

# Genomic Analysis of a Pathogenicity Island in Uropathogenic *Escherichia coli* CFT073: Distribution of Homologous Sequences among Isolates from Patients with Pyelonephritis, Cystitis, and Catheter-Associated Bacteriuria and from Fecal Samples

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Urinary tract infection is the most frequently diagnosed kidney and urologic disease and *Escherichia coli* is by far the most common etiologic agent. Uropathogenic strains have been shown to contain blocks of DNA termed pathogenicity islands (PAIs) which contribute to their virulence. We have defined one of these regions of DNA within the chromosome of a highly virulent *E. coli* strain, CFT073, isolated from the blood and urine of a woman with acute pyelonephritis. The 57,988-bp stretch of DNA has characteristics which define PAIs, including a size greater than 30 kb, the presence of insertion sequences, distinct segmentation of K-12 and J96 origin, GC content (42.9%) different from that of total genomic DNA (50.8%), and the presence of virulence genes (*hly* and *pap*). Within this region, we have identified 44 open reading frames; of these 44, 10 are homologous to entries in the complete K-12 genome sequence, 4 are nearly identical to the sequences of *E. coli* J96 encoding the HlyA hemolysin, 11 encode P fimbriae, and 19 show no homology to J96 or K-12 entries. To determine whether sequences found within the junctions of the PAI of CFT073 were common to other uropathogenic strains of *E. coli*, 11 probes were isolated along the length of the PAI and were hybridized to dot blots of genomic DNA isolated from clinical isolates (67 from patients with acute pyelonephritis, 38 from patients with cystitis, 49 from patients with catheter-associated bacteriuria, and 27 from fecal samples). These sequences were found significantly more often in strains associated with the clinical syndromes of acute pyelonephritis (79%) and cystitis (82%) than in those associated with catheter-associated bacteriuria (58%) and in fecal strains (22%) ( $P < 0.001$ ). From these regions, we have identified a putative iron transport system and genes other than *hly* and *pap* that may contribute to the virulent phenotype of uropathogenic *E. coli* strains.

*Escherichia coli* is by far the most common cause of urinary tract infection (UTI), particularly in uncomplicated cases. Strains causing these infections possess traits that distinguish them from commensal strains of *E. coli* and other pathogenic strains such as those causing diarrhea and meningitis. Characteristically, uropathogenic strains of *E. coli* are composed of a restricted number of O serogroups, produce hemolysin, P fimbriae, and aerobactin, exhibit serum resistance, and are encapsulated (6, 7, 14, 15, 18, 31, 35). The presence of these features, not found in the typical fecal strain, implies that uropathogenic strains possess a defined set of virulence determinants that allow the bacterium to colonize the urinary tract, avoid host defenses, and elicit histological damage to the uroepithelium, allowing in some cases passage of the bacterium into the bloodstream.

Indeed, such clustered sets of virulence genes, termed pathogenicity islands (PAIs), have been defined for three strains of uropathogenic *E. coli*: 536 (3), J96 (39), and CFT073 (17). Typically, these sequences are large (>30-kb) blocks of DNA inserted within or near tRNA genes (12, 33, 39), contain direct

repeats and insertion sequences, have a GC content that differs from that of the rest of the genome, and encode defined virulence determinants (3, 19, 26). We have previously shown that such sequences are widespread among uropathogenic isolates (17). One probe from the PAI of strain CFT073 hybridized with genomic DNA from approximately 80% of acute pyelonephritis and cystitis strains but only 19% of fecal strains.

Previously in our laboratory the boundaries of a PAI were identified for *E. coli* CFT073, a highly virulent strain isolated from the blood and urine of a woman with acute pyelonephritis (17). In this report, we provide an analysis of the nucleotide sequence for a 57,988-bp region. Previously unrecognized open reading frames (ORFs), as well as homologs of characterized genes of other species, were found. In addition, we identified the distribution of these sequences that span the PAI among the majority of other uropathogenic strains of *E. coli*.

## MATERIALS AND METHODS

**Bacterial strains.** *E. coli* CFT073 was isolated from the blood and urine of a woman admitted to the University of Maryland Medical System for the treatment of acute pyelonephritis (27). This *hly*<sup>+</sup> *pap*<sup>+</sup> *sfa*<sup>+</sup> *pil*<sup>+</sup> strain is highly virulent in the CBA mouse model of ascending UTI (28) and is cytotoxic for cultured human renal proximal tubular epithelial cells (27). It is phenotypically positive for the production of P fimbriae, hemolysin, and type 1 fimbriae. *E. coli* DH5 $\alpha$  (34) was used as a recipient for gene bank and recombinant clones.

Four collections of *E. coli* strains were established from humans with appro-

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appropriate clinical syndromes. The first consists of 67 isolates from the urine or blood of patients (43 women and 24 men) who were admitted to the University of Maryland Medical System with acute pyelonephritis (bacteriuria of  $\geq 10^5$  CFU/ml, pyuria, fever, and no other source of infection) (27). The second collection consists of 38 isolates from the urine of women with cystitis. These isolates were kindly provided by A. Stapleton (University of Washington) and B. Foxman (University of Michigan) (10). The third collection consists of 49 isolates from the urine of 26 patients with long-term urinary catheters in place (41). Each was isolated during the first week of a new epidemiologically defined episode of *E. coli* bacteriuria. The fourth collection consists of 27 control strains of *E. coli* from the feces of healthy women (20 to 50 years old) who had not had a symptomatic UTI or known bacteriuria within the previous 6 months and who had not experienced diarrhea or received antibiotics within the preceding 1 month (28).

**Cosmid library.** A cosmid library was constructed with partially *Sau3A*-digested genomic DNA isolated from *E. coli* CFT073. DNA was ligated into *Bam*HI-digested pHC79. The ligation mixture was packaged in vitro with the Gigapack lambda packaging kit (Stratagene) and used to infect *E. coli* DH5 $\alpha$ . Transformants were selected on Luria agar containing ampicillin (200  $\mu$ g/ml).

**Preparation of templates for nucleotide sequencing.** Three overlapping cosmid clones (8-3f, 18-2f, and 5-4a), prepared from genomic DNA from *E. coli* CFT073 (28) and found previously to carry a PAI (17), were used to prepare subclones for nucleotide sequencing by deletion, subcloning of specific restriction fragments, and PCR amplification of specific sequences.

**Nucleotide sequencing and analysis.** Double-stranded DNA was used as a template for sequencing by the dideoxy-chain termination method (36). Primers used in these studies are listed in Table 1. Reactions were run with reagents from a Prism Ready Reaction Dye Deoxy Termination kit (Applied Biosystems) in conjunction with *Taq* polymerase. A model 373A DNA sequencer (Applied Biosystems) was used, and sequences were determined in both directions. DNA-sequencing software (version 2.1; Hitachi) was used for analysis of the DNA sequence for base composition, identification of ORFs and restriction sites, and other basic analyses. Apparent homologies between ORFs both outside and inside the PAI were sought in GenBank by using the Wisconsin Package (version 8.1; Genetics Computer Group, Inc.).

**DNA probes and dot blot hybridization.** DNA restriction fragments isolated from cosmid clones 8-3f and 5-4a and subclones 8HS9 and 5HS11B were used as gene probes to determine whether homologous sequences were present in genomic DNA preparations of *E. coli* strains isolated from clinical sources. *E. coli* CFT073 and DH5 $\alpha$  were used as positive and negative controls. Fragments were labeled by using the Amersham enhanced chemiluminescence system. For dot blots, *E. coli* strains were cultured in Luria broth (80  $\mu$ l) in 96-well microtiter plates. Bacterial suspensions were lysed and pipetted onto a nucleic acid transfer membrane. Samples were neutralized with Southern blot neutralization buffer (34). Hybridization was done with 11 probes encompassing the length of the PAI (Fig. 1).

Probes including the *prxA*, *modD*, *yc73*, and L8 genes were PCR amplified from cosmid clone 8-3f (17), by using primer pairs 9-14, 13-15, 16-19, and 20-21, respectively. *Sal*I-digested 8-3f and *Bam*HI/*Sma*I-digested 8HS9 (a *Sal*I/*Hind*III-digested 8-3f subclone) resulted in probes of sizes 2.6 and 3.2 kb. *Hind*III digestion of cosmid 8-3f allowed the isolation of a 3.2-kb fragment from within the hemolysin gene cluster. A *Hind*III digest of cosmid clone 5-4a (17) included fragment sizes of 7.0 and 6.0 kb. The 5.0-kb probe was obtained from a *Sma*I digest of 5HS11B (a *Hind*III/*Bam*HI-digested 5-4a subclone). Finally, the 2.1-kb probe was isolated by PCR amplification of cosmid 5-4a, by using primers 38 and 42 (Table 1). Autoradiographs were developed as described by Kafatos et al. (16). Southern blots were prepared by standard methods (34) and developed with the Amersham enhanced chemiluminescence system as specified by the manufacturer.

**Nucleotide sequence accession numbers.** The sequences of the ORFs within the boundaries of the left and right junctions of the PAI of *E. coli* CFT073 have been assigned GenBank accession no. AF081283, AF081284, AF081285, and AF081286.

## RESULTS

**Novel genes inside the PAI.** To determine whether newly described genes were present within the boundaries of the PAI, a 61-kb region was subjected to nucleotide sequencing. In this region, which included an apparent 58-kb PAI, we have identified 44 ORFs (Fig. 1). These are listed along with homologs and their accession numbers in Table 2. Four small gaps ( $\sim 2$ ,  $\sim 1$ ,  $\sim 1$ , and  $< 1$  kb) that presented difficulties in sequencing, PCR amplification, and subcloning are also identified (Fig. 1). Among the sequenced ORFs are genes that appear to be involved in iron utilization and transcriptional regulation (see below).

**Notable homologs.** Four ORFs inside the left junction (defined as the sequences associated with lower numbers on the

TABLE 1. Primers used for nucleotide sequence determination

Primer	Orientation	Coordinate (5' end)	Sequence (5'-3')
1	F	637	TGTCGGCGTTCGTTGTC
2	F	1997	TGGGCAGCAGATCGCTTGGG
3	R	2099	CTGGTGTGGGGCTATT
4	F	2740	GCCAGCGCCAGGTTACTT
5	R	3630	GCCATTAGTGGCGTCTT
6	R	4350	CCAGCCCATACGACGATA
7	F	4922	GGCAGGCTTTGCTGTTTC
8	R	5305	TGGAATGGCGGGCAGC
9	F	5891	CCGCTATCCGCTTTCACA
10	F	6314	CGTGCCGCTGTTCTGATT
11	F	7309	CTGGTGTACCCTGACTT
12	F	7818	ACGACTGCTGGGTAAAG
13	F	7863	CGTTTTTTGAGTCTCATAGA
14	R	7926	GCAACCCGACGCTCTAT
15	R	8732	GCTGTGCTCGGAAGATAT
16	F	8965	TAAATCAACGTGAGCATAA
17	R	9182	GGCTTCCCGCTCCACTCT
18	R	9324	CGATTTATGGCGTTGATT
19	R	9553	GATTTGTCAGCCTTTACCT
20	F	9716	GCTGTCGGCAATGGCGTT
21	R	10224	GGTTGGGGCAAACCAATTATG
22	F	19470	TGTTTCCCGTTGATACTA
23	R	20703	GCTGTGGGCTCGCTCCTC
24	F	28625	TCATATCTTCTCTGTCA
25	F	29644	GCCGCTCATCACTTTGTT
26	F	30351	GCTGCTATTACCTTCTTC
27	F	43124	AAATCAGCCACCACAGC
28	R	44277	TTGTGCGTGTCTTCTTCA
29	R	44841	CGTAATGACTGGGAGAGA
30	F	45318	GGCGAAGTATCCACATT
31	F	45971	GACGTTGTTGGTTGATG
32	F	46775	CGGGAGAATGAAATGAAA
33	F	47414	GAAACAACCCAGCAATA
34	R	48820	TTGTAATCCTCAGAAGA
35	R	49564	CCGCGTTGCACTTTGTTCT
36	R	50019	ATACGCCTTTTCAGATGT
37	R	50677	CAACGCCTTTTCTTAT
38	F	50866	TATCTTCTGACGCTATGC
39	F	51441	CGAAGTAAAGTGAAAG
40	F	52199	TTTCGCTAGGTATCACA
41	F	52944	TGGTCACACCGCTTTTCA
42	F	53624	CCCTGACGCTGTTGTGTG
43	F	54164	GCCGAGACAATCATCACA
44	R	55298	CGTAAGGCCAGCTGATGGTG
45	R	55755	TTTCGCGTTTCTACCACAA
46	F	56090	TGGTGATAGCGTCTGGTA
47	F	56622	GCCACTTCGACACACACC
48	F	57074	TGGCGTAAAGCGGAAAC
49	R	58040	TCAAGTCGCGTGTATGTC
50	R	58552	CTGAGCATCCGGCTAACC
51	R	58736	TCAGGCAAGCAATGTTTG
52	R	59005	CCAGCAGGAAGTTGCGGA

*E. coli* K-12 linkage map), *prxA*, *modD*, *yc73*, and *fepC*, represent an apparent iron transport system. The genes are contiguous and are predicted to be transcribed in the same direction; however, this has not been demonstrated experimentally. In other systems (25), the *modD* gene is part of a gene cluster involved in molybdenum transport, although no specific function has been ascribed to this gene. Another iron acquisition gene homolog, comprising the R4 ORF (terminology used in Fig. 1 and Table 2), appears to encode a homolog of an exogenous ferric siderophore receptor (9).

Just inside the right junction of the PAI (defined as the sequences associated with higher numbers on the *E. coli* K-12 linkage map) is an apparent antiterminator with homology to a

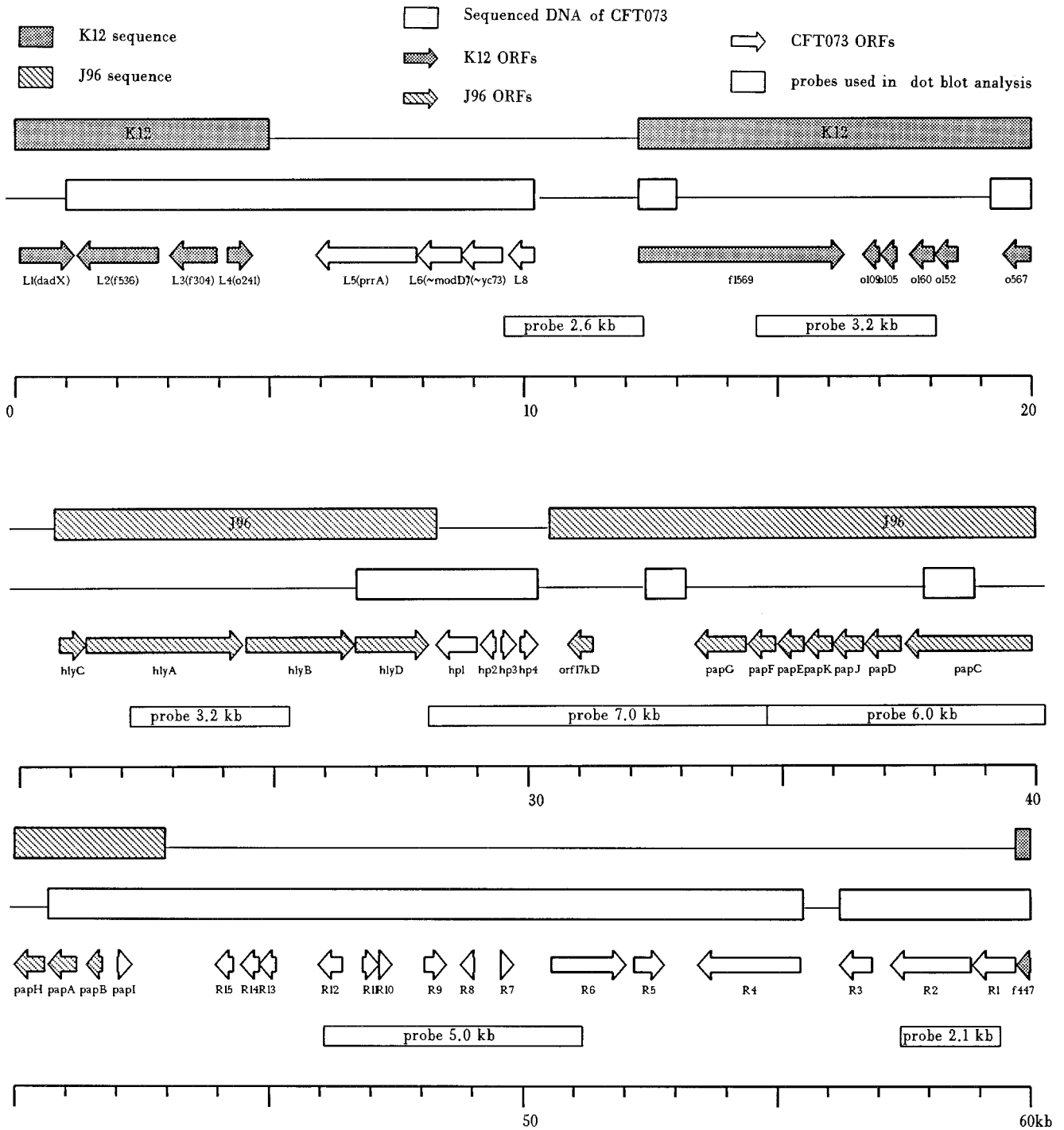


FIG. 1. Features of the pathogenicity island of *E. coli* CFT073. ORFs defined by nucleotide sequencing are shown as arrows. The ORF designations, below the arrows, are defined in Table 2. The direction of each arrow indicates the predicted direction of transcription. The top line of shaded boxes indicates that ORFs in this region are highly homologous to or identical with ORFs previously identified in *E. coli* K-12 (dark shading) (2) or uropathogenic J96 (light shading) (1, 8, 20, 21, 29, 30, 32, 40). The unshaded boxes on the second line indicate regions in which the complete nucleotide sequence was obtained in both directions. Sample sequencing was conducted only in areas not covered by unshaded boxes; these regions matched K-12 or J96 sequences with respect to restriction endonuclease sites or nucleotide sequence identity. The positions of restriction fragments or PCR products that were used as probes for DNA hybridization are shown below the ORFs. The scale at the bottom is shown in kilobases. Cosmid clone 8-3f includes ~35 kb of the PAI beginning 3 kb to the left of the depicted left junction (0 kb) and extending to the center of the *pap* operon (~39 kb). Subclone 8HS9 includes PAI sequences from ~10 to 17 kb. Cosmid clone 5-4a includes ~37 kb extending from the center of the hemolysin gene cluster (~23.5 kb) to 5 kb to the right of the right junction (58 kb). Subclone 5H11B extends from ~38 to 49 kb.

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TABLE 2. ORFs identified by nucleotide sequencing of a PAI of *E. coli* CFT073

ORF <sup>a</sup>	Coordinates <sup>b</sup>	Homolog	Description of product encoded <sup>c</sup>	Probability <sup>d</sup>	Accession no. of homolog
L1	85–1155	<i>dadX</i>	Alanine racemase	1.3e <sup>-254</sup>	P29012
L2	2822–1212 C	f536	536-aa ORF product of <i>E. coli</i>	0.0	AE000217
L3	3957–3043 C	f304	304-aa ORF product of <i>E. coli</i>	3.3e <sup>-216</sup>	AE000217
L4	4165–4668	o241	241-aa ORF product of <i>E. coli</i>	4.8e <sup>-113</sup>	AE000217
L5	7879–5909 C	<i>prnA</i>	TonB-dependent outer membrane receptor	0.0	U85771
L6	8762–7905 C	<i>modD</i>	Molybdenum transport protein	1.2e <sup>-200</sup>	U85771
L7	9571–8759 C	<i>orf2</i>	Similar to <i>H. influenzae</i> <i>yc73</i> protein	1.6e <sup>-112</sup>	U85771
L8	10203–9705 C	<i>fepC</i>	Ferric enterobactin transport ATP-binding protein	1.5e <sup>-31</sup>	F64113
Gap GL (~2 kb)					
f1569	12502–16056		1,569-aa ORF product of <i>E. coli</i>		ECAE000350
o109	16749–16420 C		109-aa ORF product of <i>E. coli</i>		ECAE000349
o105	17087–16770 C		105-aa ORF product of <i>E. coli</i>		ECAE000349
o160	17816–17334 C		160-aa ORF product of <i>E. coli</i>		ECAE000349
o152	18283–17825 C		152-aa ORF product of <i>E. coli</i>		ECAE000349
o567	20882–19181 C		567-aa ORF product of <i>E. coli</i>		ECAE000349
Gap LH (~1 kb)					
<i>hlyC</i>	21357–21869		Chromosomal hemolysin C		M10133
<i>hlyA</i>	21881–24952		Chromosomal hemolysin A		M10133
<i>hlyB</i>	25023–27146		Chromosomal hemolysin B		M10133
<i>hlyD</i>	27165–28601		Chromosomal hemolysin D	2.7e <sup>-285</sup>	Y13891
HP1	29731–28913 C		IS600 hypothetical 31-kDa protein	7.0e <sup>-195</sup>	P16940
HP2	30069–29767 C		IS600 hypothetical 11-kDa protein	2.1e <sup>-60</sup>	P16939
HP3	30172–30462		Unknown protein of plasmid Ti	2.1e <sup>-07</sup>	M25805
HP4	30547–30894		Hypothetical 15.6-kDa protein	7.7e <sup>-42</sup>	P50359
Gap HP (~1 kb)					
<i>papG</i>	34008–33001 C		P-pilus F13 tip protein		X61239
<i>papF</i>	34555–34052 C		P-pilus F13 tip protein		X61239
<i>papE</i>	35151–34630 C		P-pilus F13 tip protein		X61239
<i>papK</i>	35709–35176 C		P-pilus F13 tip protein		X61239
<i>papJ</i>	36300–35719 C		P-pilus F13		X61239
<i>papD</i>	37053–36337 C		P-pilus F13		X61239
<i>papC</i>	39649–37139 C		P-pilus F13		X61239
<i>papH</i>	40295–39708 C		P-pilus F13		X61239
<i>papA</i>	40925–40359 C		P-pilus F13		X61239
<i>papB</i>	41446–41132 C		P-pilus F13		X61239
<i>papI</i>			P-pilus F13		X61239
R15	43029–42643 C		Putative transposase of <i>E. coli</i>	1.6e <sup>-73</sup>	U06468
R14	43529–43174 C		Transposase (IS629)	8.6e <sup>-74</sup>	P16942
R13	43855–43529 C		12.7-kDa protein of <i>E. coli</i>	3.1e <sup>-51</sup>	U06468
R12	45171–44695 C		Transposase (ISAE1)	3.4e <sup>-06</sup>	A47041
R11	45575–45887		Possible precursor polypeptide	0.88	X00729
R10	45884–46144		Hypothetical 8.6-kDa protein in <i>dinG/rarB</i> 3' region	6.2e <sup>-06</sup>	P41038
R9	46789–47217	f200	200-aa ORF product of <i>E. coli</i>	3.0e <sup>-08</sup>	AE000137
R8	47771–47503 C		Neurotensin receptor	0.32	P30989
R7	48288–48534		Maltopentaose-forming amylase	0.038	D10769
R6	49286–50752		Transposase of <i>Chelatobacter heintzii</i>	3.7e <sup>-06</sup>	L49438
R5	50914–51501	<i>orfB</i>	Hypothetical protein B of <i>Bacillus</i>	1.1e <sup>-46</sup>	S23889
R4	54177–52168 C		Exogenous ferric siderophore receptor	2.2e <sup>-162</sup>	U56084
Gap GR (<1 kb)					
R3	56231–55869 C		β-Cystathionase	7.1e <sup>-100</sup>	U65013
R2	57811–56231 C	<i>malX</i>	Phosphotransferase system maltose- and glucose-specific IIabc component	3.5e <sup>-155</sup>	P19642
R1	58724–57843 C		<i>sacpa</i> operon antiterminator	2.6e <sup>-53</sup>	P26212
f447	60267–58885 C				

<sup>a</sup> For positions of ORFs, see Fig. 1.

<sup>b</sup> Base pair number beginning at left junction of PAI; C indicates that ORF is found on the complementary strand.

<sup>c</sup> Description of the best homolog. aa, amino acid.

<sup>d</sup> Probability value (*P*) of <0.05 is considered significant.

gene in the *sac* operon of *Bacillus subtilis* (11) (Fig. 1; Table 2). Based on studies with homologs (37, 38), this gene may act on the next gene, R2, which encodes a homolog of the maltose- and glucose-specific component IIa of a phosphoenolpyruvate-dependent phosphotransferase system. The adjacent gene R3

encodes a homolog of β-cystathionase (cystathionine-β lyase), the gene product of *metC*, which converts cystathione to homocysteine (4).

**Insertion sequences and transposons.** Six ORFs, designated HP1, HP2, R15, R14, R12, and R6, are related to insertion se-

TABLE 3. Strains isolated from clinical sources that hybridized with DNA probes isolated from the PAI of strain CFT073

Strain		% of isolates positive by dot blot hybridization with given probe <sup>a</sup>										
Source	No.	<i>prpA</i>	<i>modD</i>	<i>yc73</i>	<i>fepC</i>	2.6	3.2	3.2 ( <i>hly</i> )	7.0	6.0	5.0	2.1
Patient with pyelonephritis	67	87	93	90	87	84	75	49	79	81	64	79
Patient with cystitis	38	87	90	90	90	87	79	79	76	84	63	81
Patient with catheter-associated bacteriuria	49	67	69	61	61	63	65	41	57	61	41	48
Fecal sample	27	26	33	19	22	22	37	11	15	22	11	19

<sup>a</sup> Positions of probes are shown in Fig. 1.

quences and transposons which are common features of PAIs (26). The HP1 and HP2 ORFs represent the IS600 hypothetical 31- and 11-kDa proteins (24), respectively. R14 represents the transposase from insertion sequence IS629 (22). These two insertion sequences, which are members of the IS3 family and are found elsewhere in the K-12 genome (5), were originally identified in a strain of *Shigella sonnei*, a species closely related to *E. coli* (23).

**Other features of the PAI.** The PAI of strain CFT073 displays two other features common to such blocks of virulence genes. First, the GC content of the sequences, not including those of K-12 origin, is 42.9%. This value is significantly different from a value of 50.8% for the *E. coli* genome (2). Also, there appears to be segmentation with respect to unique PAI sequences and sequences of K-12 origin. Approximately 7 kb downstream of the left junction of the PAI, there is an 8-kb sequence, identical to that found in the K-12 genome, carrying six ORFs (Fig. 1). The hemolysin gene cluster *hlyCABD* follows this block.

**PAI sequences are associated with virulent strains.** To determine whether sequences found within the junctions of the PAI are common to other uropathogenic strains of *E. coli*, 11 probes were isolated along the length of the PAI (Fig. 1) and used to hybridize dot blots of genomic DNA isolated from clinical isolates (Table 3). A high percentage of isolates from patients with acute pyelonephritis or cystitis reacted with the probes (mean = 81%). These proportions were significantly higher than for strains from patients with catheter-associated bacteriuria (mean = 58%) or from fecal strains (mean = 22%) ( $P < 0.001$ ) indicating that genomic sequences homologous to those of PAI sequences from strain CFT073 are found in most other strains recovered from patients with cystitis or pyelonephritis.

**Hybridization signatures of clinical isolates.** Since uropathogenic strains clearly carry sequences that are homologous to sequences in the first CFT073 PAI, we examined which probes reacted with each isolate in the strain collection. Each strain was assigned a signature based on a positive or negative hybridization with 11 probes (Table 4). Of the 35 strains that reacted with all 11 probes, 18 were pyelonephritis-associated strains and 17 were cystitis-associated strains; no fecal strains fell into this category ( $P < 0.007$ ). The 72 strains that reacted with either 10 or all 11 probes included 39 of 67 (58%) pyelonephritis-associated strains, 23 of 38 (61%) cystitis-associated strains, 10 of 49 (20%) strains from patients with catheter-associated bacteriuria, but only 1 of 27 (4%) fecal strains ( $P < 0.0001$ ). Of the 12 strains that reacted with none of the probes, 11 were fecal strains and 1 was from a patient with catheter-associated bacteriuria. The 20 strains that reacted with either none or only 1 probe included only 1 of 67 (1%) pyelonephritis-associated strains, none of the cystitis-associated strains, 3 of 49 (6%) strains from patients with catheter-associated bacteriuria, but 16 of 27 (59%) fecal strains ( $P < 0.0001$ ). These results indicate that PAI sequences present in *E. coli* CFT073

are also found in other uropathogenic strains, especially those isolated from patients suffering with pyelonephritis and cystitis. These sequences are not generally found in fecal strains and are found less frequently in isolates from patients with catheter-associated bacteriuria.

## DISCUSSION

We have characterized a 61-kb region of DNA from *E. coli* CFT073, a pyelonephritis- and bacteremia-associated isolate, by isolation of overlapping cosmid clones, restriction endonuclease mapping, subcloning, hybridization, and nucleotide sequencing (Fig. 1). In this region, we have identified what can be defined as a PAI that includes 44 ORFs (Table 2). PAIs are typically larger than 30 kb (3), have a GC content lower than that of neighboring DNA (19), and have gene clusters posi-

TABLE 4. Hybridization signatures for clinical strains

Signature <sup>a</sup>	No. of positive probes	No. of isolates from:				Total no. of strains
		Patients with pyelonephritis	Patients with cystitis	Patients with catheter-related bacteriuria	Fecal sample	
11111111111	11	18	17	1	0	36
11111101111	10	8	1	7	0	16
11111011111		5	0	3	1	9
11111111101		4	3	1	0	8
11111111110		3	2	0	0	5
11111101110	9	2	1	3	1	7
11111001111		1	0	3	0	4
11111101101		2	0	1	0	3
11111100001	7	2	0	1	0	3
11111000001	6	1	0	5	1	7
11111100000		2	1	0	0	3
10000111110		0	1	2	0	3
11111000000	5	1	0	3	0	4
00000101100	3	0	0	3	1	4
00000001100	2	0	0	4	0	4
00000100000	1	1	0	2	3	6
00000000000	0	0	0	1	11	12

<sup>a</sup> Signatures (denoting positive [1] or negative [0] hybridization with 11 probes) represented by only one or two strains are as follows: for patients with pyelonephritis, 11111111100, 11111001101, 11110011110, 01111011001, 01111000111, 11110101100, 01110001111, 01001101101, 11101000101, 11110001100, 00000111110, 10000100011, 01010000001, 10000000011, 01110000000, 00000100001; for patients with cystitis, 11111010111, 11111011101, 11111110001, 1111100101, 01111011101, 11111010001, 11111010000, 11111000100, 00000111111, 01110000001, 00000000101, 00000000001; for patients with catheter-associated bacteriuria, 11111100111, 01011011111, 11111101100, 11111101001, 1111100101, 11111101000, 11110001110, 0111100001, 11011010100, 10000101110, 01101000011, 11110000001, 01000101100, 10010000000, 10000000100, 01000100000, 00000010000, 10000000000; fecal samples, 11111100111, 01111101111, 10000101110, 11111010000, 11001000001, 01000010100, 01000010000, 00000001000.

tioned near each other which contribute to a single virulence property (26). The first PAI of CFT073 is 58 kb in size (Fig. 1), has a GC content of 42.9% (compared to 50.8% in K-12 genomic DNA), and includes the genes encoding HlyA hemolysin and P fimbriae.

Distinct segmentation and insertion sequences are also common features of PAIs, which suggests that DNA rearrangements mediated by illegitimate or *recA*-dependent recombination have occurred or that DNA has been inserted by transposons, phage, or integrons (19, 26). This 61-kb region shows some evidence of rearrangement with the K-12 genome, as two blocks of four and six ORFs within this region closely match entries in the complete K-12 genome sequence (Fig. 1, darkly shaded boxes). Another four genes, in the *hylCABD* cluster, encode the HlyA hemolysin and are nearly identical to sequences from *E. coli* J96, a notable uropathogenic strain. Also carried on this stretch of DNA is the *pap* operon, comprising of 11 ORFs (*papIBAHCDJKEFG*) encoding one of the two P fimbriae expressed by this strain (28); this *pap* operon encodes a class II PapG adhesin (data not shown). Insertion elements and transposases occur six times in the region, perhaps predisposing these sequences to rearrangement. Remaining are 19 ORFs, which are also listed in Table 2, beginning at the left junction of the PAI and moving toward the right junction.

Nucleotide sequences of the first 58-kb PAI of CFT073 reveal newly described genes. These gene sequences, as assayed by DNA hybridization, are present in virulent uropathogenic strains and are generally absent in nonvirulent strains. Therefore, we postulate that these newly described genes may represent virulence determinants that contribute to the pathogenesis of cystitis and acute pyelonephritis caused by uropathogenic *E. coli*. As we continue our studies on the pathogenicity islands of CFT073, we will select mutants with phenotypes that may relate to virulence such as iron uptake, metabolite uptake, and transcriptional regulation. We will undertake allelic exchange mutagenesis of specific genes and test these mutants by *in vitro* assays and, *in vivo*, by using the CBA mouse model of ascending UTI (13). By creating such mutants, we hope to determine which genes contribute to the virulence phenotype of uropathogenic *E. coli*.

It is likely, however, that strain CFT073 contains another PAI. For two other uropathogenic strains that have been studied closely, 536 and J96, each strain was found to contain two separate PAIs. For strain 536, PAIs of 190 and 70 kb are inserted at 97 min (within *leuX*) and 82 min (within *selC*), respectively (reviewed in reference 19). For strain J96, PAIs of 110 and >170 kb are inserted at 94 min (within *pheR*) and 64 min (within *pheV*), respectively (39). Because we know that strain CFT073 contains two complete *pap* operons encoding distinct P fimbriae (28) and that only one operon is found in the PAI described in this report (17), it is likely that these sequences are found in a separate PAI along with the genes encoding F1C fimbriae (*foc* [unpublished observation]). That CFT073 has another PAI of unknown size raises the possibility that a significant number of virulence genes, not detected on the PAI described here, also contribute to the virulence of this strain.

Finally, an important question raised by these studies is whether there are distinctions between strains isolated from patients with cystitis and strains from patients with pyelonephritis. Based on hybridization of chromosomal DNA with PAI probes, we are unable to make such a distinction. With the exception of the *hly* hemolysin-containing probe, there were no significant differences ( $P > 0.2$ ) between the percentages of cystitis and pyelonephritis isolates that reacted with each of the

11 PAI probes (Table 3). In contrast, other hybridization studies which used specific probes (e.g., *pap*, *hly*, *sfa*, and *foc*) have revealed that a higher percentage of pyelonephritis-associated isolates than of cystitis-associated isolates reacted with these probes (6). It is now clear that uropathogenic isolates generally express adhesins that aid in colonization and toxicity (hemolysin and cytotoxic necrotizing factor), but this group of strains clearly does not have a single phenotype (6). It will be interesting to see whether cystitis and pyelonephritis strains are, in the future, delineated on the basis of specific genotypes.

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