In vivo detection of *Escherichia coli* type 1 fimbrial expression and phase variation during experimental urinary tract infection

Carsten Struve and Karen Angeliki Krogfelt

Adhesion mediated by fimbriae is thought to play an important role in the pathogenesis of urinary tract infections (UTI) by *Escherichia coli*. The majority of clinical isolates of *E. coli* from UTI are able to express type 1 fimbriae. However, the importance of these fimbriae as a virulence factor has been controversial. To investigate the expression of type 1 fimbriae *in vivo* during UTI, mice were transurethrally infected with uropathogenic *E. coli* C175-94 and type 1 fimbrial expression was determined directly by two independent methods at 2 h, 1 d and 3 d after infection. By use of an assay combining *in situ* rRNA hybridization and immunofluorescence, all bacterial cells detected in urine, bladders and kidneys from mice sacrificed 1 and 3 d after onset of infection were found to express type 1 fimbriae. In contrast, the majority of cells in the suspension used for infection of mice and specimens from mice sacrificed 2 h after inoculation were found to be non-fimbriated. Similar results were obtained with a PCR assay revealing the orientation of the invertible promoter driving the transcription of type 1 fimbrial genes. Whilst the promoter in both ON and OFF positions could be amplified from the suspension used for infection and specimens from mice sacrificed 2 h after inoculation, at 1 and 3 d after onset of infection only the promoter in the ON orientation could be amplified. These results show that introduction of *E. coli* C175-94 into the mouse urinary tract leads to markedly enhanced expression of type 1 fimbriae.

Keywords: type 1 fimbriae, phase variation, urinary tract infection, *in situ* hybridization, immunofluorescence

INTRODUCTION

*Escherichia coli* is the pathogen responsible for most cases of uncomplicated urinary tract infections (UTI), one of the most frequent human infectious disorders (Sobel, 1991). Fimbriae mediate the ability of *E. coli* cells to adhere to the uroepithelium, thereby resisting elimination by the flow of urine. Adhesion is therefore considered an important step in the pathogenesis of UTI (Reid & Sobel, 1987). Among the different types of fimbrial adhesins, P fimbriae have been identified as an important virulence factor among uropathogenic *E. coli* strains (Donnenberg & Welch, 1996). The role of type 1 fimbriae in pathogenesis has been less clear, especially due to the ubiquity of the fimbriae, as they are found on approximately 70% of all *E. coli* isolates and most P-fimbriated *E. coli* strains also possess type 1 fimbriae (Orndorf & Bloch, 1990; Hagberg *et al*., 1981). Furthermore, *in vitro* adherence studies of either purified type 1 fimbriae or type-1-fimbriated *E. coli* strains to formaldehyde-fixed human tissue and to exfoliated epithelial cells from the urine gave contradictory results (Fujita *et al*., 1989; Virkola 1987; Ørskov *et al*., 1980).

Recent studies of UTI with animal models show that type-1-fimbriated *E. coli* cells are better colonizers of the urinary tract than their isogenic mutants (Connell *et al*., 1996; Langermann *et al*., 1997; Sokurenko *et al*., 1998; Mulvey *et al*., 1998). Type 1 fimbriae mediate adhesion to structures containing D-mannose, but have also been shown to mediate binding to non-glycosylated peptide epitopes (Sokurenko *et al*., 1994). The *fim* gene cluster, consisting...
of nine genes, includes fim\textit{A}, which encodes the major
fimbrial subunit (Klemm & Krogfelt, 1994). The
expression of type 1 fimbriae is characteristically phase-
variable; i.e. individual cells can switch between
fimbriated and non-fimbriated states. The phase
variation is mediated by a 314 bp invertible DNA element
(phase switch) containing the promoter of \textit{fim}\textit{A}
(Abraham \textit{et al}., 1985). When the switch is in one
position (ON) the promoter drives the transcription of
\textit{fim}\textit{A} and fimbriae are produced, whilst inversion of the
switch to the OFF position leads to arrest of transcrip-
tion and non-fimbriated cells. Two recombinases
encoded by the \textit{fim} cluster, \textit{FimB} and \textit{FimE}, mediate the
inversion of the invertible element (Klemm, 1986).
Several elements regulating the expression of \textit{FimB} and
\textit{FimE} and the sensitivity of the invertible element
towards switching, including environmental factors
such as temperature and growth media, have been
identified (Old & Duguid, 1970; Schwan \textit{et al}., 1992;
Blomfield \textit{et al}., 1993; Gally \textit{et al}., 1993; Newman \textit{et al}.,

Only a few studies have investigated the expression of
type 1 fimbriae in \textit{vivo} during UTI. These studies have either
used indirect methods, i.e. determined fimbrial
expression in bacteria isolated from specimens from
infected animals after growth in vitro (Hagberg \textit{et al}.,
1983; Schaeffer \textit{et al}., 1987), or investigated bacterial
cells in urine collected from patients suffering from UTI
(Oek \textit{et al}., 1981; Harber \textit{et al}., 1982; Ljungh & Wadström,
\textit{et al} (1983) investigated the expression of type 1
fimbriae in urine and bladder lavages from infected
mice by electron microscopy and in a recent study, a
PCR assay was used to determine the orientation of the
phase switch in specimens from infected mice (Lim

In the present study, we used two independent assays
to detect directly the expression of type 1 fimbriae in \textit{vivo}
during experimental UTI: a combination of \textit{in situ}
rRNA hybridization and immunofluorescence for de-
tection of fimbrial expression, and a PCR assay for
determination of the orientation of the phase switch of
\textit{E. coli} bacteria in tissues of the urinary tract to establish
the kinetics of phase variation of the fimbriae during
infection.

\section*{METHODS}

\textbf{Bacterial strains and media.} \textit{E. coli} C175-94, a clinical isolate
from a patient with UTI, was used throughout the study. It
belongs to serotype O8:K48:H9, expresses type 1 fimbriae but
not P or S fimbriae and does not produce haemolysin. \textit{Yersinia}
\textit{enterocolitica} SSI 7558 serotype O3 was used as control for
specificity in the development of the combined \textit{in situ}
hybridization and immunofluorescence assay. Bacteria were
grown in either L broth (Statens Serum Institut; SSI) or on
bromothymol blue agar plates (SSI).

\textbf{Preparation of inoculum for infection of mice.} Bacteria were
grown in filter-sterilized human urine and passed three
times as described by Sharma \textit{et al} (1991). They were then
incubated overnight at 37 \textdegree C with shaking, centrifuged at
6500 g for 10 min and the pellet suspended in PBS (130 mM
NaCl, 8 mM Na\textsubscript{2}HPO\textsubscript{4}, 2 mM KH\textsubscript{2}PO\textsubscript{4}, pH 7.4) (SSI) to a
concentration of approximately \(10^{10}\) c.f.u. ml\textsuperscript{-1}.

\textbf{Mouse model of ascending UTI.} Outbred female albino mice,
Ssc:CF1, 30±2 g (SSI) were used. The mice were anaes-
thetized by intraperitoneal administration of 0.08 ml mixture of
Hypnorm (0.315 mg fentanyl citrate ml\textsuperscript{-1} and 10 mg
fluanison ml\textsuperscript{-1}; Janssen Animal Health) and diazepam (5 mg
ml\textsuperscript{-1}) in a ratio of 10:3. Twenty-four mice were inoculated
transurethrally with 0.05 ml of the bacterial suspension by use of
plastic catheters (SSI). As control, six mice were inoculated
with PBS. The catheter was carefully pushed in horizontally
through the urethral orifice until it reached the top of the
bladder, whereupon the bacterial suspension was slowly
injected into the bladder. The catheter was immediately
removed after inoculation and the mice subjected to no further
manipulations until sacrifice.

The mice were sacrificed in groups of 10 by cervical dislocation
2 h, 1 d and 3 d after inoculation. Urine was collected from
each mouse by gentle compression of the abdomen and then
the bladder and kidneys were removed aseptically. The organs
were bisected and one half of each specimen was either used
for preparation of tissue sections or as sample for PCR. The
other half was homogenized manually in 500 \mu l collagenase
solution (500 units ml\textsuperscript{-1}; Sigma) and serially diluted for
quantitative culture on bromthymol blue agar plates (SSI).

\textbf{Preparation of samples for combined \textit{in situ} hybridization and
immunofluorescence assay.} Immediately after death, bladders and
kidneys from inoculated mice were transferred to a phosphate-buffered 4 %
solution of formalin. The fixed organs were then dehydrated and embedded in paraflin prior
to preparation of 5 \mu m thick sections mounted on glass slides. Prior to hybridization, paraflin was recovered from the
sections by treatment with xylene 3 times for 10 min followed
by 10 min dehydration in 96 % ethanol. After drying at room
temperature, the sections were circumscribed with a hydro-
phobic pen (Dako).

Urine from inoculated mice, inoculum suspension and over-
night cultures of C175-94 were centrifuged at 6500 g; the pellets were
resuspended in 4 % formalin solution and kept at 4 \textdegree C for 20 min. The fixed cells were isolated by centrifugation and washed three times with PBS. The washed cells were sus-
pended in PBS and 15 \mu l samples of the suspension were then transferred to wells in a poly-l-lysine-coated 6-well slide and
left to dry at room temperature.

\textbf{Oligonucleotide probe and antisera.} Probe EC1531 (5′-
CACC GTAG GCCTG CATCA 3′) specific for \textit{E. coli} 23S
rRNA, labelled with CY3, was used for visualization of \textit{E. coli}
cells as previously described (Poulsen \textit{et al}., 1994).

For detection of expression of type 1 fimbriae, rabbit
antisera raised against purified type 1 fimbriae was used
(Krogfelt & Klemm, 1988). Rabbit antisera against \textit{Y.
enterocolitica} O3 antigen (SSI) was used for detection of \textit{Y.
enterocolitica} cells. FITC-conjugated swine anti-rabbit
immunoglobulin (Dako) was used as secondary antibody.

\textbf{Combined \textit{in situ} hybridization and immunofluorescence
assay.} Hybridization was performed by adding hybridization
solution [10 % formamide, 0-1 M Tris pH 7-2, 0-9 M NaCl] containing 2-5 ng probe EC1531 \mu l\textsuperscript{-1} to tissue sections or slide
wells. After incubation in humidified chambers overnight
at 37 \textdegree C, the slides were washed in washing solution (0.1 M Tris
pH 7-2, 0.9 M NaCl) for 15 min, followed by incubation with antibody diluted in washing solution containing 2.5 ng probe
EC1531 \mu l\textsuperscript{-1} for 45 min. The slides were then washed 3 times
overnight. The experiment was repeated three times.

For the inoculum suspension and each urine sample, three different slide wells were evaluated. The total number of cells in three microscope fields, typically 30–300 cells, was counted and compared to the number of cells expressing type 1 fimbriae, respectively.

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Preparation of samples for PCR. Inoculum suspension, urine, bladders and kidneys sampled for PCR were collected in Eppendorf tubes containing 500 µl PBS, immediately boiled for 5 min and kept at –80 °C until use. After thawing, the organs were homogenized manually. Organ homogenates, urine samples and inoculum suspension were boiled for 20 min, centrifuged at 10000 g for 15 min and the supernatants were removed and kept at –20 °C until use as templates for PCR reaction.

PCR amplification and restriction enzyme digestion. PCR primers A (5′-GAGAAGAGGTGGTATTTACCTTTTG-3′) and B (5′-AGAGCGCTGTAGAAGCTGG-3′) were used for amplification of a 559 bp DNA element covering the fim switch region as described by Roesch & Blomfield (1998). The Expand High Fidelity PCR System (Boehringer Mannheim) was used for the PCR reaction. The reaction mixtures were prepared according to the instructions of the manufacturer in a total of 50 µl 1.5 mM MgCl2 solution containing 2 µl template. The reaction conditions were as follows: an initial denaturation at 94 °C for 1 min; followed by 40 cycles of denaturation at 94 °C for 1 min, annealing at 58 °C for 70 s and elongation at 72 °C for 70 s. The reaction was completed by a final elongation step at 72 °C for 3 min.

The PCR products were digested by the restriction endonuclease HindIII (New England Biolabs) according to the manufacturer’s instructions. The digested PCR products were electrophoresed in 3% agarose gels, stained by ethidium bromide and visualized by UV illumination.

Growth in mouse urine, bladder mucus and L broth. Urine was collected from six mice, sterilized by UV illumination and dialysed against 0.9% NaCl overnight.

Mucus was scraped from the bladders of six mice using a rubber spatula. Dialysed urine, crude mucus suspension and L broth were inoculated with E. coli C175-94 corresponding to a concentration of 10^4 c.f.u. ml⁻¹ and incubated at 37 °C overnight. The experiment was repeated three times.

RESULTS

Experimental UTI

Mice were successfully infected with E. coli C175-94 as evaluated by viable counts of bacteria in kidneys from mice sacrificed 2 h, 1 d and 3 d after inoculation. Two hours after inoculation, bacteria were recovered in low numbers from the majority of kidneys, reflecting transfer of bacteria from the bladder to the renal pelvis. One day after inoculation, all mice were found to have positive bladder cultures, the number of bacteria ranging from 10^2 to 10^6 c.f.u. and the median being 7.2 × 10^5 c.f.u. per bladder. Six of 8 mice had positive kidney cultures, 10 of 16 kidneys were infected and the number of bacteria ranged from 10^2 to 10^6 c.f.u., the median being 3.6 × 10^5 c.f.u. per kidney. Three days after inoculation, all mice were found to have infected bladders, the number of bacteria ranging from 10^2 to 10^6 c.f.u. and the median being 2.1 × 10^5 c.f.u. per bladder. Positive kidney cultures were found in 6 of 8 mice, 10 of 16 kidneys were infected, the number of bacteria ranged from 10^2 to 10^6 c.f.u. and the median was 6.0 × 10^5 c.f.u. per kidney. All specimens from mice inoculated with PBS were found to be sterile 2 h, 1 d and 3 d after inoculation.

Development of combined in situ hybridization and immunofluorescence assay

For simultaneous visualization of E. coli cells and detection of type 1 fimbriae expression, an assay combining in situ rRNA hybridization and immunofluorescence was developed.

In situ hybridization was initially carried out as described previously (Poulsen et al., 1994), but the presence of SDS in the hybridization solution was found to cause non-specific binding of antibodies to the bacterial cells. After omission of SDS, the hybridization process was found to have no effect on the subsequent use of antibodies. Washing out of the oligonucleotide probe during the immunofluorescence assay was avoided by use of washing buffer with a high salt concentration and addition of the oligonucleotide probe during incubation with antibody.

The specificity of the combined assay was tested on a mixture of E. coli C175-94 and Y. enterocolitica cells. The two bacterial species are easy to distinguish by phase-contrast microscopy due to their different cell morphology. When probe EC1531, specific for E. coli 23S rRNA, and antiserum against Y. enterocolitica were used in the combined assay, the two species were clearly distinguishable by fluorescence microscopy (results not shown). No Y. enterocolitica cells were stained by the CY3-labelled EC1531 probe and no E. coli cells were stained by FITC-labelled antibody, confirming that the specificity of the hybridization and the immunostaining remained intact when the two techniques were combined.

Expression of type 1 fimbriae during experimental UTI

The expression of E. coli type 1 fimbriae during UTI was investigated by use of the combined in situ hybridization and immunofluorescence assay on tissue sections and urine from infected mice sacrificed 2 h, 1 d and 3 d after inoculation.

A mean of 25% (range 15–40%) of the E. coli C175-94 cells were found to express type 1 fimbriae when grown in vitro, as shown by immunostaining of the bacterial
Fig. 1. Monitoring expression of type 1 fimbriae by combined *in situ* rRNA hybridization and immunostaining of (a, b) inoculum suspension used for infection of mice, and (c, d) bladder section from mouse sacrificed 1 d after onset of infection. (a, c) *In situ* hybridization with CY3-labelled probe specific for *E. coli* 23S rRNA. (b, d) Immunostaining with antiserum against type 1 fimbriae. Bars: (a, b) 10 µm; (c, d) 7.5 µm.
suspension used for infection of mice (Fig. 1a, b). Likewise, the majority of bacteria present in urine 2 h after inoculation were found to be non-fimbriated (results not shown). In bladder sections from mice sacrificed 2 h after inoculation, bacteria were found randomly distributed as single cells either in contact with the bladder mucosa or free in the lumen; both fimbriated and non-fimbriated cells were observed (results not shown). Due to the low number of bacteria present in kidneys from mice sacrificed 2 h after inoculation, it was not possible to detect any bacteria in the kidney sections.

One and three days after onset of infection, expression of type 1 fimbriae was detected in all bacterial cells observed in urine (results not shown). In bladder sections, bacteria were observed forming microcolonies associated with the bladder epithelium (Fig. 1c, d). The microcolonies were only seen in some parts of the bladder, leaving most of the bladder mucosa uncolonized. Many of the bacterial cells observed in bladder sections were found to have undergone changes in cell morphology and were radically elongated, often with filamentous cell structures. As in urine, all bacterial cells observed in bladder sections from mice sacrificed 1 and 3 d after onset of infection were found to express type 1 fimbriae (Fig. 1c, d). In kidneys, bacteria were detected in the renal pelvis and were often embedded in the mucosa or in close contact with inflammatory cells, making the bacterial cells difficult to identify by phase-contrast microscopy. By use of the combined assay, bacterial cells were easily identified by in situ hybridization and the expression of type 1 fimbriae could be determined. In kidneys, all bacterial cells detected were found to express type 1 fimbriae (results not shown).

Orientation of the E. coli fim switch during experimental UTI

The orientation (ON or OFF) of the invertible fim switch, which drives the transcription of type 1 fimbriae genes, was determined by PCR amplification of a 559 bp element containing the switch, followed by restriction endonuclease digestion of the PCR product (Fig. 2a). Hinfl cuts the invertible DNA element asymmetrically resulting in different sized fragments depending on whether the switch is in the ON or OFF orientation (Fig. 2b). PCR was performed on urine and organ homogenates from infected mice. As expected, Hinfl digestion of the PCR product from the bacterial suspension used for infection of mice resulted in four fragments, corresponding to the switch being in both the ON and OFF positions (Fig. 2c); i.e. both cells transcribing and not transcribing fim genes were present in the inoculum solution. Bacteria from urine, bladder and kidney samples of mice sacrificed 2 h after inoculation were also found to have the switch in both the ON and OFF position (Fig. 2c). In contrast, only fragments corresponding to the switch being in the ON position could be detected after digestion of PCR products from urine, bladders and kidneys of mice sacrificed 1 and 3 d after onset of infection (Fig. 2c); i.e. during established UTI all bacterial cells in urine, bladder and kidneys of infected animals transcribed type 1 fimбриae genes.

Fig. 2. Orientation of the E. coli type 1 fimbriae phase switch during experimental UTI. (a) Schematic representation of the phase switch region including the invertible DNA element containing the promoter (P) of fimA. The orientation of the phase switch was determined by PCR amplification of the switch region followed by digestion of the PCR product with Hinfl. (b) Theoretically expected results after PCR amplification of the switch region. Due to asymmetric location of the Hinfl cleavage site within the invertible element, different fragments are obtained depending on the orientation of the switch. (c) In vivo results of PCR amplification of the fim phase switch. Orientation of the switch in inoculation suspension (i) and urine (U), bladders (B) and kidneys (K) from mice sacrificed 2 h, 1 d and 3 d after inoculation. Lane M, molecular size markers; lane UC, uncut PCR product.
Influence of growth media on expression of type 1 fimbriae

To investigate whether components of mouse urine or bladder mucus had any effect on the expression of type 1 fimbriae, overnight cultures of C175-94 grown in these media were immunostained with antiseraum against type 1 fimbriae and compared with cells grown in L broth. Since mouse urine was found to be bacteriocidal for C175-94, urine dialysed against 0.9% NaCl overnight was used.

No difference in the expression of type 1 fimbriae was observed between bacterial cultures grown in dialysed mouse urine, crude bladder mucus or L broth (results not shown). Approximately 25% (range 15–40%) of the mouse urine, crude bladder mucus or L broth (results observed between bacterial cultures grown in dialysed No difference in the expression of type 1 fimbriae was was used. 

Whilst animal models of UTI have been an important instrument in the assessment of virulence determinants of uropathogenic bacteria, the actual expression of specific virulence factors during infection has only been investigated in few studies. In the present study, two independent methods were used for direct determination of the expression of E. coli type 1 fimbriae in the urinary tract of mice infected with the uropathogenic E. coli C175-94. It is shown here that type 1 fimbriae are indeed expressed in vivo and that the phase switch is always towards the ON orientation during infection.

For simultaneous detection of E. coli cells and expression of type 1 fimbriae in tissue sections from infected mice, an assay combining in situ rRNA hybridization and immunofluorescence was developed. Even though both in situ rRNA hybridization and immunofluorescence are frequently used techniques, to our knowledge, only one study has so far described a combination of the two methods. In that work, a combination of in situ hybridization and immunofluorescence was used for the identification of Clavibacter michiganensis sepedonicus, but only in laboratory cultures (Li et al., 1997). In the present study, the specificity of the in situ hybridization and immunofluorescence assay when the two techniques were combined was confirmed by using a mixture of E. coli and Y. enterocolitica cells. The simultaneous identification of specific bacterial species and detection of expression of specific antigens by use of the combined assay is a useful tool in studies of complex bacterial environments, for example the gut, skin and environmental biofilms.

The expression of type 1 fimbriae during UTI was investigated by submitting urine, bladders and kidneys from mice sacrificed at different time points after inoculation with E. coli C175-94 to the combined assay of in situ hybridization and immunofluorescence. E. coli cells were detected in bladders as well as in kidneys. The presence of radically elongated bacteria in bladders has been reported previously and may be due to the high osmolarity of mouse urine (Hagberg et al., 1986). Two hours after infection, both fimbriated and non-fimbriated bacterial cells were detected in the specimens, but on days 1 and 3 after infection all bacterial cells in urine, bladders and kidneys were found to be heavily fimbriated. In contrast, approximately a quarter of the cells in the suspension used for infection of mice were fimbriated. Thus, the introduction of E. coli C175-94 into the mouse urinary tract selects for a markedly enhanced expression of type 1 fimbriae. The presence of only fimbriated bacteria in bladders and kidneys cannot be explained by specific selection for fimbriated cells via elimination of non-fimbriated cells by the flow of urine as only fimbriated cells were detected in urine 1 and 3 d after infection. If the presence of only fimbriated cells in bladders and kidneys was due solely to selection by enhanced adhesion of the fimbriated cells, then it would be expected that non-fimbriated cells would be detected in the urine. Since the phase switch occurs regularly, adhering fimbriated cells would occasionally give rise to non-fimbriated cells which should be present in the urine. Thus, our results show that the phase switch is directed to the ON position by an environmental signal in the urinary tract.

The above results were confirmed by PCR amplification of the invertible phase switch which drives the transcription of type 1 fimbriae. Only the switch in the ON position could be amplified from urine, bladders and kidneys from mice sacrificed 1 and 3 d after onset of infection, whilst the switch in both ON and OFF positions could be amplified from specimens from mice sacrificed 2 h after infection, as well as from the suspension used for inoculation of mice. Our results suggest that in the mouse urinary tract some unknown factors influence the type 1 fimbriae phase switch in such a way that the switch is always ON, and consequently fimbriae are continuously produced. Surprisingly, neither growth in bladder mucus or mouse urine promoted type 1 fimbriae expression in C175-94 compared to when the bacteria was grown in L broth. However, as dialysis of mouse urine was necessary for growth of C175-94, the presence of a dialysable factor promoting type 1 fimbriae expression in urine cannot be excluded.

This study clearly indicates that expression of type 1 fimbriae is markedly enhanced during UTI, as has been shown previously for the expression of type 1 fimbriae during E. coli colonization of the gut (Krogfelt et al., 1991). Enhanced fimbrial expression during experimental UTI has also been reported for Proteus mirabilis, which expresses MR/P fimbriae (Zhao et al., 1997). Studies investigating expression of type 1 fimbriae in E. coli isolated directly from urine of humans suffering from UTI by the use of either immunofluorescence, electron microscopy or agglutination assays generally find that bacteria in urine produce fewer type 1 fimbriae compared to strains subcultured in vitro (Ofek et al., 1981; Harber et al., 1982; Ljungh & Wadström,
In many of these studies, the expression of type I fimbriae may be affected by the processing of the samples, for example cooling and storage of the urine, or collection of urine from a specific patient group, for example patients with indwelling catheters. In bladder lavages of infected mice, bacteria were predominately fimbriated, but non-fimbriated bacteria were also present (Hultgren et al., 1985). In a recent study, it was found by use of PCR amplification of the phase switch that 2 d after onset of infection in mice, the phase switch was ON in approximately one-third of the bacteria infecting bladders and kidneys (Lim et al., 1998). However, in the suspension used for infection of mice the phase switch was almost exclusively in the OFF position, i.e. the expression of type I fimbriae was actually found to be enhanced during infection. It is important to mention that the strain used by Lim et al. also produced P fimbriae, haemolysin and 5 fimbriae (Mobley et al., 1993); the presence of these virulence factors may make the expression of type I fimbriae less prominent during infection of the urinary tract. Generally, it must be expected that the expression of a given virulence factor is influenced by the presence of other virulence determinants. Therefore, the patterns of fimbrial expression during infection might vary among different strains. That type I fimbriae are expressed during UTI is also supported by a recent study using the type I fimbria adhesin (FimH) for vaccination of mice (Langermann et al., 1997). The vaccinated mice showed markedly enhanced resistance against development of UTI following inoculation with E. coli.

In conclusion, we have shown by direct detection of type I fimbriae as well as by assessment of the orientation of the phase switch regulating expression of type I fimbrial genes that expression of type I fimbriae is markedly enhanced during UTI caused by E. coli C175-94. The type I fimbriae were expressed in all E. coli cells present in the urine, the bladder and the kidneys, and the switch was directed to the ON orientation, suggesting regulation of the switch by an environmental factor. Thus our results indicate that type I fimbrial expression is important during UTI.

This is, to our knowledge, the first study where the expression and phase variation of type I fimbriae during infection have been assessed in vivo.

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REFERENCES


In vivo phase variation of Escherichia coli type 1 fimbrial genes in women with urinary tract infection. Infect Immun 66, 3303–3310.


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