## The FimH A27V Mutation Is Pathoadaptive for Urovirulence in Escherichia coli B2 Phylogenetic Group Isolates

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Correlations between FimH mutations and virulence were established by studying a collection of human commensal and extraintestinal pathogenic *Escherichia coli* natural isolates. Pathoadaptive (A27V and, to a lesser extent, A119V) and "commensal-adaptive" (A202V) mutations were evidenced in B2 phylogenetic group strains. *fimH* phylogenetic analysis indicates that these pathoadaptive mutations occurred several times.

One of the mechanisms in bacterial evolution towards virulence is the allelic variation of existing genes that has been called pathoadaptive (19). One of the most studied pathoadaptive mutations is the allelic variation within the adhesive subunit of type I fimbriae, FimH, of Escherichia coli (15, 16, 18). This bacterium is a normal inhabitant of the intestinal tract of vertebrates but can also be responsible for intestinal and extraintestinal diseases (mainly urinary tract infections [UTI] and septicemia). Indeed, all E. coli isolates are capable of binding to trimannose receptors, allowing bacteria to bind buccal epithelial cells. In addition, 70% of UTI isolates bind to both trimannose and monomannose residues (17), allowing them to adhere strongly to uroepithelial cells. This change in tissue tropism is due to a small number of amino acid changes in FimH, an otherwise highly conserved molecule (1, 7, 14, 15). In vitro mutagenesis and analysis of the FimH sequences of a small number of natural isolates had allowed the identification of the amino acids responsible for an increase in binding monomannose residues (13, 17, 18). Moreover, previous studies have shown a link between long-term evolutionary history (strain phylogeny) and virulence in *E. coli*, as extraintestinal *E*. coli pathogens (including UTI strains) belonged preferentially to one of the four main E. coli phylogenetic groups, i.e., the B2 phylogenetic group (9, 12). Thus, the simultaneous action on virulence of both single-mutation effects and the accumulated divergence in genetic background over millions of years led us to study more carefully the diversity and evolutionary history of the *fimH* gene from natural strains belonging to the various E. coli phylogenetic groups (6). We sequenced the fimH gene from the first postsignal peptide codon to codon 232 in an epidemiologic collection encompassing both 44 commensal and 69 pathogenic strains. The strains were isolated mainly in France and in Benin and included two previously published collections (11, 12). fimH amplification was performed by standard PCR from bacterial lysate using primer fimHF (5'-TGA TGGGCTGGTCGGTAAATG-3') and fimH3R (5'-CGATA

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The Pearson correlation coefficients between the different FimH amino acid mutations and the characteristics of the strains, i.e., their origin of isolation (normal feces, urine, blood, miscellaneous infections), their phylogenetic groups, and the lethality in mice (0 to 1 mouse, 2 to 8 mice, and 9 to 10 mice killed of 10 mice inoculated), were calculated, and the statistical significance was evaluated by a Fisher test. Among the coding sequence, 33 amino acids were found to be variable, but

TABLE 1. FimH alleles of the studied natural isolates

Allele <sup>a</sup>	No. of strains (%)	Mutation at amino acid residue <sup>b</sup> :							In vitro binding affinity
		27	66	70	78	119	163	202	to yeast mannan cells (strain) <sup>c</sup>
Ι	32 (26.23)	А	G	Ν	S	А	V	А	Low (KB21)
II	23 (18.85)	V							High (PC31)
III	23 (18.85)			S	Ν				Low (F18)
IV	11 (9.01)					V			Low (CI3)
V	9 (7.34)			S	Ν		А		Low (MI11-2)
VI	8 (6.56)							V	
VII	4 (3.28)		S					V	
VIII	3 (2.46)		С						
IX	3 (2.46)	V		S	Ν	V			
Х	2 (1.64)			Η					
XI	1 (0.82)	Т							
XII	1 (0.82)			S					
XIII	1(0.82)				Ν				
XIV	1 (0.82)			S	Ν	V			
XV	1 (0.82)		S	S	Ν				

<sup>&</sup>lt;sup>*a*</sup> Alleles are defined as combinations of amino acid mutations. Only mutations retreived more than seven times were used to define the alleles. The wild-type allele (allele I) is arbitrarily defined as the most frequently isolated.

<sup>c</sup> As determined in reference 17.

<sup>&</sup>lt;sup>b</sup> Blank entries indicate identity with the wild type, whereas mutations are indicated.

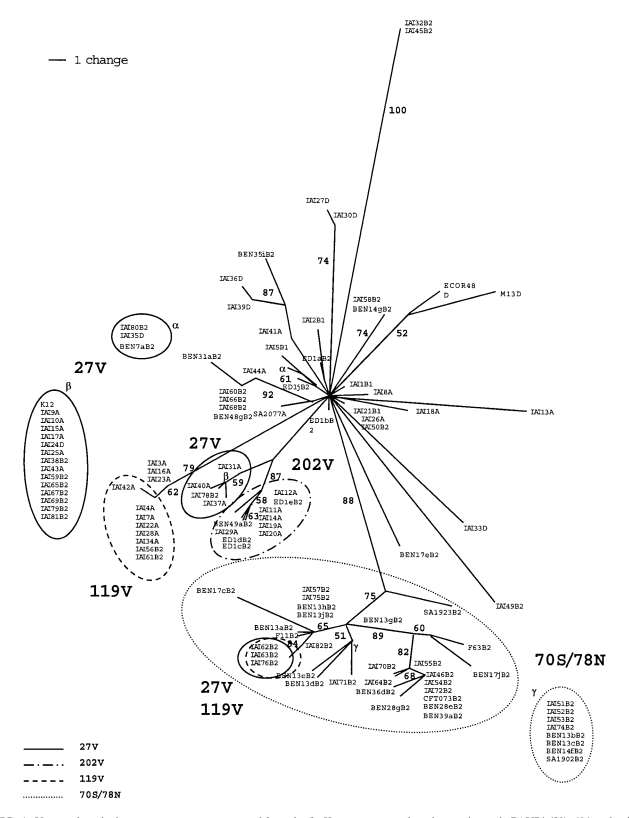


FIG. 1. Unrooted semistrict consensus tree reconstructed from the *fimH* gene sequences by using parsimony in PAUP\* (20); 696 nucleotides, of which 62 are informative for parsimony (l = 241; consistency index, 0.456; retention index, 0.877). The numbers at branches are the bootstrap proportions obtained from 100 replicates. Only bootstraps with proportions above 50% are given. The phylogenetic groups to which the strains belong, determined as described in reference 2, are indicated: A, B1, D, and B2. Similar topology was obtained with a neighbor-joining approach (data not shown). Strains within circles correspond to the strains with the A27V, N70S/S78N, A119V, and A202V mutations. K12 and CFT073 sequences have been extracted from GenBank.

only 7 of those, which were found variable in more than seven strains, were considered in the statistical analyses (Table 1). Except for A202V, all amino acid changes considered here (A27V, G66S/C, N70S, S78N, A119V, V163A) were previously described as polymorphic and potentially pathoadaptive for *E. coli* (15). Fifteen different FimH alleles (I to XV) were found in our collection of natural isolates (Table 1). While most amino acid mutations were widespread among the 15 alleles, the V163A mutation was found only in allele V in combination with both N70S and S78N mutations. Moreover, mutations N70S and S78N were almost always found together (0.865; P < 0.0001) and were the most common amino acid mutations. These two mutations are localized within  $\beta$ -sheet 6 in the three-dimensional structure of FimH (1).

When all strains were considered, significant correlations were found only between N70S/S78N mutations, the B2 phylogenetic group (0.467; P < 0.0001), and lethality in mice (0.316; P < 0.0009). In addition, a link between virulence in mice and B2 phylogenetic group was observed in our data (0.529; P < 0.0001), in accordance with previous works (8, 12). To know whether N70S and S78N mutations were correlated with lethality in mice independently from phylogenetic groups, we looked for correlations between FimH amino acid mutations, origin, and virulence in B2 group strains (68 strains, of which 28 are commensal and 40 are pathogenic). No significant correlation was observed for the N70S/S78N mutations, indicating that the previously observed correlation with lethality in mice was due to the B2 specificity of these mutations and not to the pathogenic nature of the strain.

Among the B2 strains, a highly significant correlation was found between the A27V mutation and UTI strains (0.378; P <0.002). To a lesser extent, significant correlations were also found between the A119V mutation and UTI strains (0.273; P < 0.03) and between the A202V mutation and strains isolated from feces (0.307; P < 0.016). Because of the small sample size used to calculate these last two coefficients, additional B2 strains should be tested to confirm these data. In addition, a correlation was found between the A27V mutation and virulence in mice (0.330; P < 0.009). This mutation is localized in a loop between  $\beta$ -sheets 3 and 4 (1), and previous in vitro binding assays had shown a potential role of A27V mutation in the capacity of the strain to bind to monomannose (17) (Table 1). No correlation was found when non-B2 group strains were analyzed (data not shown). Taken together, these data strongly suggest a pathoadaptive role of the A27V mutation in the natural environment.

Having demonstrated that FimH has both mutations that are linked to the global phylogeny of the strains (N70S/S78N) and mutations that are linked to the environment (A27V, A119V, and A202V), we then compared the *fimH* gene and the strain phylogenies to understand the *fimH* evolutionary history. The unrooted tree reconstructed from the *fimH* sequences showed a grouping that is incongruent with the phylogenetic groups to which the strains belong (Fig. 1), an argument for the occurrence of horizontal gene transfers during *fimH* evolution (3, 5). The mutations separating the monophyletic group of B2 strains supported by a bootstrap value of 88% from the remaining strains, which also encompass some B2 strains (Fig. 1), were determined with MacClade (10). They are not scattered all over the sequence but are clustered between nucleotides 183 and 351 (data not shown), a feature typical of gene transfer (3). It could be assumed that the monophyletic group of B2 strains which possesses the N70S/S78N mutations corresponds to the true B2, whereas the remaining B2 strains have suffered from horizontal gene transfers. In addition, the phylogenetic tree (Fig. 1) indicated that the nucleotide changes corresponding to the A27V and A119V mutations (C80T and C356T, respectively) were each flanked by different sequences (three and two sequences, respectively). Thus, pathoadaptive mutations had arisen several times during the evolution.

This study is the first large-scale analysis of FimH pathoadaptive mutations from a collection of well-characterized natural isolates. Altogether, our data indicated that pathoadaptive mutations are selected for in nature in B2 phylogenetic group strains. Since B2 strains have numerous extraintestinal virulence determinants, virulence can be considered a multifactorial trait as previously suggested (8), with a probably essential effect of the genetic background of the strain in the rise of particular mutations such as pathoadaptive ones. Furthermore, the concept of commensal-adaptive mutation could be proposed for the A202V mutation, which was preferentially present in B2 group strains isolated from feces. It would be interesting to test whether FimH A202V strains adhere better to gut epithelium. Moreover, one could propose that the balancing selection for these patho- and commensal-adaptive mutations could explain the presence of B2 group strains in normal feces (4, 21).

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