

Genetic Characterization of *Escherichia coli* Type 1 Pilus Adhesin Mutants and Identification of a Novel Binding Phenotype

TERRI S. HAMRICK, SANDRA L. HARRIS, PATRICIA A. SPEARS, EDWARD A. HAVELL,
JOHN R. HORTON, PERRY W. RUSSELL, AND PAUL E. ORNDORFF*

Department of Microbiology, Pathology and Parasitology, College of Veterinary Medicine,
North Carolina State University, Raleigh, North Carolina 27606

Received 22 February 2000/Accepted 1 May 2000

Five *Escherichia coli* type 1 pilus mutants that had point mutations in *fimH*, the gene encoding the type 1 pilus adhesin FimH, were characterized. FimH is a minor component of type 1 pili that is required for the pili to bind and agglutinate guinea pig erythrocytes in a mannose-inhibitable manner. Point mutations were located by DNA sequencing and deletion mapping. All mutations mapped within the signal sequence or in the first 28% of the predicted mature protein. All mutations were missense mutations except for one, a frameshift lesion that was predicted to cause the loss of approximately 60% of the mature FimH protein. Bacterial agglutination tests with polyclonal antiserum raised to a LacZ-FimH fusion protein failed to confirm that parental amounts of FimH cross-reacting material were expressed in four of the five mutants. The remaining mutant, a temperature-sensitive (*ts*) *fimH* mutant that agglutinated guinea pig erythrocytes after growth at 31°C but not at 42°C, reacted with antiserum at both temperatures in a manner similar to the parent. Consequently, this mutant was chosen for further study. Temperature shift experiments revealed that new FimH biosynthesis was required for the phenotypic change. Guinea pig erythrocyte and mouse macrophage binding experiments using the *ts* mutant grown at the restrictive and permissive temperatures revealed that whereas erythrocyte binding was reduced to a level comparable to that of a *fimH* insertion mutant at the restrictive temperature, mouse peritoneal macrophages were bound with parental efficiency at both the permissive and restrictive temperatures. Also, macrophage binding by the *ts* mutant was insensitive to mannose inhibition after growth at 42°C but sensitive after growth at 31°C. The *ts* mutant thus binds macrophages with one receptor specificity at 31°C and another at 42°C.

Type 1 pili are filamentous proteinaceous appendages produced by several members of the *Enterobacteriaceae*. In *Escherichia coli*, type 1 pili have been studied extensively with regard to their genetics, biosynthesis, and ability to bind mannose-containing receptor molecules on a variety of eucaryotic cells (reviewed in reference 29). Although the pili are made principally of a single protein monomer, the product of the *fimA* gene, several minor protein components are also incorporated (12, 34). These are most often found at the ends of pili and are organized into fibrillar structures (15). One of the minor components, the product of the *fimH* gene (FimH), binds directly to the receptor (18). Whereas the specificity of the interaction of the *fimH* product can be influenced by other fimbrial components (23), several studies have linked certain naturally occurring *fimH* allelic types to the specificity of receptor binding (43, 46) and the strength of receptor binding (45). Additional experiments have suggested that some *fimH* allelic differences can contribute to tissue tropism (35, 43, 44). Gene fusion experiments have indicated that FimH binding capacity resides in the amino one-third to one-half of the protein (17, 48). However, point mutations in various parts of the coding region can effect a change in specificity for particular types of ligands (46). This is consistent with what is found with other bacterial adhesins (4). FimH has been used to express foreign antigens by inserting heterologous gene segments into the *fimH* gene (33) and used on its own as an effective

immunogen in preventing experimental urinary tract infections in mice (20).

We have been particularly interested in the factors influencing the specificity of FimH binding and how FimH affects proper pilus structure (reviewed in 29). Some years ago, we isolated a number of *fimH* point mutants (13). These mutants were isolated following enrichment for individuals that formed pellicles when grown in static broth (a property associated with FimH) in the presence of a mannose analogue (α -methylmannoside) that normally inhibits pellicle formation. All mutants isolated in this manner were defective in their ability to bind guinea pig erythrocytes. Additionally, several of the mutants produced pili with altered morphology.

In this report, we further characterize five of the *fimH* mutants isolated in the previous study. These five mutants represent all allelic classes of the 11 mutants initially isolated (13). In particular, we concentrate upon one of the mutants that was conditionally defective in erythrocyte agglutination, showing that erythrocyte and macrophage binding are differentially affected by the mutation. The results indicate that FimH-mediated attachment to different types of eucaryotic cells can occur through different mechanisms.

MATERIALS AND METHODS

Bacteria, bacteriophage, plasmids, and media. The bacterial strains, all *E. coli* K-12 derivatives, bacteriophage, and plasmids used in this investigation are listed in Table 1. Media consisted of L agar and L broth (28), λ -broth (28), tetrastazolium agar (42), and MinA broth and agar (28). Antibiotics were added as previously described (30) unless otherwise noted.

Genetic techniques. Transformation with plasmid DNA followed the method described by Lederberg and Cohen (21). P1 transduction followed the method described by Miller (28).

Recombinant DNA techniques. Restriction endonuclease cleavage, plasmid and chromosomal DNA isolation, fragment purification, end filling, ligation, and

* Corresponding author. Mailing address: College of Veterinary Medicine, 4700 Hillsborough Street, Raleigh, NC 27606. Phone: (919) 513-6207. Fax: (919) 513-6455. E-mail: Paul_Orndorff@ncsu.edu.

TABLE 1. *E. coli* strains, bacteriophage, and plasmids used in this study

Strain, phage, or plasmid	Description	Source or reference
<i>E. coli</i>		
JM101	$\Delta(lac-proAB) supE thi/F' lacI^q\Delta M15 traD36 proAB$	27
EC901	<i>leu-6 argE3 proA2 lacY1 his-4 thi-1 galK2 ara-14 xyl-5 srl-31 hsdR4 recA13 srl::Tn10</i>	38
ORN103	<i>thr-1 leu-6 thi-1 $\Delta(argF-lac)U169 xyl-7 ara-13 mtl-2 gal-6 rspL tonA2 minA minB$</i> $\Delta(fimEACDFGH)$	31
ORN172	<i>thr-1 leuB thi-1 $\Delta(argF-lac)U169 xyl-7 ara-13 mtl-2 gal-6 rpsL tonA2 supE44$</i> $\Delta(fimBEACDFGH)::kan pilG1$	50
ORN206	JM101 except $\Delta(fimBEACDFGH)$ and <i>recA13 Kan^r</i>	P1 transduction from ORN172, then EC901, then ORN103
ORN155	ORN115 except (Tn5 inserted adjacent [3'] to <i>fimH</i>)	13
ORN160	ORN155 except <i>fimH241</i>	13
ORN162	ORN155 except <i>fimH204</i> (has identical lesion to <i>fimH236</i>)	13
ORN163	ORN155 except <i>fimH218</i> (has identical lesion to <i>fimH236</i>)	13
ORN164	ORN155 except <i>fimH236</i>	13
ORN165	ORN155 except <i>fimH244</i>	13
ORN157	ORN155 except <i>fimH205(ts)</i>	13
ORN158	ORN155 except <i>fimH208</i>	13
ORN115	<i>thr-1 leuB thi-1 $\Delta(argF-lac)U169 malA1 xyl-7 ara-13 mtl-2 gal-6 rpsL fhuA2 supE44$</i> <i>pilG1</i>	47
ORN133	Same as ORN115 except <i>fimH'-kan Mal⁻</i>	26
ORN183	Same as ORN157 except <i>Mal⁺</i>	13
ORN175	Same as ORN115 except <i>Mal⁺</i>	16
ORN204	Same as ORN133 except <i>Mal⁺</i>	11
Bacteriophage P1	<i>vir</i>	Laboratory collection
Plasmids		
pBR322	ColE1, Ap ^r Tc ^r	3
pACYC184	P15A, Cm ^r Tc ^r	5
pUR288	<i>lacZ</i> fusion vector Ap ^r	37
pSH2	pACYC184 <i>fimBEAICDFGH</i> Cm ^r	10
pORN118	pSH2 except <i>PstI</i> site in <i>fimH</i> changed to <i>XhoI</i> Cm ^r	25
pORN123	pBR322 $\Delta PvuII$ Ap ^r	32
pORN148	pORN123 <i>fimD' fimF fimG fimH</i> Ap ^r	36
pORN303	pUR288 with <i>lacZ-fimH</i> fusion	This study
pORN304	Bal31 <i>fimH</i> deletion mutant of pORN118; terminal 22 codons removed	This study
pORN305	Bal31 <i>fimH</i> deletion mutant of pORN118; terminal 82 codons removed	This study
pORN306 ^a	<i>KpnI-SalI</i> deletion mutant of pSH2; terminal 276 codons of <i>fimH</i> removed	This study
pORN307 ^b	<i>PvuII-SalI</i> deletion mutant of pSH2; entire <i>fimH</i> gene deleted	This study
pORN142	ColE1, <i>fimH</i> with adjacent Tn5 insertion	13
pORN241	pORN142 except <i>fimH241</i> allele	13
pORN236	pORN142 except <i>fimH236</i> allele	13
pORN204	pORN142 except <i>fimH204</i> allele	13
pORN218	pORN142 except <i>fimH218</i> allele	13
pORN244	pORN142 except <i>fimH244</i> allele	13
pORN205	pORN142 except <i>fimH205</i> allele	13
pORN208	pORN142 except <i>fimH208</i> allele	13

^a The *KpnI* site in *fimH* was first converted to an *XhoI* site by partial digestion of the pSH2 plasmid with *KpnI* (pSH2 has two *KpnI* sites), followed by end filling with DNA polymerase I and *XhoI* linker ligation as described by Orndorff and Falkow (31).

^b The *PvuII* site in *fimH* was first converted to an *XhoI* site by partial digestion of the pSH2 plasmid with *PvuII* (pSH2 has five *PvuII* sites) and ligation of *XhoI* linkers as described by Orndorff and Falkow (31).

subcloning were performed as previously described (36). DNA sequencing was performed on double-stranded plasmid DNA as previously described (50) using 20-oligonucleotide primer pairs bracketing various regions of the *fimH* and *lacZ* genes. PCR product sequencing of chromosomal and plasmid DNA was carried out as described by Russell and Orndorff (36). Bal31 exonuclease digestions were performed as described by Maniatis et al. (24).

Construction and induction of the *lacZ-fimH* fusion, and isolation of the fusion protein. A *lacZ-fimH* fusion was created by *KpnI-PstI* digestion of pORN148 followed by end filling and ligation to *Bam*HI-digested, end-filled plasmid pUR288. This created plasmid pORN303. The fusion product contained approximately 92% of the mature FimH product. Pilot experiments revealed that the most efficient production of fusion protein was obtained by diluting an overnight culture of the strain containing the fusion plasmid (ORN206/pORN303) 1:50 in fresh warm L-broth containing ampicillin (100 μ g/ml) and 5 mM isopropyl- β -D-thiogalactopyranoside (IPTG), followed by 4 h of growth with shaking at 37°C. After this period, bacteria were harvested by centrifugation (7,500 \times g for 10 min), and the pellet was resuspended in 1 \times protein sample

buffer (19) and boiled for 5 min. After a brief centrifugation to remove the insoluble material, various amounts of the supernatant were subjected to electrophoresis on a 10% discontinuous polyacrylamide gel (19). The fusion protein band was detected following Coomassie brilliant blue staining at the position expected from the predicted size of the fusion protein induced by IPTG. For preparative gels (prepared as above except 1.5-mm thick), the approximate location and amount of the fusion protein band were determined by staining marker strips containing high-molecular-weight protein standards (Bio-Rad) with Coomassie brilliant blue stain.

Immunological methods. Rabbit polyclonal antiserum against the LacZ-FimH fusion protein was raised by injecting a New Zealand White rabbit (ca. 2 kg) with a macerated acrylamide gel slice that contained approximately 100 μ g of a LacZ-FimH fusion protein in complete Freund's adjuvant. Booster doses of 100 μ g of fusion protein in incomplete Freund's adjuvant were administered approximately every 2 weeks with blood drawn beginning after the fourth boost and every 2 weeks thereafter, for a total of five bleedings. Antiserum was stored at

-20°C, and aliquots were ammonium sulfate precipitated (7) and concentrated approximately fivefold in phosphate-buffered saline (PBS) prior to use.

Immunological and functional detection of FimH. The presence of FimH was determined immunologically by bacterial agglutination reactions performed in 96-well round-bottomed microtiter plates. Bacteria from overnight cultures were isolated by brief centrifugation (1 to 2 min in a microcentrifuge) and concentrated two- to fivefold in PBS (approximately 4×10^9 to 10×10^9 cells per ml) depending upon the experiment. Twenty-five microliters of antiserum was serially twofold diluted in microtiter wells, and 25 μ l of the bacterial suspension was added to each well. Microtiter plates were incubated at room temperature for 1 h and then refrigerated (4°C) until the negative control wells (containing a *fimH* insertion mutant) had settled (typically 24 to 48 h).

FimH function was assayed by the ability of *E. coli* to agglutinate guinea pig erythrocytes. Agglutination tests were conducted in 96-well round-bottomed microtiter plates in which overnight cultures, concentrated twofold, were serially twofold diluted and the contents of each well were mixed by adding 25 μ l of a 4% suspension of fresh guinea pig erythrocytes. Incubation proceeded as above until erythrocytes in the negative control wells (containing the *fimH* insertion mutant as above) had settled (typically 15 to 24 h).

Recombination mapping of *fimH* mutations. Transformants that contained a chromosomal *fimH* mutant allele received plasmids containing deletion derivatives of *fimH* via transduction. Transductants (800 to 1,000 colonies) were recovered from agar plates using a cotton swab. The material from each swab was expressed into approximately 1.0 ml of PBS, and 0.3 ml was added to a microcentrifuge tube containing 100 μ l of settled fresh guinea pig erythrocytes. The bacteria and erythrocytes were mixed by inversion and incubated (to allow binding) for 10 min. The red blood cells were isolated by centrifugation for approximately 1 s in a microcentrifuge, and the supernatant was aspirated. The pellets were resuspended and washed with 1.0 ml of PBS five more times. The final pellet was resuspended in 0.5 ml of distilled water (to lyse the erythrocytes), and 2.0 ml of L-broth with chloramphenicol was added. This mixture was incubated overnight with shaking at 37°C to expand the population that remained bound to the erythrocytes. The following day, red blood cell debris was removed from 1.0 ml of the culture by a 2-s microcentrifugation step, and the bacteria in the supernatant were subsequently isolated by centrifugation for 2 min. The isolated bacteria were washed in 1.0 ml of PBS and resuspended in 0.75 ml of PBS. Fifty microliters of settled guinea pig erythrocytes was then added. The subsequent incubation, washing, and enrichment were repeated an additional two times. After the last outgrowth, cultures were streaked onto L-agar plates, and 20 individual colonies were scored for their ability to agglutinate erythrocytes.

Erythrocyte binding assay. Overnight cultures of *fimH* point mutants and positive and negative control strains (described below) were harvested by microcentrifugation, resuspended in PBS, and mixed 1:1 by volume to give a final concentration of approximately 5×10^6 cells/ml. Sixty microliters of each mixture to be tested was added to an Eppendorf tube. Ten microliters was removed, diluted, and titered on maltose-tetrazolium plates to obtain the ratio of the two strains. To the bacteria remaining in the tube, 0.1 ml of settled guinea pig erythrocytes was added. The resulting suspension was then gently mixed and incubated at room temperature for 10 min. Adherent bacteria were removed by a brief (approximately 3 s) centrifugation to pellet erythrocytes and erythrocyte-bound bacteria. A portion of the supernatant was diluted and plated, and the resulting ratio was compared to the starting ratio.

In each assay, a *fimH* point mutant was mixed with a *fimH* insertion mutant (strain ORN204). The insertion mutant provided a negative control within each assay. To ensure that the changes in ratio accurately reflected differences in binding ability, pilot experiments included mixtures of two parental strains and two *fimH* insertion mutants. The strain combinations used in this case were ORN115 with ORN175 and ORN133 with ORN204. These strains were also mixed in parent-insertion mutant pairs. The strains in all mixtures were distinguished by their different maltose utilization phenotype on maltose-tetrazolium agar (refer to Table 1). Normalization of mutant binding values was done in part to reduce the effects of artificial variability between assays (e.g., erythrocyte age and concentration).

Macrophage binding assay. Resident (unelicited) peritoneal macrophages from male BALB/c mice 8 to 12 weeks of age were used in these experiments. Macrophages were harvested, delivered into 48-well cluster culture plates, and incubated overnight as described previously (11). Macrophages (approximately 1×10^5 to 2×10^5 cells/well in 0.5 ml of tissue culture medium) were exposed to approximately 10^6 *E. coli* (grown overnight in λ -broth, and harvested and mixed pairwise as described [11]) that were added in 25 μ l of PBS for 10 min at 37°C. After incubation, wells were washed four times (each wash was with 0.5 ml of PBS). After the final wash, 0.5 ml of PBS containing 0.1% Triton X-100 was added to each well to lyse the macrophages. Approximately 5 min after the Triton X-100 additions, the contents of the wells were diluted and plated. The exposure of bacteria to Triton X-100 had no effect on bacterial viability. Control wells that contained no macrophages were used to assess nonspecific binding of bacteria. In no instance was the level of binding appreciable (>10% of that of macrophage-containing wells). When α -methyl-mannoside (α mm) was added to inhibit *E. coli* binding, a small volume of a 1.0 M α mm stock solution (in PBS) was added to a final concentration of 50 mM. Control wells had an equivalent

volume of PBS added. PBS solutions used to remove unbound *E. coli* also contained 50 mM α mm.

In each assay (typically performed at least in duplicate), the strain mixtures were the same as those indicated for the erythrocyte binding assay just described. That is, binding efficiency was determined relative to a *fimH* insertion mutant. In order to assess the degree of macrophage binding inhibition effected by α mm, comparisons between parent and point mutant were normalized to a *fimH* insertion mutant (whose binding is insensitive to α mm) by using mixtures containing the strain to be assayed and an insertion mutant. This normalization controlled for well-to-well variation in macrophage number and allowed internal calibration of the degree of α mm binding inhibition.

Statistical and DNA sequence analysis methods. Standard deviation of the mean was calculated with the Microsoft Excel STDEV function. Standard error was calculated as the standard deviation divided by the square root of the number of experiments. The significance of mean differences was determined by Student's *t* test. Both tests were provided by the Microsoft Excel version 4 statistics package. Statistically significant differences were defined as a *P* of <0.05. Genetics Computer Group (version 7) SIGCLEAVE analysis using the cleavage rules of von Heijne (49) was used to assess possible alterations in the FimH signal cleavage site due to signal sequence mutations in *fimH*.

Nucleotide sequence accession numbers. The sequences of the five representative *fimH* mutant alleles have been deposited in GenBank with accession numbers as follows: *fimH241*, AF154925; *fimH236*, AF154926; *fimH244*, AF154927; *fimH205*, AF154928; and *fimH208*, AF154929.

RESULTS

Mapping point lesions in *fimH*. DNA sequencing of the plasmid-borne *fimH* alleles used to create the chromosomal mutants described by Harris et al. (13) provided the sequence of both strands of the entire *fimH* gene. The sequence of 11 alleles revealed that 5 were unique, each having a single lesion at the sites marked in Fig. 1. (The area sequenced included the *fimH* coding region plus a minimum of 20 bp on either side.) Sequencing of PCR amplicons from the chromosomal *fimH* alleles in all 11 strains revealed that they contained the same lesion as the plasmid-borne allele. (The entire gene was not sequenced in these cases [data not shown].) As a separate test that the lesions sequenced were responsible for the hemagglutination-negative phenotype, recombination mapping was carried out. This mapping involved enrichment for hemagglutination-positive individuals that were produced as a result of recombination between a chromosomal *fimH* allele having one of the point mutations and a set of in vitro-generated deletion derivatives of *fimH* residing on plasmids. The results of this mapping procedure (Fig. 2) were in good agreement with the lesion location indicated by DNA sequencing. Also, since the mutant alleles were not resequenced in their entirety after their introduction into the chromosome via homologous recombination (13), this mapping procedure indicated that unappreciated sequence differences in or around the chromosomal *fimH* gene were not responsible for the mutant phenotype.

Variety of phenotypes displayed by *fimH* point mutants. Whereas all *fimH* mutants failed to agglutinate guinea pig erythrocytes, certain *fimH* point mutants displayed additional properties (summarized in Table 2). Two of these properties involved the generation of aberrant pilus morphologies (class II and class III), and a third involved a conditional agglutination phenotype, all of which were initially noted by Harris et al. (13). Class II mutants (represented here by strain ORN164) had longer than normal pili. Class III mutants (represented here by strain ORN158) had much longer and very sparse pili. The lesion associated with the class II phenotype predicted a change in the last amino acid at the signal sequence site from serine to leucine. The class III mutant had a frameshift lesion predicted to generate a truncated FimH product approximately 40% of the normal size. In addition to the aberrantly fimbriated mutants, a mutant with a conditional (temperature sensitive [ts]) erythrocyte agglutination phenotype was isolated (Table 2). This mutant (strain ORN157, carrying the *fimH205*

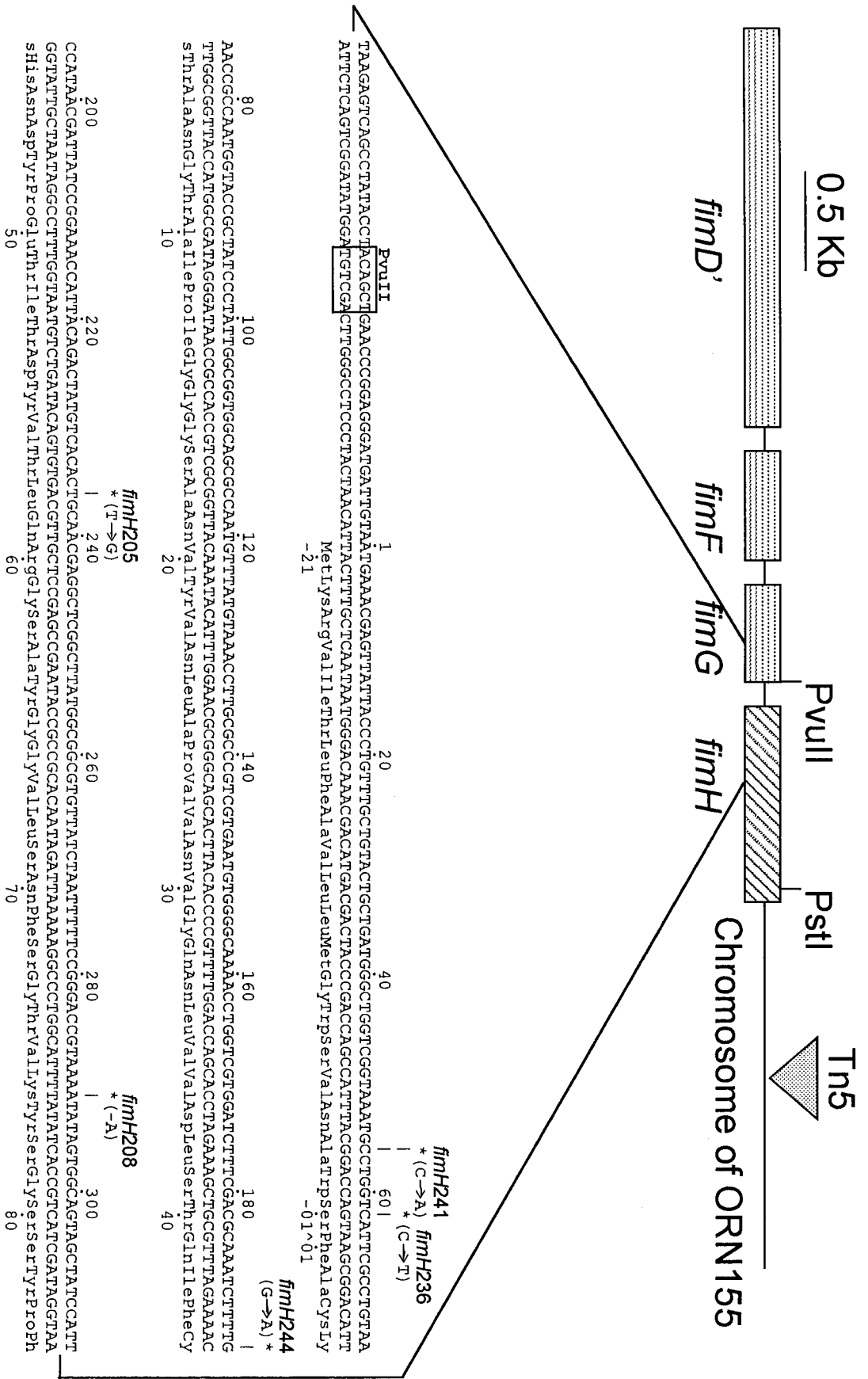


FIG. 1. Diagram of the *fimH* gene (diagonally striped rectangle) in the chromosome of strain ORN155. The entire *fim* gene cluster (*fimBEACDEFGH*) is located at approximately 98 min on the *E. coli* genetic map (more precise coordinates can be obtained from Genbank, accession no. AE000502, and reference 2). All genes are transcribed left to right. The Tn5 insertion adjacent to *fimH* is not drawn to scale. Approximately one-third of the 5' end of the *fimH* gene is expanded to show the sequence. Nucleotide numbering follows the convention in Genbank. Amino acids are numbered, with positive numbers denoting residues in the mature protein. Negative numbers indicate the amino acids in the signal sequence. The caret (^) below the sequence denotes the signal processing site (12). Mutation sites are designated by asterisks (*) and allele number. Arrows show the base change involved. A minus sign indicates a deletion. Amino acid changes accompanying the mutations and the phenotypes conferred by the lesions are summarized in Table 2.

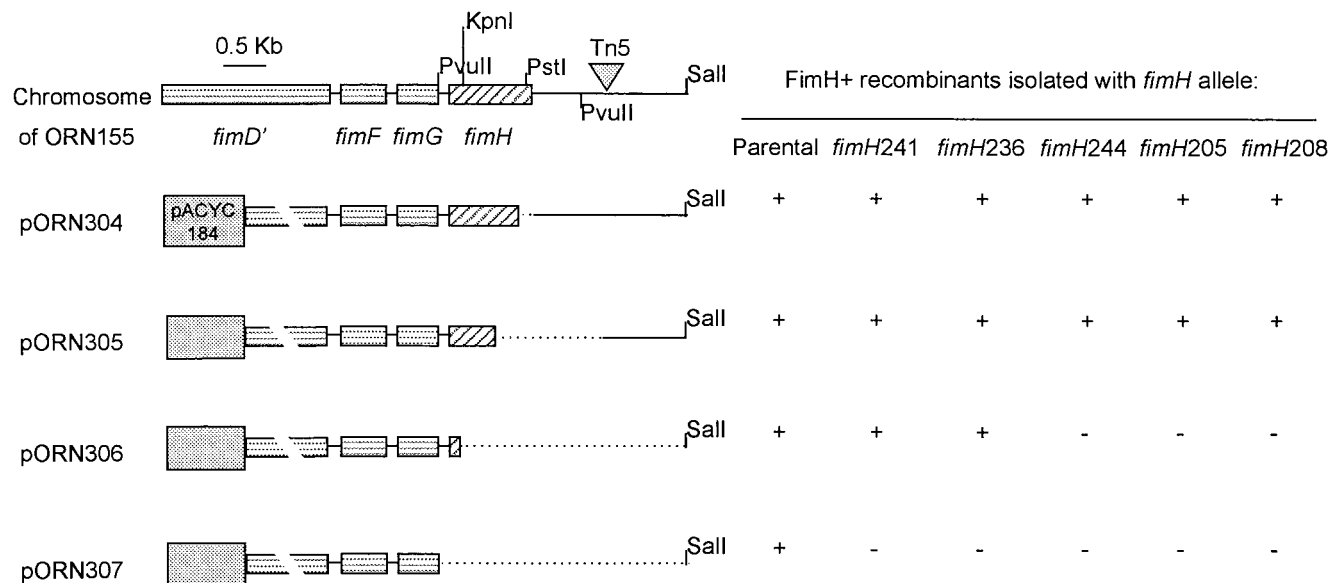


FIG. 2. Mapping of mutations shown in Fig. 1 by recombination. Mutants with chromosomal *fimH* alleles shown on the right side of the figure were incubated with the plasmids shown at the left. Recombinants capable of hemagglutination were recovered following enrichment as described in the text. The *fimH* gene is denoted by the diagonally striped rectangle. The dotted line indicates the region deleted. Specific deletions (codons deleted) from the 3' end of FimH are listed in Table 1. Deletions were created by removing either the restriction fragments indicated or by using Bal31 exonuclease digestions starting from the *XhoI* site in pORN118 as described in the text.

allele) had morphologically similar pili at both the permissive and restrictive temperatures but failed to agglutinate erythrocytes at the restrictive temperature (13).

Is the defect in the *fimH* products one of localization or function? In the initial characterization of *fimH* point mutants (13), it was inferred that the *fimH* product was being expressed and properly localized because the mutants formed pellicles in static broth, a property eliminated by *fimH* insertion mutations (13). However, in these tests, we could not conclude that parental amounts of mutant FimH were being expressed because

the level of FimH expression needed for pellicle formation could have been lower than that needed for erythrocyte agglutination.

In order to provide a measurement of the *fimH* product that was not linked to function, we produced and employed polyclonal rabbit antiserum raised against a *lacZ-fimH* translational fusion. The reactivity of four of the five mutants was distinguishably less than that of the parental strain (data not shown). Only the ts mutant having the *fimH205* allele showed reactivity indistinguishable from the parent's (see next sec-

TABLE 2. Properties of *fimH* mutants

Lesion location ^a (nt)	No. of isolates ^b	Morphologic class ^c	Representative <i>fimH</i> allele (strain no.)	Type of lesion, amino acid change if missense (amino acid position) ^d	Phenotype or other relevant properties
56	5	I	<i>fimH241</i> (ORN160)	Missense, Ala (-3)→asp	Amino acid change in signal sequence near processing site
62	3 ^e	II	<i>fimH236</i> (ORN164)	Missense, Ser (-1)→Leu	Amino acid change in signal sequence at processing site
194	1	I	<i>fimH244</i> (ORN165)	Missense, Cys (44)→Tyr	Amino acid change in first of four cysteine residues
236	1	I	<i>fimH205</i> (ORN157)	Missense, Leu (58)→Arg	Amino acid change produces a conditional erythrocyte agglutination phenotype
291	1 ^f	III	<i>fimH208</i> (ORN158)	Frameshift, deletion of 1 nucleotide	Frameshift lesion predicts a truncated protein approximately 40% of normal size

^a Nucleotide (nt) residue altered. Refer to Fig. 1 for coordinates. The parental *fimH* allele is identical to that of the fully sequenced *E. coli* K-12 strain MG1655, referenced in the text.

^b Number of isolates refers to the number of independently isolated *fimH* mutants having the same mutation by DNA sequencing and recombination analysis. Five identical mutant alleles were found at position 56 (*fimH241*, *fimH229*, *fimH222*, *fimH247*, and *fimH246* [13]). Three identical alleles were found at position 62 (*fimH336*, *fimH204*, and *fimH218* [13]).

^c Morphologic class refers to the classes observed by Harris et al. (13). Class I mutants have normal-appearing pili, class II mutants have longer than normal pili, and class III mutants have fewer and longer pili. See text for additional details.

^d See Fig. 1 for base pair changes, lesion location, and amino acid position.

^e Originally, only one mutant having the representative allele (*fimH236*) was scored as being a class II mutant (13). Retrospective analysis of available electron micrographs revealed that at least one of the two additional mutants was simply overlooked in the original screen (data not shown).

^f In addition to strain ORN158, one additional mutant was scored as being a class III mutant in our original phenotypic screening (13). Upon examination of Southern blots using DNA encoding *fimH* as a probe, and PCR sequencing of the chromosomal *fimH* allele in this mutant, it was concluded that there had been an aberrant introduction of this allele into the chromosome (our unpublished observations).

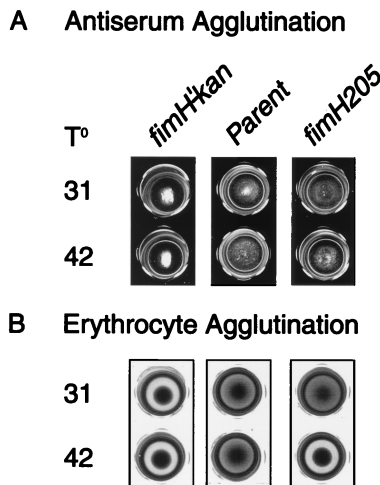


FIG. 3. Microtiter agglutination reactions of strain ORN157 bearing the *fimH205* allele compared to the parental strain (ORN155) and strain ORN133, a *fimH* insertion mutant (*fimH'*-kan). (A) Bacterial agglutination of mutants grown at the temperatures indicated, 31 or 42°C, concentrated twofold from overnight cultures, and incubated with FimH antiserum (1:4 dilution) as described in the text. (B) Microtiter guinea pig erythrocyte agglutination by the same strains obtained under the same growth condition. Incubation conditions are described in the text.

tion). The decreased reactivity of the four mutants made further functional comparisons to the parent problematical, because we could not conclusively state the reason for the reduced antiserum reactivity (e.g., poor cross-reactivity, reduced FimH expression, or exposure in the pilus fiber). For these reasons, further characterization of the binding properties of these mutants was curtailed, and results focused on the *fimH205* mutant.

FimH expression in the *ts* mutant at the restrictive and permissive temperatures. Bacterial agglutination of ORN157 (carrying the *ts* *fimH205* allele) using dilutions of FimH-specific antiserum revealed no noticeable difference between the parent and mutant in terms of antibody reactivity after growth at both the permissive and restrictive temperatures (summarized in Fig. 3A). In contrast, erythrocyte agglutination effected by the mutant was decidedly reduced (Fig. 3B).

Role of temperature in FimH-mediated erythrocyte agglutination in *fimH205* mutants. Two ways that a temperature shift could effect a change in FimH function are (i) via a spontaneous (instantaneous) conformational change in existing FimH molecules and (ii) via a conformational change in nascent or newly synthesized *fimH* product (requiring time for new synthesis). We tested these two possibilities by examining the kinetics with which the hemagglutination phenotype (Hag) changed after a temperature shift in the presence and absence of sufficient chloramphenicol to halt protein synthesis. Our results (Fig. 4) revealed that the change from Hag⁺ to Hag⁻ (and vice versa) was not instantaneous and required new protein synthesis. Thus, it appeared that a temperature shift could not induce a structural change in existing FimH molecules.

Erythrocyte and macrophage binding by *fimH205* mutants. Guinea pig erythrocyte binding was greatly influenced by temperature in the *fimH205* mutant: *fimH205* mutants bound erythrocytes with statistically the same effectiveness as a *fimH* insertion mutant at 42°C but had the binding effectiveness of the parental strain at 31°C (Fig. 5). In contrast, the ability of the *fimH205* mutant to bind to macrophages was statistically the same as that of the parent at both 31 and 42°C (Fig. 5). For

comparison, the results of this binding assay with the four other *fimH* mutants (the ones that did not agglutinate in FimH antiserum at parental levels) are shown. These four mutants bound with the same effectiveness (statistically) as the *fimH* insertion mutant.

Macrophage binding by the *ts* mutant grown at the restrictive temperature was mannose insensitive. The ability of the nonmetabolizable mannose analog α mm to inhibit macrophage binding was assessed with a *fimH205* mutant (ORN183) grown at 31 and 42°C. The degree to which the addition of 50 mM α mm inhibited binding was compared with binding by the parental strain (Fig. 6). Binding inhibition was quantitated by normalizing the reduction in binding effected by α mm on the parent and mutant relative to a *fimH* insertion mutant present as an internal control in all assays. For the parental strain, α mm inhibited binding approximately 3.5-fold regardless of the temperature (31, 37, and 42°C tested). For the mutant, however, α mm failed to change the binding significantly in 42°C-grown cells. After growth at 31°C, α mm inhibited macrophage binding of the mutant to a degree that was statistically the same as inhibition in the parent.

DISCUSSION

The results reported herein define the location of lesions and phenotypic properties of five *fimH* point mutants isolated after site-directed mutagenesis (13). Although 11 independently isolated *fimH* mutants were initially identified, pilot experiments revealed that some had identical *fimH* lesions. The five described here represent one of each allelic type isolated. One mutant with a novel conditional erythrocyte binding phenotype was further defined as having an altered binding specificity after growth under permissive and restrictive conditions.

The *fimH* mutants examined here were obtained by enriching for individuals that formed pellicles in static broth (a property associated with FimH) in the presence of α mm (13). α mm is a mannose analogue that inhibits pellicle formation as well as erythrocyte agglutination by type 1 pili. Since pellicle formation is eliminated in *fimH* insertion mutants (13), we expected that most, if not all, of the *fimH* mutants would express parental levels of FimH and that the lesions would thus define regions of FimH required for erythrocyte binding. However, in the present study, bacterial agglutination tests with polyclonal antiserum raised against a LacZ-FimH fusion protein did not confirm that parental amounts of FimH were being produced in four of the five mutants. Whereas it is possible that some of these mutants have lesions that define areas of FimH responsible for erythrocyte binding and are poorly reactive with antibody as a consequence, we could not rule out the possibility that their failure to agglutinate (or bind to) erythrocytes was due entirely to inadequate FimH exposure or expression.

Two of the four mutants that did not express normal amounts of FimH-cross-reacting material (those with the *fimH236* and *fimH208* alleles) displayed signs of defective pilus biogenesis (13). We assume that their phenotypes result from the aberrant routing of the defective products in the pilus biogenesis process. In the case of the *fimH236* mutation, the lesion lies in an area of the gene encoding the signal sequence (12), and the altered amino acid would thus not be part of the mature protein. However, the *fimH236* lesion may result in an alteration in the site at which the signal is cleaved (49), creating a defect in the amino-terminal portion of the mature protein. The nature of the *fimH208* lesion (a frameshift lesion approximately one-third of the way into the gene produced a garbled sequence of 33 amino acids before terminating, leaving a prod-

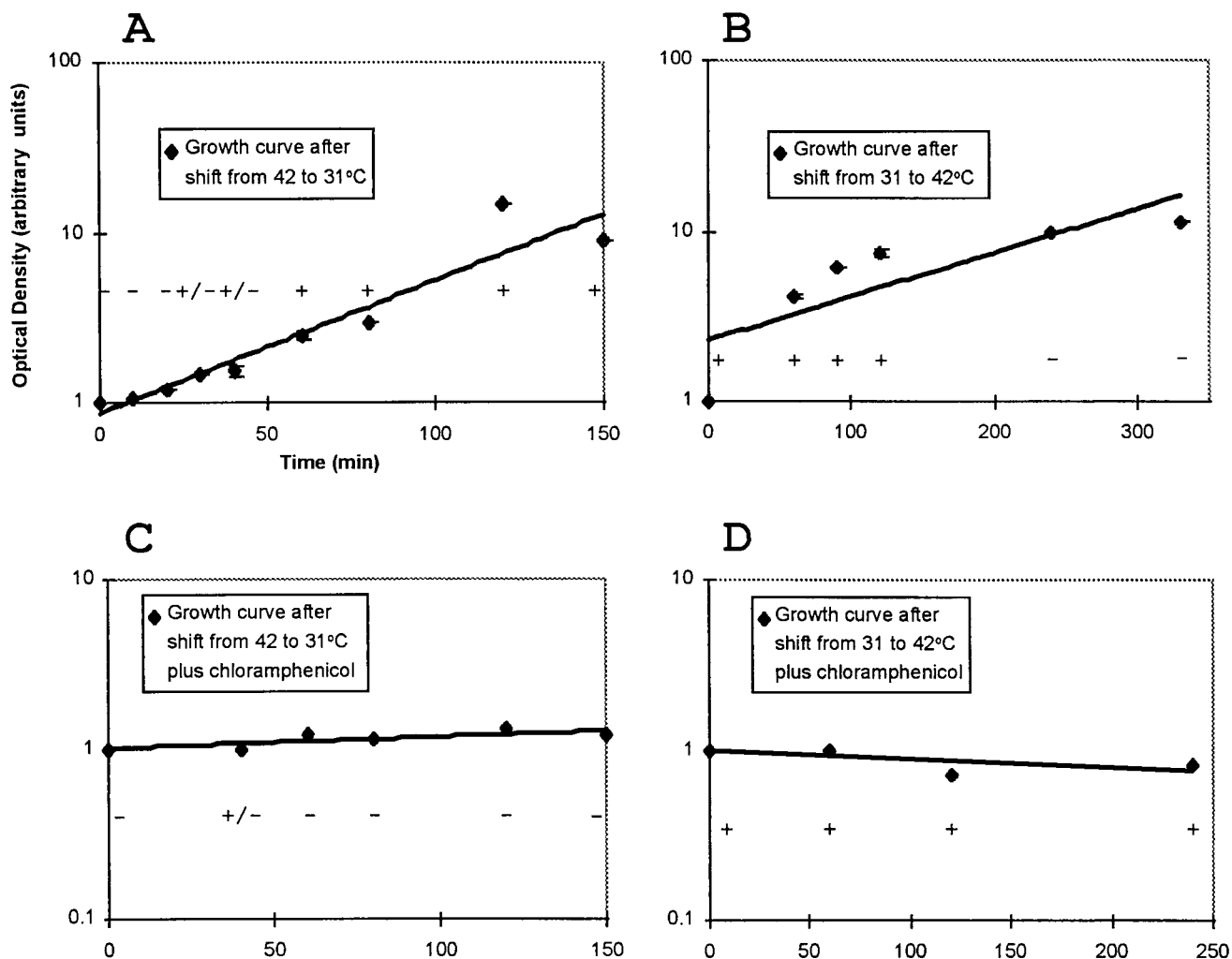


FIG. 4. Kinetics of the appearance of a hemagglutination-positive or -negative phenotype after a shift in temperature from 31 to 42°C and vice versa. (A and B) Emergence and disappearance, respectively, of the hemagglutination phenotype following a temperature shift. (C and D) Same shifts but in the presence of chloramphenicol (20 $\mu\text{g}/\text{ml}$). Diamonds represent points on growth curves taken from at least two experiments. Optical density at 600 nm values were normalized to the starting value in each experiment. Consequently, the units are arbitrary. The line shown through the points is a trend line drawn from a logarithmic regression analysis of the points. Hemagglutination was scored as a positive (+), weakly positive (+/-), or no hemagglutination (-) depending on the strength of the agglutination reaction. To test hemagglutination, samples were removed at the times indicated, diluted (or concentrated) to a constant optical density, and tested for hemagglutination of guinea pig erythrocytes as described in the text. Error bars represent standard error of the mean.

uct approximately 40% of normal size, 33% of which was garbled) produces a mutant with the most dramatically altered pilus morphologic phenotype (13). Such mutations may affect the ability of the product to properly interact with the FimC chaperone protein (14) or with minor pilus components needed for pilus assembly (15, 36, 39). The recently acquired crystalline structure of the FimC-FimH complex should prove helpful (6) in this regard.

By far the most straightforward mutant to characterize was the *ts fimH205* mutant. This mutant had parental levels of FimH and had a very pronounced phenotype. The lesion defining this allele converted a leucine to an arginine approximately one-fifth of the way through the mature FimH protein (amino acid position 58). The location and possible importance of this particular amino acid have been previously noted in studies by Sokurenko et al. (45, 46), who examined naturally occurring *fimH* alleles. In one of their studies, the *fimH* allele from *E. coli* K-12 strain CSH50 had arginine at the 58 position (GenBank accession no. A36976). This allele conferred the

ability to bind to periodate-treated fibronectin in a mannose-inhibitable fashion. In fact, only one additional amino acid change in the CSH50 *fimH* allele, at position 201 (a histidine in place of a threonine), kept the two alleles from being identical. (Our parental *fimH* allele was identical to that of *E. coli* K-12 strain MG1655 that has been completely sequenced [2]; GenBank accession number U14003.) Sokurenko et al. (46) commented on the similarity of the *fimH205* mutants to carry out protein-protein interactions in pellicle formation and the ability of the CSH50 FimH to bind to periodate-treated fibronectin. It was thus reassuring to find, in the present study, that the two mutants have one very specific genotypic feature in common. However, in the report of Sokurenko et al. (46), no mention was made of a conditional nature of fibronectin binding. Also, the binding reported was subject to inhibition by mannose. (The protein-protein binding conferred by the *fimH205* allele in forming a pellicle [13] and in binding to macrophages was insensitive to mannose inhibition.) A recent report by Pouttu et al. (35) has indicated that the collagen

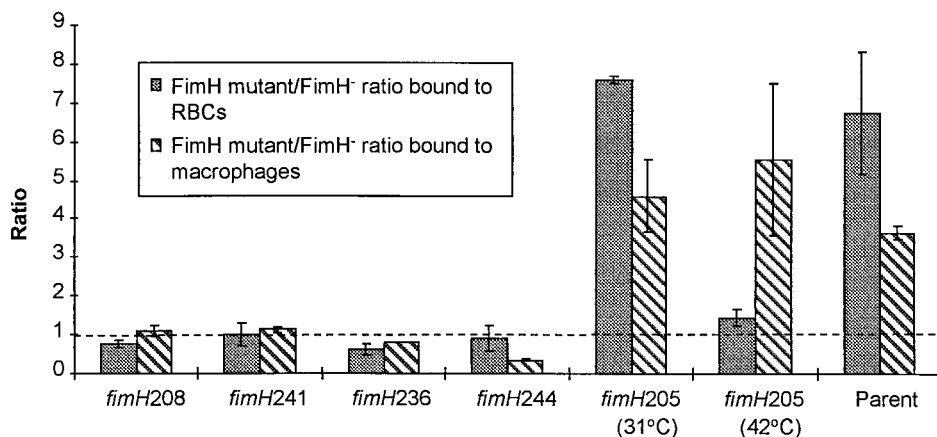


FIG. 5. Comparison of the binding ability of mutants having the *fimH* lesions indicated relative to a *fimH* insertion mutant (*fimH*⁻-*kan* strain ORN133, denoted as FimH⁻ in the legend box) and compared to the parental (ORN155) strain for their ability to bind guinea pig erythrocytes (RBCs) and resident BALB/c peritoneal macrophages. All mutant strains were grown at 37°C except for the *fimH205* mutant, which was grown at the temperatures indicated. Parental binding values (ratio of the parental strain to the *fimH* insertion mutant; far right-hand bars were not significantly affected by the growth temperatures employed (31, 37, or 42°C), and for economy, a single average value is shown. Vertical bars represent the standard error of the mean. The dashed line indicates a 1:1 ratio between the test strain and the *fimH* insertion mutant. At least two experiments were averaged.

binding ability of FimH is related to an amino acid change at residue 62. Whether this binding was mediated through protein-protein interactions in collagen was not specifically addressed. In any case, it appears that residues 58 to 62 of the FimH protein are important for determining receptor binding specificity. In addition to the above, it has been previously reported that linker insertion mutagenesis that changed residue 56 abolished mannose-sensitive binding (41).

In the present study, the nature of the *ts* defect in products from the *fimH205* allele was pointed out in experiments that measured phenotypic lag after a temperature shift. Since new protein synthesis was required to effect a change in phenotype after a temperature shift, presynthesized FimH evidently could not undergo a conformational change to assume the new activity. Rather, a new (or nascent) FimH molecule was needed to assume the new configuration responsible for the altered activity. Also, the differences in phenotypic lag when shifting

from the permissive to the nonpermissive temperature and vice versa were consistent with the idea that comparatively few pili (containing functional FimH products) were sufficient for hemagglutination. That is, acquisition of the hemagglutination-positive phenotype after a temperature shift was much more rapid than the loss of the hemagglutinating ability after a reverse shift.

Of most interest to us was the finding that the *ts* mutant carrying the *fimH205* allele had different eucaryotic cell binding specificities when grown at the permissive and restrictive temperatures. At the restrictive growth temperature, the *fimH205* mutant, while not binding to erythrocytes, still bound to macrophages through a FimH-specific interaction that was insensitive to mannose inhibition. The most likely explanation for this phenotype involves the earlier observation (10) that at the restrictive growth temperature, this mutant forms pellicles that appear to involve FimH-FimH interactions that are insensitive to mannose inhibition. This protein-protein type of interaction may be employed here in binding to macrophages. That is, macrophages may have a protein(s) on their surface capable of interacting with the mutant version of FimH. This explanation implies that erythrocytes lack such a protein(s). In related experiments by Sokurenko et al. (43), a correlation was made between the degree of affinity of FimH products (from naturally occurring *fimH* alleles) for monomannose and their affinity for cultured uroepithelial cells. It would thus appear that FimH has the potential to display a number of binding specificities.

Factors that determine FimH binding specificity have the potential to be exploited for processes that require conditional attachment and release of microorganisms from specific ligands. Already, the natural binding affinity of FimH for mannose has been used in combination with other binding domains to create chimeric adhesins (40). Another use for adhesin mutants with altered receptor specificities is in experiments examining the effects of bacterium-eucaryotic cell interactions. The effects of bacterial attachment on eucaryotic cell physiology are just beginning to be understood (1, 8, 22). The effects of specific types of attachment on eucaryotic cells are almost always deduced from experiments in which bacterial mutants that lack the attachment organelle (or the adhesive part of the

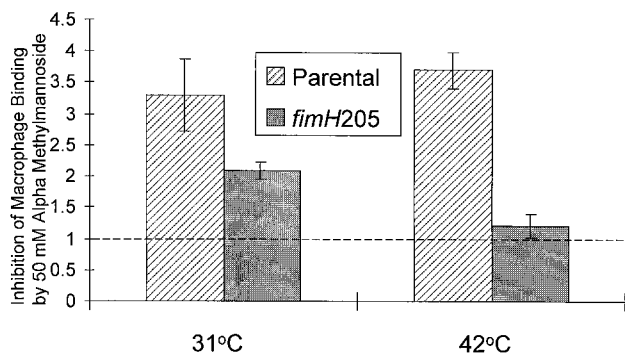


FIG. 6. Degree of inhibition of macrophage binding by α mm in the *fimH205* mutant grown at the restrictive and permissive temperatures. Macrophage binding was carried out in microtiter wells containing either the parental strain (ORN115) or a *fimH205* mutant strain (ORN183) each mixed with a *fimH*⁻-*kan* insertion mutant strain (ORN204) as described in the text. The degree to which α mm inhibited binding of the parent or *fimH205* mutant (ordinate values) was calculated from the change in ratio of these strains normalized to the change exhibited by the *fimH*⁻-*kan* insertion mutant. A value of 1 (dashed line) defines the value expected if there was no effect of α mm addition. Results presented are the averages of three separate experiments, each performed in duplicate. Error bars denote standard error of the mean.

organelle) are used as negative controls (9, 11, 16). In the case of type 1 pili, such mutants do not bind eucaryotic cells with an efficiency high enough to serve as a truly appropriate control (i.e., one that eliminates the effects of nonspecific binding rather than just the elimination of binding entirely). Attachment specificity mutants, such as the one isolated here, may allow better-controlled assays of the effect of receptor-ligand interactions to be determined.

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