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1-1-2016

Complete Genome Sequence of a CTX-M-15-Producing Escherichia coli Strain from the H30Rx Subclone of Sequence Type 131 from a Patient with Recurrent Urinary Tract Infections, Closely Related to a Lethal Urosepsis Isolate from the Patient's Sister.

Timothy J. Johnson

Maliha Aziz George Washington University

Cindy M. Liu George Washington University

Evgeni Sokurenko University of Washington

Dagmara I. Kisiela

APA Citation

Johnson, T. J., Aziz, M., Liu, C. M., Sokurenko, E., Kisiela, D. I., Paul, S., Andersen, P. S., Johnson, J. R., & Price, L. B. (2016). Complete Genome Sequence of a CTX-M-15-Producing Escherichia coli Strain from the H30Rx Subclone of Sequence Type 131 from a Patient with Recurrent Urinary Tract Infections, Closely Related to a Lethal Urosepsis Isolate from the Patient's Sister.. *Genome Announcements*, *4* (3). http://dx.doi.org/10.1128/genomeA.00334-16

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Authors

Timothy J. Johnson, Maliha Aziz, Cindy M. Liu, Evgeni Sokurenko, Dagmara I. Kisiela, Sandip Paul, Paal S. Andersen, James R. Johnson, and Lance B. Price

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Complete Genome Sequence of a CTX-M-15-Producing *Escherichia coli* Strain from the H30Rx Subclone of Sequence Type 131 from a Patient with Recurrent Urinary Tract Infections, Closely Related to a Lethal Urosepsis Isolate from the Patient's Sister

Timothy J. Johnson,^a Maliha Aziz,^{b,c} Cindy M. Liu,^{b,c} Evgeni Sokurenko,^d Dagmara I. Kisiela,^d Sandip Paul,^d Paal Andersen,^e James R. Johnson,^f Lance B. Price^{b,c}

Department of Veterinary and Biomedical Sciences, College of Veterinary Medicine, University of Minnesota, St. Paul, Minnesota, USA^a; Department of Environmental and Occupational Health, George Washington University, Washington, District of Columbia, USA^b; Division of Pathogen Genomics, Translational Genomics Research Institute (TGen), Flagstaff, Arizona, USA^c; Department of Microbiology, University of Washington School of Medicine, Seattle, Washington, USA^d; Microbiology and Infection Control, Statens Serum Institut, Copenhagen, Denmark^e; Veterans Affairs Medical Center and University of Minnesota, Minneapolis, Minnesota, USA^f

We report here the complete genome sequence, including five plasmid sequences, of *Escherichia coli* sequence type 131 (ST131) strain JJ1887. The strain was isolated in 2007 in the United States from a patient with recurrent cystitis, whose caregiver sister died from urosepsis caused by a nearly identical strain.

Received 16 March 2016 Accepted 21 March 2016 Published 12 May 2016

Citation Johnson TJ, Aziz M, Liu CM, Sokurenko E, Kisiela DJ, Paul S, Andersen P, Johnson JR, Price LB. 2016. Complete genome sequence of a CTX-M-15-producing *Escherichia coli* strain from the *H*30Rx subclone of sequence type 131 from a patient with recurrent urinary tract infections, closely related to a lethal urosepsis isolate from the patient's sister. Genome Announc 4(3):e00334-16. doi:10.1128/genomeA.00334-16.

Copyright © 2016 Johnson et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license. Address correspondence to Maliha Aziz, mlaziz@owu.edu.

Address correspondence to Maina Aziz, miaziz@gwd.edu.

scherichia coli sequence type 131 (ST131) is currently the most prevalent extraintestinal E. coli lineage (1). Its ST131-H30 sublineage accounts for most E. coli isolates that are fluoroquinolone resistant and produce extended-spectrum β -lactamases (ESBLs) (2). Carriage of *bla*_{CTX-M-15} is associated with the ST131-H30Rx sublineage within ST131-H30 (3). ST131-H30 is associated with recurrent urinary tract infections, pyelonephritis, and urosepsis (1, 4-7). We sequenced two E. coli clinical isolates from two adult sisters, one with recurrent E. coli cystitis, and the other with fatal E. coli urosepsis that developed after caring for her sister. We previously reported the complete genome sequence of the deceased sister's fatal E. coli strain (JJ1886), from the ST131-H30Rx sublineage (8). Here, we present the complete genome sequence of the surviving sister's E. coli strain (JJ1887), which is also from ST131-H30Rx and nearly identical to JJ1886. Whether the sisters' distinct clinical outcomes are attributable to host response differences, strain-specific features, or other factors, the JJ1887 genome represents a valuable added resource for understanding the pathogenicity of the ST131-H30Rx lineage.

JJ1887 was sequenced on the PacBio platform (P5-C3 chemistry), which generated 66,442 raw PacBio reads (mean read length, 10,942 bp; total nucleotides, 764,654,567). The PacBio sequence reads were manually error corrected using Illumina short reads ($125 \times$ coverage, TruSeq chemistry) using CLC Genomics Workbench version 7. The resultant data were assembled using the Hierarchical Genome Assembly Process (HGAP) version 3 in the PacBio single-molecule real-time (SMRT) Portal, which produced a circular chromosome and five closed plasmids, with a mean coverage of $118 \times$. The short-read mapping was further analyzed with Delly (9) and the Northern Arizona SNP Pipeline (NASP) (http://TGenNorth.github.io/NASP). The verified sequences were annotated using Prokka 1.10 (10) and the NCBI Prokaryotic Genome Annotation Pipeline (PGAP). Antimicrobial resistance genes were identified using ResFinder 2.1 (11), and phage regions were identified using PHAST (12).

The complete genome of the JJ1887 chromosome comprises a 5,081,061-bp chromosome with a G+C content of 50.78%. It includes 4,773 coding sequences (CDSs), 89 tRNAs, 22 rRNA features, and 7 intact prophage regions, one of which (57.4 kb) harbors $bla_{\rm CTX-M-15}$.

JJ1887 contains five plasmids, of which pJJ1887-1, -2, and -3 are identical to plasmids in JJ1886 (8). According to Plasmid-Finder 1.3 (13), pJJ1887-1 (1,552 bp) is a Col(MG828) plasmid with two CDSs; plasmid pJJ1887-2 (5,167 bp) is a Col156 plasmid with five CDSs; pJJ1887-3 (5,631 bp) is a ColE1-like plasmid with five CDSs; pJJ1887-4 (107,507 bp) has both RepFIA and Rep-FII replicons, with 125 CDSs; and pJJ1887-5 (130,603 bp) has both RepFIB and RepFII replicons, with 158 CDSs. *bla*_{TEM-1B}, *bla*_{OXA-1} in pJJ1887-4 and *bla*_{TEM-1B} in pJJ1887-5 confer resistance to β -lactams, and the *aac*(6')-*Ib-cr* gene in pJJ1887-4 codes for fluoroquinolone resistance. pJJ1887-5 also carries genes conferring resistance to aminoglycosides [*aac-(3)-IIa*], macrolides [*mph*(*A*)], tetracyclines [*tet*(*B*)], trimethoprim (*dfrB4*), sulfonamides (*sul1*), and fluoroquinolones (*qepA*).

Nucleotide sequence accession numbers. The complete sequences of the chromosome and plasmids of *E. coli* JJ1887 have been deposited in GenBank under accession numbers CP014316 to CP014321.

ACKNOWLEDGMENTS

This material is based upon work supported by the Office of Research and Development, Medical Research Service, Department of Veterans Affairs,

grant 1 I01 CX000192 01 (to J.R.J.); NIH grant RC4-AI092828 (to E.S. and J.R.J.) and R01AI106007 (to E.S.); the TGen Foundation (to L.B.P. and M.A.); the Statens Serum Institut (to P.A.); the Augustinus Fonden (to P.A.); and USAMRMC grant W81XWH-10-1-0753 (to L.B.P.).

Patricia Stogsdill, Lonny Yarmus, Rob Owens, John Quinn, and Karen Lolans facilitated the provision of *E. coli* JJ1887 and associated clinical data. Computational resources for PacBio assembly were provided through the Minnesota Supercomputing Institute.

The opinions presented here are those of the authors and do not represent the official policy or position of the U.S. Government or any of its agencies.

FUNDING INFORMATION

This work, including the efforts of James R. Johnson, was funded by Office of Research and Development, Medical Research Service, Department of Veterans Affairs (1101CX000192). This work, including the efforts of Maliha Aziz and Lance B. Price, was funded by TGEN Foundation. This work, including the efforts of Lance B. Price, was funded by USAMRMC. This work, including the efforts of Paal Skytt Andersen, was funded by Statens Serum Institut. This work, including the efforts of Lance B. Price, was funded by USAMRMC (W81XWH-10-1-0753). This work, including the efforts of Evgeni Sokurenko and James R. Johnson, was funded by HHS | National Institutes of Health (NIH) (RC4-AI092828). This work, including the efforts of Paal Skytt Andersen, was funded by HHS | National Institutes of Health (NIH) (R01AI106007). This work, including the efforts of Paal Skytt Andersen, was funded by Augustinus Fonden (Augustinus Foundation).

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