Nucleotide Sequence and Transcriptional Regulation of a Positive Regulatory Gene of *Shigella dysenteriae*

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A 1,937 bp *PstI-HindIII* fragment containing the *ipaR* locus was cloned from the large invasion plasmid of *Shigella dysenteriae* CG097, and its nucleotide sequence was completely determined. The IpaR protein (35 kDa, calculated from the DNA sequence) was synthesized in *Escherichia coli* χ1411 minicells containing the 1,937-bp *PstI-HindIII* fragment. To determine the regulatory role of *ipaR* for *ipa* genes, we applied genetic complementation experiments using chloramphenicol acetyltransferase (CAT) as reporter. Analyses of CAT activity of the recombinant plasmids containing the 5' flanking sequences of the 24-kDa-protein gene and the *ippI*, *ipaB*, *ipaC*, and *ipaD* genes defined strong promoters upstream of the *ippI* and *ipaB* genes, and the absence of any promoter activity for the *ipaC* gene. Complementation analyses showed that the CAT activity only under direction of the *ippI* promoter region increased 1.8-fold in the presence of IpaR protein. On the basis of our data, we suggest that an operon comprising *ippI*, *ipaB*, and *ipaC* is positively regulated by IpaR protein which has a trans effect on a DNA sequence upstream of the *ippI* promoter.

Shigellae cause bacillary dysentery in humans and monkeys. Invasion of epithelial cells of the colon constitutes one of the earliest steps in *Shigella* infection. Genes required for the full expression of virulence are on both the *Shigella* chromosome and a nontransmissible large plasmid (180 to 230 kb) (9, 19, 20, 23). Invasion plasmid antigens IpaB, IpaC, and IpaD are associated with the initial adhesion and invasion of colonic epithelial cells (3, 6, 11, 14, 21, 29). Expression of virulence genes is temperature regulated, wild-type strains being virulent when grown at 37°C but phenotypically avirulent at 30°C (12). Maurelli and Sansonetti (13) demonstrated that temperature regulation of virulent gene expression in *Shigella flexneri* is negatively controlled by virR, a chromosomal gene. Studies of *S. flexneri* YSH6000 and M90T have shown that a plasmid gene, *ipaR* (or *virB*), acts as a positive regulator, at the transcriptional level, for the expression of *ipaB*, *ipaC*, *ipaD*, and *ipaA* (1, 7). An identical regulatory gene, designated invE, has also been found in *S. sonnei* (27). However, such a study has not been done with *S. dysenteriae*, which causes bacillary dysentery in children with greater severity than dysentery caused by other *Shigella* species or enteroinvasive *Escherichia coli*. Furthermore, the molecular mechanism by which *ipaR* (or *virB*, or *invE*) activates the transcription of *ipaB*, *ipaC*, *ipaD*, and *ipaA* is not known.

In this work, we report the presence of the *ipaR* gene in the invasion plasmid of *S. dysenteriae* CG097 and describe its positive regulation on the expression of *ippI*, *ipaB*, and *ipaC* genes through a trans effect on the 5' flanking region of the *ippI* gene.

**MATERIALS AND METHODS**

**Bacterial strains.** *E. coli* HB101 was used as the host in recombinant DNA cloning and for chloramphenicol acetyltransferase (CAT) expression. *E. coli* JM103 was used as a host for M13 bacteriophage. *S. dysenteriae* 60R (30) was used as the host for CAT expression. A minicell-producing strain, *E. coli* χ1411, was used for analysis of plasmid-encoded proteins.

**Cloning vectors and DNA methods.** Plasmids pUC8 (BRL), pKK232-8 (Pharmacia) (5), and pACYC177 (gift of M. P. Jackson) were used for DNA cloning, and vectors M13mp18 and M13mp19 were used for DNA sequencing. Isolation of plasmid DNA, digestion of restriction enzymes, recovery of DNA fragments from agarose gel, ligation, and transformation were performed as described by Sambrook et al. (17).

**Plasmid constructions.** Plasmids pWS102, pWS103, and pWS104 previously described (29), containing *S. dysenteriae* *ipa* genes, were the source for cloning of *ipa* promoter regions in this study. The 5' flanking sequence of the 24-kDa-protein gene excised from pWS104 by *BamHI* and *HindIII* digestions (396 bp, −317 to +79, with respect to the initiation codon of the 24-kDa-protein gene), a 772-bp *HindIII* fragment of the 5' flanking sequence of the *ippI* gene (−605 to +167) from pWS104, a 1,707-bp *PvuII-SalI* DNA fragment of the 5' flanking sequence of the *ipaB* gene from pWS102 (−431 to +1171), also containing 105 bp of pUC8 preceding −431, an 850-bp *SphI-BamHI* fragment of the 5' flanking region of the *ipaC* gene from pWS102 (−406 to +426, also containing 18 bp of pUC8 following +426), and a 632-bp *HincII* fragment of the 5' flanking sequence of the *ipaD* gene from pWS103 (−423 to +209) were inserted into the multiple cloning sites upstream of the CAT gene of the pKK232-8 plasmid, to yield recombinant plasmids pWS396, pWS772, pWS1602, pWS832, and pWS632, respectively (Table 1). The direction of each inserted DNA fragment was identified by analysis of the cleavage patterns generated by the specific restriction endonucleases.

To construct the 5' deletion mutants of the *ippI* promoter region, a 562-bp *BamHI-HindIII* fragment (−395 to +167, 210-bp deletion from the 5' end of the 772-bp *HindIII* fragment) and a 414-bp *HaeIII-HindIII* fragment (−247 to +167, 358-bp deletion from the 5' end of the 772-bp *HindIII* fragment) were linked to *SmaI* and *HindIII* sites of.
pKK232-8 to yield pWS210 and pWS358, respectively (Table 1).

To clone the DNA fragment containing the ipaR gene into the pACYC177 plasmid, which is compatible with the replication of pKK232-8 for complementation analyses, a 1,972-bp PvuII fragment from pWS106 (Fig. 1; 1,875 bp from bp 58 to bp 1937 of Fig. 2 and 93 bp of pUC8 following bp 1937) was inserted into the HincII site of pACYC177 to yield pWS1879 (Table 1).

Estimation of plasmid copy number. The plasmid copy number was determined as described by Petersen and Hansen (15). Plasmids pBR322 (15 to 20 copies per chromosome) and pACYC177 (10 to 12 copies per chromosome) (17) were included as controls.

Analysis of plasmid-encoded proteins. The pUC8 plasmids, with and without inserts, were transformed into E. coli χ1411. Minicells prepared from 250-ml overnight cultures of E. coli χ1411 were isolated according to the method of Roozen et al. (16). Plasmid-encoded proteins synthesized in minicells were labeled with [35S]-methionine and analyzed by sodium dodecyl sulfate (SDS)-12% polyacrylamide gel electrophoresis (PAGE) (4).

DNA sequence determination. The DNA sequences were determined by the chain termination method of Sanger et al. (18), by using a T7 sequencing kit (Pharmacia), following cloning into M13mp18 and M13mp19.

Sequence analysis. GENEPRO (Riverside Scientific) was used to search for open reading frames and restriction sites, to predict protein sequences from nucleic acid data, and to analyze hydropathy of polypeptides.

CAT enzyme assay. Bacteria were grown in L broth to an A600 of between 0.5 and 1.0. The harvested bacterial cells were sonicated in an equal volume of 0.25 M Tris-HCl (pH 7.8) for three 10-s treatments with 1-min intervals in an ice-water bath (Fisher Sonic Dismembrator, model 300). After centrifugation (12,000 × g for 10 min at 4°C) to remove cellular debris, the total protein concentrations of extracts were determined by using a biocinchoninic acid protein assay kit (Pierce Chemical Co., Rockford, Ill.). Assays for CAT activity were performed as described previously (4), with an incubation time of 45 min at 37°C. CAT reaction was carried out in the following reaction mixture: 0.01 to 3 μg of cell extract, 20 μl of freshly made acetyl coenzyme A (Sigma) at 3.5 mg/ml, 1 μl of [14C]chloramphenicol, and 2.5 M Tris-HCl (pH 7.8) in a total final volume of 150 μl. After the reaction, the products were separated from unacetylated chloramphenicol on thin-layer chromatography plates (Sigma). Various amounts of lysate were used to ensure proper quantitation. The percentage of acetylation was determined with the AMBISS Radioanalytic Imaging System and converted to units of CAT by using pure CAT (Sigma) as a standard. One unit of CAT activity corresponds to 1 nmol of chloramphenicol acetylated per min at 37°C under optimal assay conditions.

Nucleotide sequence accession number. The nucleotide sequence shown in Fig. 2 has been given GenBank accession number X63593.

RESULTS

Cloning and DNA sequencing of the ipaR region. In the course of DNA sequencing of ipaB, ipaC, ipaD, and ipaA genes from the 9-kb fragment (pWS100) of S. dysenteriae CG097 (29), a truncated open reading frame (ORF) downstream of the ipaA gene was observed. To identify the extent of this ORF, the relevant restriction fragment (1,937-bp Pstl-HindIII) from pWS100 was cloned into pUC8, and the resulting plasmid was designated pWS106 (Table 1). Its restriction map is shown in Fig. 1. Figure 2 presents the nucleotide sequence of this 1.9-kb fragment. The DNA sequence of an open reading frame beginning at bp 597 and ending at bp 1524 shows homology with that of ipaR (or virB or invE) of both S. flexneri and S. sonnei (1, 7, 27) and was

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Table 1. Characterization of recombinant plasmids

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Vector</th>
<th>Promoter region or gene</th>
<th>Inserted DNA</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>pWS106</td>
<td>pUC8</td>
<td>ipaR, 9-kDa-protein gene</td>
<td>1,937 bp</td>
<td>pWS100*</td>
</tr>
<tr>
<td>pWS396</td>
<td>pKK232-8</td>
<td>24-kDa-protein gene</td>
<td>396 bp, -317 to +79</td>
<td>pWS104*</td>
</tr>
<tr>
<td>pWS772</td>
<td>pKK232-8</td>
<td>ippI</td>
<td>772 bp, -605 to +167</td>
<td>pWS104*</td>
</tr>
<tr>
<td>pWS210</td>
<td>pKK232-8</td>
<td>ippI</td>
<td>562 bp, -395 to +167</td>
<td>pWS104*</td>
</tr>
<tr>
<td>pWS358</td>
<td>pKK232-8</td>
<td>ippI</td>
<td>414 bp, -247 to +167</td>
<td>pWS104*</td>
</tr>
<tr>
<td>pWS1602</td>
<td>pKK232-8</td>
<td>ipaB</td>
<td>1,707 bp, -431 to +1171 and 105 bp from pUC8</td>
<td>pWS102*</td>
</tr>
<tr>
<td>pWS1832</td>
<td>pKK232-8</td>
<td>ipaC</td>
<td>850 bp, -406 to +426 and 18 bp from pUC8</td>
<td>pWS102*</td>
</tr>
<tr>
<td>pWS210</td>
<td>pKK232-8</td>
<td>ippD</td>
<td>632 bp, -432 to +209</td>
<td>pWS103*</td>
</tr>
<tr>
<td>pWS1879</td>
<td>pACYC177</td>
<td>ipaR, 9-kDa-protein gene</td>
<td>1,879 bp and 93 bp of pUC8</td>
<td>pWS106, this work</td>
</tr>
<tr>
<td>pWS1009</td>
<td>pACYC177</td>
<td>9-kDa-protein gene</td>
<td>731 bp</td>
<td>pWS106, this work</td>
</tr>
</tbody>
</table>

* Plasmids pWS100, pWS102, pWS103, and pWS104 were described previously (29).
designated ipaR. The amino acid sequence of the IpA protein deduced from the DNA sequences contains 309 amino acids with a molecular mass of 35 kDa. The putative -10 and -35 promoter regions and ribosome binding sites sequences are located upstream from the initiation codon of ipaR. At the 5' end of the 1,937-bp PstI-HindIII fragment, we observed a nucleotide sequence encoding 45 amino acids which overlapped the 3' end of the ipaA gene (Fig. 2) (25).

As shown in Fig. 2, the nucleotide sequence of the 1,937-bp PstI-HindIII fragment (pWS106) reveals two additional small ORFs located upstream of the ipaR gene. The first ORF begins 3 bp downstream from the terminator codon of ipaA (bp 144 to 379) and encodes 78 amino acids with a calculated molecular mass of 9 kDa. The second ORF begins 24 bp downstream from the terminator codon of the first ORF (bp 405 to 582) and codes for 59 amino acids (molecular mass 7 kDa; its amino acids not shown). These two small ORFs were in the same orientation as the ipaR ORF. The putative rho-independent terminator was found at the 3' end of the 9-kDa ORF but not at the 3' end of the 7-kDa or IpA ORF. Interestingly, a typical signal sequence of 19 amino acids was found on the amino-terminal sequence of the 9-kDa polypeptide but not on that of the 7-kDa or IpA polypeptide. This signal sequence contains nine hydrophobic residues preceded by two basic residues, lysines, with serine and alanine at the +1 and +3 positions before the cleavage site (i.e. MIKEKSLVACCG_LAYS' + KLS). In addition, the 9-kDa polypeptide contains two hydrophobic transmembrane domains as predicted by our hydrophathy analysis. Thus, we speculate that this polypeptide is targeted onto the bacterial membrane by using the same translocation mechanism as those of OmpA, OmpC, OmpF, and alkaline phosphatase of E. coli (28).

Minicell analysis of ipaR region products. Figure 3 shows the result of SDS-PAGE of [35S]-methionine-labeled proteins, isolated from the minicell-producing strain of E. coli χ1411 carrying the plasmid pUC8 with and without insert. SDS-PAGE analysis of proteins made in E. coli χ1411 minicells containing pWS106 showed two polypeptides of 35 and 9 kDa (Fig. 3, lane B). The observed molecular masses of these polypeptides agree with the molecular masses of the products calculated from the nucleotide sequence. However, the 7-kDa polypeptide was not seen in the extract of the minicells containing pWS106. No Shine-Delgarno sequence is found upstream of the 7-kDa ORF (Fig. 2), and this probably accounts for its absence in the minicell analysis (Fig. 3, lane B).

Characterization of ipa promoters. On the basis of the
homology of \textit{ipaR} DNA sequences of \textit{S. dysenteriae} CG097 with those of \textit{ipaR} (\textit{virB}, or \textit{invE}) of both \textit{S. flexneri} and \textit{S. sonnei}, which positively regulate the expression of \textit{ipaB}, \textit{ipaC}, \textit{ipaD}, and \textit{ipaA} (1, 7, 27), and in order to determine whether or not IpaR acts on the promoter regions of \textit{S. dysenteriae} \textit{ipa} genes, we first identified the promoters of \textit{ipa} genes by using CAT as the reporter gene. DNA fragments, containing the 5' flanking region of each \textit{ipa} gene with the promoter region and a portion of amino acid codons from the amino terminus, were inserted upstream of the CAT gene of the promoter cloning vector pKK232-8 plasmid in the same orientation as the CAT gene, resulting in the recombinant plasmids pWS396, pWS772, pWS1602, pWS832, and pWS632 (Table 1). These pKK232-8-derivative plasmids containing the 5' flanking region of \textit{ipa} genes were found to be stable in \textit{E. coli} HB101 and \textit{S. dysenteriae} 60R, and the insertion did not affect the plasmid copy number (10 to 12 copies per chromosome). The function of promoters of the \textit{ipa} genes was determined by measuring CAT activity of these recombinant plasmids in \textit{E. coli} HB101 (Fig. 4). Promoterless vector pKK232-8 was included as a negative control (Fig. 4, lane 13). pWS396 (carrying the promoter region of the 24-kDa-protein gene) and pWS632 (carrying the promoter region of the \textit{ipaD}) produced a high level of CAT activity (Fig. 4, lanes 2 and 11). Low CAT activity was detected in pWS772 (carrying the promoter region of \textit{ippI}) and pWS1602 (carrying the promoter region of \textit{ipaB}) (Fig. 4, lanes 4 and 7). CAT was not expressed in pWS832 (carrying the 5' flanking region of \textit{ipaC}) (Fig. 4, lane 9). Similar results were obtained when these recombinant plasmids were introduced into \textit{S. dysenteriae} 60R (data not shown). Strain 60R does not contain the invasion plasmid (30), so it was used as \textit{S. dysenteriae} background. These results suggest that there are strong promoters in the DNA fragments containing the 5' flanking sequences of both the 24-kDa-protein gene and \textit{ipaD} and weak promoters in \textit{ippI} and \textit{ipaB}. Furthermore, it appears that the \textit{ipaC} gene does not have a promoter.

\textbf{IpaR protein has a \textit{trans} effect on the 5' flanking sequence of the \textit{ippI} gene.} To determine whether the \textit{ipaR} product has a \textit{trans} effect on the \textit{ipa} promoter regions identified above, we cloned the 1,972-bp \textit{PvuII} DNA fragment containing the \textit{ipaR} locus from pWS106 into the low-copy-number plasmid pACYC177 (P15A of replicon, stringent), yielding pWS1879 (Table 1). pWS1879 containing the \textit{ipaR} locus is maintained in the same copy number as pACYC177 and stably coexists with pWS396, pWS772, pWS1602, or pWS632 in \textit{E. coli} HB101 and \textit{S. dysenteriae} 60R. As shown in Fig. 4, the CAT activity increased 1.8-fold in complementation of pWS1879 (\textit{ipaR}) with pWS772 (lane 5) in HB101 compared with that of pWS772 only (lane 4). No increased CAT activity was detected in complementation of pWS1879 with pWS396, pWS1602, or pWS632 (Fig. 4, lanes 3, 8, and 12, respectively). Yet no CAT activity was detected in HB101 harboring both pWS832 (carrying the 5' flanking region of \textit{ipaC}) and pWS1879 (\textit{ipaR} locus) (Fig. 4, lane 10). Similar results were observed in the \textit{S. dysenteriae} 60R background (data not shown). Since pWS1879 is derived from pACYC177, which is maintained in lower copy number than pBR322 (17) and is not amplified by chloramphenicol, it is unlikely that the \textit{trans} effect of IpaR protein is a result of overexpression of the \textit{ipaR} gene. Thus, we suggested that IpaR protein has a \textit{trans} effect only on the promoter region of \textit{ippI} but not on the promoter regions of the 24-kDa-protein gene, \textit{ipaB}, and \textit{ipaD}.

Nucleotide sequence and minicell analysis of the 1,937-bp \textit{PstI}-HindIII fragment revealed the presence of a 9-kDa polypeptide upstream of the \textit{ipaR} ORF (Fig. 2 and 3). To determine whether the 9-kDa polypeptide is involved in the regulation, a 731-bp \textit{PvuII-EcoRV} fragment, containing only the 9-kDa ORF, was obtained from pWS106 and cloned into the \textit{HincII} site of the pACYC177 plasmid, yielding pWS1009 (Table 1). No increased CAT activity was observed in complementation of pWS1009 with pWS772 (Fig. 4, lane 6, and Fig. 5), indicating that IpaR but not the 9-kDa polypeptide is involved in the regulation of expression of \textit{ipa} genes.

\textbf{Functional properties of the DNA sequence located in the 5' flanking region of the \textit{ippI} gene.} The above-described results suggest the presence of a DNA sequence responsible to IpaR protein in the 5' flanking sequence of the \textit{ippI} gene. To further map the location of this DNA sequence between \text{−}605 bp and the initiation codon of \textit{ippI}, two deletion mutant plasmids, pWS210 (\text{−}395 to +167) and pWS358 (\text{−}247 to +167), were constructed (Table 1) and their CAT activity was measured in the presence or absence of pWS1879 (\textit{ipaR} gene). The activation of IpaR was abolished in these two
FIG. 4. Fluorography of the products of [14C]chloramphenicol obtained in CAT assays (see Materials and Methods) of E. coli HB101 containing various recombinant plasmids in the presence or absence of pWS1879 (ipaR locus). Lane 1, 0.5 U of chloramphenicol transferase (Sigma); lane 2, pWS396; lane 3, pWS396/pWS1879; lane 4, pWS772; lane 5, pWS772/pWS1879; lane 6, pWS772/pWS1879; lane 7, pWS1602; lane 8, pWS1602/pWS1879; lane 9, pWS832; lane 10, pWS832/pWS1879; lane 11, pWS632; lane 12, pWS632/pWS1879; and lane 13, pKK232-8/pWS1879. Cell extract used: lanes 2 to 8, 11, and 12, 1 μg; lanes 9, 10, and 13, 3 μg. CAT activity was determined as described in Materials and Methods. chl, chloramphenicol; 1-chl, 1-acetyl-chloramphenicol; 3-chl, 3-acetyl-chloramphenicol; 1,3-chl, 1,3-acetyl-chloramphenicol. 

mutant plasmids (Fig. 5). However, pWS210 (with deletion of 210 bp, −395 to +167) produced a level of CAT activity (over fivefold) higher than that of pWS772, whereas a lower CAT activity was detected in pWS358 (deletion of 358 bp, −247 to +167) than in pWS772 (Fig. 5). 

DISCUSSION

Comparison of the nucleotide sequence of the ipaR gene of S. dysenteriae CG097 with those of S. flexneri M90T and YSH6000 and S. sonnei revealed a difference of six bases among these species (1, 7, 27). At position 569, the adenine (A) in the ipaR or virB gene of S. flexneri or the invE gene of S. sonnei was converted into guanine (G) in the ipaR gene of S. dysenteriae. This alteration occurred within the putative −10 promoter region (i.e., TATAAAT in M90T, YSH6000, and S. sonnei; TATGAAT in CG097). Furthermore, the change of this single base (A to G) produced two direct repeats (CAATATGAATCAATATGAAT) in the ipaR promoter region of S. dysenteriae CG097. At bp 1106, the change of G in S. flexneri M90T or YSH6000 or S. sonnei to C in S. dysenteriae CG097 resulted in a new amino acid, 

FIG. 5. Effect of 5′ deletions on CAT expression of pWS772 (containing the promoter region of the ippl gene). pWS772, as well as its 5′ deletion mutants pWS210 and pWS358, was complemented with (+) or without (−) pWS1879 (ipaR locus) or pWS1009 (9-kDa locus). CAT activity was determined as described in Materials and Methods. chl, chloramphenicol; 1-chl, 1-acetyl-chloramphenicol; 3-chl, 3-acetyl-chloramphenicol; 1,3-chl, 1,3-acetyl-chloramphenicol.
histidine, in the IpaR of CG097 instead of the glutamine in IpaR of M90T, YSH6000, and S. sonnei. At bp 780 and bp 1278, thymine in YSH6000 and S. sonnei was substituted for cytosine in CG097 and M90T. However, these two nucleotide changes did not change the corresponding amino acid; in all cases it was leucine. Downstream from the 3' end of IpaR, two additional bases had changed: at bp 1691, A in M90T and YSH6000 was converted into C in CG097 and S. sonnei, and at bp 1763, G in M90T, YSH6000, and S. sonnei was converted into A in CG097.

The promoter cloning vector pKK232-8 has been used to identify and measure relative activities of various promoters (8, 22, 24), since it has three stop codons between the multiple cloning sites and the AUG of the cat gene, which prohibits readthrough into the cat gene (5). Functional analyses of CAT activity of the promoter-CAT derivatives of pKK232-8 containing the 5'-flanking sequences of the 24-kDa protein gene or ipipl, ipaB, ipaC, or ipaD defined the strong promoters upstream of the 24-kDa-protein gene and ipaD and the weak promoters upstream of the ipipl and ipaB genes (Fig. 4). CAT activity was not detected in plasmid pWS832 (carrying the ipaC sequence) in S. flexneri, indicating that the ipaC gene does not contain a promoter, although a putative promoter sequence was found upstream of initiation codon of the ipaC (2, 21, 26, 29). Complementation analyses showed that IpaR positively regulates the expression of CAT under the direction of the ipipl promoter region but not under the direction of the 24-kDa-protein gene, ipaB, or ipaD promoter region. Since the 24-kDa-protein gene, ipipl, ipaB, ipaC, and ipaD of S. dysenteriae CG097 are closely linked and arranged clockwise in the large invasion plasmid (29) and the 24-kDa-protein gene and ipaD are both expressed independently from their respective promoters (Fig. 4, lanes 2 and 11), we deduce that the ipipl, ipaB, and ipaC genes are organized as an operon, and it appears that the ipipl promoter is activated by IpaR in trans to express coordinate these three genes, ipipl, ipaB, and ipaC. Although the ipaB gene contains a weak promoter (Fig. 4, lane 7), it may be nonfunctional in vivo, because IpaR has no effect on it. This is further supported by the work of Venkatesan et al. (26) who detected a 3.3-kb RNA by Northern (RNA) blotting in S. flexneri which encompasses the ipipl, ipaB, and ipaC genes. Although the 2.4- and 1.4-kb RNA bands which hybridized with individual ipaB and ipaC probe were also observed, Venkatesan et al. claimed that these two RNAs could conceivably be derived from the 3.3-kb transcript by an unknown processing scheme (26).

A putative −10 sequence (TATAAA) and a putative −35 sequence (TTGACA) which almost match with TATAAT (−10 sequence) and TTGACA (−35 sequence) of E. coli were found at positions −318 bp and −337 bp, respectively. A sequence of dyad symmetry (TGTtatTTCACTTAA AACA) is found between −605 and −395. IpaR polypeptide shows significant sequence homology with the bacteriophage P1 partition protein ParB (7, 27). The ParB has been identified as a DNA-binding protein, and its recognition site consists of a 34-bp DNA sequence including a sequence of dyad symmetry (10). Therefore, it is presumed that IpaR protein may bind to TGTtatTTCACTTAA AACA sequence and render the binding of RNA polymerase to the ipipl promoter sequences efficient and constitutive. Alternatively, as a positive regulator, IpaR protein may interact either with DNA sequence or with other proteins such as transcriptional factor(s) or repressor. The interaction of IpaR protein with specific DNA sequence or another protein(s) remains to be elucidated.

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