pseudodimer that is closely associated with BitD. Similar to the Bit periplasmic iron transport system, the Yfe periplasmic iron transport system of Yersinia pestis possesses two cytoplasmic membrane permeases, YfE and YfED (8). The conserved EAA loop motifs (33) observed in BitE and BitF probably interact directly with the BitD protein.

Iron can exist in two redox states which are dependent upon pH and oxygen availability. Naturally abundant iron is not readily available to microbes in aerobic aqueous environments at neutral pH, where it exists as insoluble ferric complexes. Some gram-negative bacteria such as E. coli solubilize and scavenge ferric iron [Fe(III)] by means of excreted siderophores molecules with high affinities for Fe(III). In anaerobic environment, the iron is in the ferrous [Fe(II)] form (45). An Fe(II) uptake system has been characterized in E. coli (30). Although B. hyodysenteriae has been shown to express at least three iron-regulated proteins when grown under iron-restricted conditions (40), it was shown to be unable to use siderophores such as catechol and hydroxamate. Like other anaerobes, this anaerobic spirochete may not secrete siderophore. Uptake of the metal can occur via a ferrous transport system, from host iron-binding proteins such as degraded products of hemoglobin, or from ferrated transferrin or ferrated lactoferrin. It could be envisioned that degraded products of hemoglobin such as heme or heme are readily available because the hemolytic activity of B. hyodysenteriae could serve as a source of iron. It was observed for a nontypeable H. influenzae strain that mutation of hitC from the hitABC operon prevents this strain to utilize these sources of iron, demonstrating that this system is implicated in the uptake of iron from transferrin or iron chelates (58). It is conceivable that the Bit system described in this report is similarly implicated in iron transport from these host iron sources. The presence of different periplasmic iron-binding proteins in the Bit system tends to support this hypothesis.

We propose a model for the BitABC system. Periplasmic iron-binding proteins (BitA, BitB, or BitC) bind iron from different sources, Fe(III) or heme, and change conformation. Then the protein-iron complex interacts with the cytoplasmic membrane complex formed by the ABC transporter protein (BitD) and the permeases (BitE and BitF). The importation of iron into the cytoplasm of the bacterial cell is then coupled with ATP hydrolysis.

The intracellular level of iron is carefully controlled in the bacterial cell. Storage of iron will reduce the growth of bacteria, whereas a high concentration of the metal can be toxic. Therefore, the expression of iron acquisition systems is regulated in response to iron, being increased under iron limitation. Iron-dependent regulation of genes involved in iron acquisition is under the transcriptional control of the holorepressor composed of the Fur protein and the cofactor Fe(II) in a number of gram-negative bacteria (16, 62). The operator consensus palindromic sequence (iron box or Fur box) has been identified on promoter regions of a number of iron-controlled genes including iron transport-related genes (17). Analysis of the results indicated that the presence or absence of a putative Fur box in different recombinant clones did not affect the level of β-galactosidase expression of the fhuC::lacZ fusion. Although the presence of the bit sequences did not titrate out the E. coli Fur protein from the FURTA system, we cannot conclude that there are no Fur-binding sequences in their insert. We can only conclude that if these sequences are present, the E. coli Fur protein does not recognize them. Fur regulation might not act on the regulation of bit gene expression as demonstrated for the N. gonorrhoeae iron periplasmic import system, fbpABC (17). In spirochetes, no Fur-regulated genes have been identified. Interestingly, a DtxR homologue has been cloned in Treponema pallidum, indicating that regulation of iron in spirochetes might function similarly to some gram-positive bacteria. The first example of a gram-positive iron-regulated promoter-operator was that of the diphtheria toxin gene, toxA, by an Fe(II)-dependent DNA-binding repressor, DtxR (11). It is now clear that DtxR functions as a global iron-sensitive regulatory element in the control of gene expression in some gram-negative bacteria (15). It would be interesting to investigate if the DtxR homologue is present in B. hyodysenteriae.

A correlation has been established between the synthesis of an iron periplasmic transport system and virulence of bacterial pathogens such as N. gonorrhoeae (1, 10), H. influenzae (2, 58), Serratia marcescens (5, 71), A. pleuropneumoniae (13), Yersinia enterocolitica (56a), and Y. pestis (8). Furthermore, the ability to compete for iron in the host environment correlates with pathogenicity of Neisseria species. Indeed, the iron periplasmic binding protein Nibp is expressed at antigenically detectable levels only in the microbial pathogens N. gonorrhoeae and N.
meningitidis but not in commensal Neisseria species such as N. sicca or N. perflava when propagated under iron stress (9, 42). The Bit system is detected only in B. hyodysenteriae and is absent in B. innocens and B. pilosicoli. The Bit system might correlate with the pathogenicity of B. hyodysenteriae. Inactivating the Bit system in B. hyodysenteriae will enable us to better understand its role in the pathogenicity of this spirochete.

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