

## SHORT COMMUNICATION

# Draft genome sequences of five recent human uropathogenic *Escherichia coli* isolates

Sargurunathan Subashchandrabose<sup>1</sup>, Tracy H. Hazen<sup>2</sup>, David A. Rasko<sup>2</sup> & Harry L. T. Mobley<sup>1</sup>

<sup>1</sup> Department of Microbiology and Immunology, University of Michigan Medical School, Ann Arbor, MI, USA

<sup>2</sup> Department of Microbiology and Immunology, Institute for Genome Sciences, University of Maryland School of Medicine, Baltimore, MD, USA

This publication briefly describes the draft genomes of five recent human uropathogenic (UPEC) *Escherichia coli* isolates. UPEC are of increasing importance to human health. The genomes of these new isolates are clearly and simply described and will be of great utility and interest to this research community.

## Keywords

uropathogenic *Escherichia coli*; draft genomes; type III secretion system.

## Correspondence

Harry L.T. Mobley, Frederick G. Novy  
Professor and Chair, Department of Microbiology and Immunology, University of Michigan Medical School, 5641 Medical Science Bldg. II, 1150 West Medical Center Dr., Ann Arbor, MI 48109-0620, USA.  
Tel.: +1 (734) 764 1466  
fax: +1 (734) 763 7163  
e-mail: hmobley@med.umich.edu

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Urinary tract infections (UTIs) are one of the most common bacterial infections afflicting humans (Russo & Johnson, 2003). Uropathogenic *Escherichia coli* (UPEC) is the etiological agent of a majority of cases of uncomplicated UTIs in otherwise healthy individuals (Hooton, 2012). UPEC are a heterogeneous group of bacteria that are closely related to avian pathogenic *E. coli* and neonatal meningitis *E. coli* (Russo & Johnson, 2000). UPEC is believed to exhibit a commensal-like lifestyle within the gastrointestinal tract and induces pathological changes only upon entry into extra-intestinal sites such as the urinary tract and bloodstream (Russo & Johnson, 2000). Available genomes of UPEC strains reveal a complex mosaic structure encompassing a core genome interrupted by regions that carry the hallmarks of horizontally transferred genetic elements (Welch *et al.*, 2002; Brzuszkiewicz *et al.*, 2006). Frequently, such islands

## Abstract

This study reports the release of draft genome sequences of five isolates of uropathogenic *Escherichia coli* (UPEC), isolated from patients suffering from uncomplicated cystitis in 2012 in Ann Arbor, Michigan. Phylogenetic analyses revealed that these strains belonged to *E. coli* phylogroups B2 and D and are closely related to known UPEC strains. Comparative genomic analysis revealed that more conserved proteins were shared between these recent isolates and UPEC strains causing cystitis than those causing pyelonephritis. Additional genomic comparisons identified that three isolates encode a type III secretion system (T3SS) and a putative T3SS effector gene cluster along with an invasin-like outer membrane protein. The presence of T3SS genes is a rare occurrence among UPEC strains. These genomes further substantiate the heterogeneity of the gene pool of UPEC and provide a foundation for comparative genomic studies using recent clinical isolates.

contain genes that contribute to uropathogenesis and can be considered as pathogenicity islands (Lloyd *et al.*, 2007, 2009).

UPEC remains a major burden on human health and is becoming increasingly recalcitrant to routinely used therapeutic agents (Gupta *et al.*, 2011; Hooton, 2012). Several virulence factors, such as type 1 and P fimbriae, flagella, capsule, and toxins, have been identified in UPEC (Brumbaugh & Mobley, 2012). Multiple fitness mechanisms, including co-opting metabolic enzymes to enable survival and colonization in the mammalian urinary tract, have been delineated in UPEC (Alteri & Mobley, 2012). However, to translate the knowledge on UPEC pathogenesis to develop novel therapeutic and prophylactic agents, a comprehensive understanding of the cues encountered by UPEC during human infection is required (Hagan *et al.*, 2010). In an effort to better define those cues, we are currently profiling the



transcriptomes of UPEC derived directly from patients with clinical UTI. Due to the genetic heterogeneity observed among UPEC strains, we sequenced the genomes of an additional five clinical isolates. The majority of UPEC reference genomes have been derived from isolates that are decades old, and it is possible that human activity, including both intended and unintended exposure to antibiotics, has changed the selective pressures on this bacterium in recent years.

UPEC isolates (HM26, HM27, HM46, HM65, and HM69) were isolated from urine of female patients diagnosed with cystitis at the University of Michigan Health Service clinic. The age of patients ranged from 18 to 25 years, with a median age of 22 years. Briefly, urine samples were cultured in MacConkey agar, and lactose-fermenting colonies were screened using a Vitek 2 system to conclusively identify *E. coli*. Quantitative cultures of urine samples revealed high levels of UPEC bacteriuria ( $> 10^5$  CFU mL<sup>-1</sup>) in all samples. All samples, except HM26, were obtained from patients suffering from isolated instances of UTIs; HM26 was isolated from a patient suffering from recurrent UTI (four episodes in the 6 months preceding sample collection). Antimicrobial susceptibility profile for each isolate was determined using a Vitek 2 system, and HM69 was found to be resistant to trimethoprim/sulfamethoxazole, the primary therapeutic agent for uncomplicated cystitis (Gupta *et al.*, 2011). None of these isolates were resistant to ciprofloxacin, another commonly used primary therapeutic agent to treat UTIs (Gupta *et al.*, 2011). Somatic (O) and flagellar (H) antigen types were determined at the *E. coli* reference center at Pennsylvania State University and are as follows: HM26 (O2:H18), HM27 (O4:H5), HM46 (O166:H15), HM65 (O2:H6/41), and HM69 (O15:H18). These isolates represent a typical diversity of *E. coli* isolated from uncomplicated cystitis in humans in all characteristics.

Genomic DNA was extracted from bacteria grown in lysogeny broth using DNeasy kit (Qiagen). The genome sequence of each isolate was generated at the Institute for Genome Sciences Genome Resource Center (<http://www.igs.umaryland.edu/resources/grc/index.php>) on Illumina HiSeq2500 using paired-end libraries with 300-bp inserts (Table 1). The draft genomes were assembled using both the Velvet assembler (Zerbino & Birney, 2008) with k-mer values determined using VELVETOPTIMISER, version 2.1.4

(<http://www.vicbioinformatics.com/software.velvetoptimiser.shtml>), and the EDENA, version 3, assembler (Hernandez *et al.*, 2008). Contigs from the two assemblies were merged using Minimus (Sommer *et al.*, 2007), and contigs longer than 200 bp were used for further analysis. The resulting genome assemblies contained an average of 125 contigs per genome (range 43–285) (Table 1). Nucleotide sequences were annotated using the Rapid Annotation using Subsystem Technology (RAST) server (Aziz *et al.*, 2008). The numbers of predicted genes from the draft genomes (Table 1) were similar to the previously sequenced *E. coli* genomes with an average of 5165 genes per genome (range 4904–5420). The presence of select known urovirulence factors in these isolates can be found in Table 2.

We probed the phylogenetic relationship between our recent isolates with a collection of representative *E. coli* and *Shigella* strains (Fig. 1) using a whole-genome phylogeny-based approach as previously described (Sahl *et al.*, 2011). Briefly, draft genome sequences were aligned to sequenced reference strains (Fig. 1) using Mugsy (Angiuoli & Salzberg, 2011). Aligned regions were extracted, and a maximum-likelihood phylogenetic tree with 100 bootstrap replicates was inferred from the aligned regions using RAxML, version 7.2.8 (Stamatakis, 2006), and visualized using FIGTREE, version 1.3.1 (<http://tree.bio.ed.ac.uk/software/figtree>). Phylogroups B2 and D encompass most known UPEC strains (Russo & Johnson, 2000), and the isolates identified in this study are also members of these two phylogroups. Three isolates (HM26, HM46, and HM69) cluster with a cystitis strain, UMN026, and an enteroaggregative *E. coli* strain 042 in phylogroup D (Fig. 1), whereas HM27 and HM65 cluster with extensively studied prototypical UPEC strains, CFT073, UTI89, and 536, which are members of phylogroup B2 (Fig. 1). Based on the whole-genome phylogeny, the recent UPEC isolates appear to be similar to previously identified UPEC isolates.

Initial screening of the draft genome sequences for features, not found in previously sequenced strains, revealed that the isolates in phylogroup D, HM26, HM46, and HM69, all contain a significant number of genes encoding proteins involved in plasmid conjugation and transfer functions, suggesting that these isolates harbor plasmids. All three isolates contain an IncF-type machinery,

**Table 1** Sequencing statistics and genome characteristics

Isolate	Phylogroup	No. of reads	Genome size (bp)	%GC	No. of genes*	Sequence coverage	Conserved genes <sup>†</sup>	Divergent genes <sup>‡</sup>
HM26	D	11 660 136	5 271 678	50.64	5199	223	3137	1677
HM27	B2	12 778 406	5 166 851	50.51	5150	250	3158	1640
HM46	D	12 952 066	4 967 159	50.78	4904	263	3084	1456
HM65	B2	11 907 652	5 162 282	50.49	5154	233	3177	1741
HM69	D	11 722 448	5 374 332	50.70	5420	220	3109	1973
Ave		12 204 142	5 188 460	50.62	5165	238	3133	1697

\*Gene numbers are based on RAST.

<sup>†</sup>Number of genes that are conserved (BSR of  $\geq 0.8$ ) among UPEC strains CFT073, 536, F11, UTI89, UMN026, and all the HM isolates.

<sup>‡</sup>Number of genes that are divergent (BSR of  $< 0.8$  to  $> 0.4$ ) among UPEC strains CFT073, 536, F11, UTI89, UMN026, and all the HM isolates.

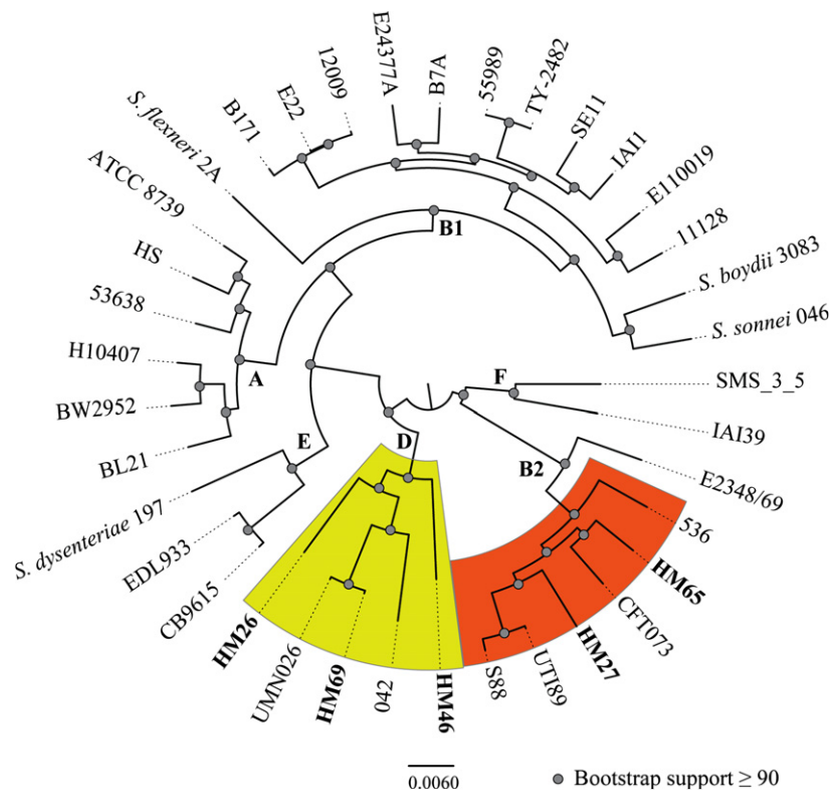
**Table 2** Prevalence of select urovirulence factors

Virulence factor	HM26	HM27	HM46	HM65	HM69
<b>Iron uptake</b>					
Enterobactin R	+	+	+	+	+
Salmochelins R	+	+	–	+	–
Aerobactin R	–	–	–	–	+
Yersiniabactin R	+	+	–	+	+
ChuA, Heme R	+	+	+	+	+
Hma, Heme R	–	+	–	–	–
<b>Toxins</b>					
Hemolysin A	–	+	–	+	–
Cnf	–	–	–	+	–
<b>Fimbriae</b>					
Type 1	+	+	+	+	+
Pap	–	+	–	+	–
F1C	–	+	–	+D	–

R, receptor; D, disrupted; +, found in genome sequence; –, not present in genome sequence.

and HM26 and HM69 encode an additional Incl1 machinery, suggesting that multiple plasmids may be present in these isolates. Further analysis of the genes present in these plasmids is required to elucidate whether these plasmids contain genes involved in antibiotic resistance and virulence.

BLAST score ratio (BSR), an *in silico* approach to conduct comparative proteomic analyses based on proteins predicted to be encoded in a genome (Rasko *et al.*, 2005), was used to compare the proteins encoded in the newly sequenced strains with well-characterized UPEC strains. The BSRs were calculated as the ratio of raw BLASTP score for the query to the raw BLASTP score of the reference strain. BSR cutoffs of  $\geq 0.8$  and  $< 0.8$  to  $> 0.4$  were used to determine whether a gene is conserved or divergent, respectively. A BSR value of 0.8 corresponds to c. 85–90% identity over 90% length of a protein sequence, indicating a highly conserved sequence (Rasko *et al.*, 2005). An average of 3133 proteins were conserved, and



**Fig. 1** Phylogenomic analysis of the five recent UPEC isolates sequenced in this study compared with a collection of diverse *Escherichia coli* and *Shigella* genomes available in the public domain. Genomes used for whole-genome alignments are available in GenBank at the following GenBank accession numbers: EDL933 (NC\_002655), CB9615 (NC\_013941), 11128 (NC\_013364), E2348/69 (NC\_011601), B171 (AAJX00000000), 12009 (NC\_013353), E22 (AAJV00000000), E110019 (AAJW00000000), 53638 (AAKB00000000), HS (NC\_009800), ATCC 8739 (NC\_010468), BL21 (NC\_012947), BW2952 (NC\_012759), H10407 (NC\_017633), TY-2482 (AFOG01000000), *Shigella flexneri* 2A 2457T (NC\_004741), *Shigella sonnei* 046 (NC\_007384), *Shigella boydii* 3083 (NC\_010658), B7A (AAJT02000000), E24377A (NC\_009801), IA11 (NC\_011741), SE11 (NC\_011415), 55989 (NC\_011748), CFT073 (NC\_004431), S88 (NC\_011742), UTI89 (NC\_007946), 536 (NC\_008253), 042 (FN554766), UMN026 (NC\_011751), *Shigella dysenteriae* Sd197 (NC\_007606), IA139 (NC\_0117500), and SMS-3-5 (NC\_010498). The genomes were aligned using Mugsy (Angiuoli & Salzberg, 2011), and a maximum-likelihood phylogeny with 100 bootstrap replicates was inferred using RAxML, version 7.2.8 (Stamatakis, 2006). Bootstrap support at all branch points were  $\geq 90$ .

1697 proteins were divergent between the recent UPEC isolates and the established UPEC strains CFT073, 536, F11, UT189, and UMN026 (Table 1). When compared to cystitis strain F11, these isolates contained 3342 and 1477 proteins that were conserved and divergent, respectively. Three thousand two hundred and thirty-six proteins were conserved, and 1783 proteins were divergent between these isolates and pyelonephritis strain CFT073. Taken together, BSR analysis indicates that the isolates sequenced during this study are more closely related to cystitis strains than to pyelonephritis strains.

Type III secretion system (T3SS) is used by bacteria to inject effectors directly into host cells (Ren *et al.*, 2004). T3SS has been the subject of extensive investigation in enteropathogenic and enterohemorrhagic strains of *E. coli* (Wong *et al.*, 2011). However, T3SS genes are not commonly found in UPEC isolates; a previous study revealed that three of 76 cystitis isolates, collected in Japan, had genes encoding components of a T3SS (Miyazaki *et al.*, 2002). In contrast, in this study including only five isolates, three isolates (HM26, HM46, and HM69) revealed genes that encode the structural components of a T3SS near tRNA *glyU*. These phylogroup D UPEC isolates also contain a putative effector island (*eip* island) adjacent to tRNA *selC* that encodes potential T3SS effectors and an invasin-like outer membrane protein. Regions near *glyU* and *selC* tRNAs are common sites for insertion of horizontally transferred genetic elements. Gene encoding the invasin-like protein is unusually large for a bacterial gene (10 548 bp), and the predicted protein contains 19 repeats of bacterial immunoglobulin-like domain. A PSORTB search indicates that this protein possibly localizes in the outer membrane. Both T3SS structural genes and the *eip* island are reminiscent of the ETT2 locus and the *eip* island found in an enteroaggregative *E. coli* strain 042 (Ren *et al.*, 2004; Sheikh *et al.*, 2006; Chaudhuri *et al.*, 2010) and in a cystitis strain UMN026 (Lescat *et al.*, 2009). The ETT2 locus is distinct from the T3SS found in the locus of enterocyte effacement (*lee*) pathogenicity island in enteropathogenic *E. coli* (Ren *et al.*, 2004). ETT2 genes have been implicated in the pathogenesis of sepsis caused by *E. coli* (Ideses *et al.*, 2005; Ayres *et al.*, 2012). Surprisingly, a UPEC strain was determined as the cause of hemolytic uremic syndrome (Tarr *et al.*, 1996) in a patient, and that isolate exhibited a phenotype typically associated with T3SS-specific effectors. Efforts are under way to test the role of genes encoding T3SS structural and effector proteins and the invasin-like protein in uropathogenesis.

In summary, we present the genome sequences for five recent isolates of UPEC. Many of the genes previously implicated in the pathogenesis of UPEC were identified in these isolates. Our results also reveal that ETT2 genes are found in three of five UPEC strains sequenced during this study. The availability of these additional genome sequences will be a valuable resource to the UPEC research community for further comparative genomic analyses.

Nucleotide sequence accession numbers: This whole-genome shotgun sequencing project has been deposited at DDBJ/EMBL/GenBank under the accession numbers

APNW00000000, APNU00000000, APNY00000000, APNX00000000, and APNV00000000 corresponding to the UPEC isolates HM26, HM27, HM46, HM65, and HM69, respectively. The versions described in this paper are the first version: APNW01000000, APNU01000000, APNY01000000, APNX01000000, and APNV01000000.

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