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Complete Closed Genome Sequence of Nontoxigenic Invasive Corynebacterium diphtheriae bv. mitis Strain ISS 3319

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ABSTRACT The genome sequence of the human pathogen *Corynebacterium diphtheriae* bv. mitis strain ISS 3319 was determined and closed in this study. The genome is estimated to have 2,404,936 bp encoding 2,257 proteins. This strain also possesses a plasmid of 1,960 bp.

corynebacterium diphtheriae is the etiological agent of diphtheria, a toxigenic infection of the upper respiratory tract. Besides diphtheria, systemic infections caused by nontoxigenic strains have been reported (1–5). The molecular basis of toxin-independent pathogenicity of *C. diphtheriae* is poorly understood (6). Therefore, we sequenced the whole genome of *C. diphtheriae* bv. mitis ISS 3319, isolated from the throat of 9-year-old patient (7, 8). The strain lacks the tox gene but is able to adhere to and invade human epithelial cells and effectively colonize spleen and kidneys of mice (9–11).

Sequencing of libraries was carried out with the Oxford Nanopore MinION instrument and the Illumina HiSeq platform using a 250-bp paired-end protocol. The Illumina and MinION reads were assembled together using Unicycler with Pilon polishing, with a final hybrid assembly of two contigs of 1,604,547 and 800,125 bp. Two gaps of 124 bp and 140 bp were identified in the genome using BLAST searches of >500 bp from the ends of both the contigs against the published assembly of the same strain (GenBank accession no. JAQN00000000). The contigs were merged using the missing sequence after aligning them to the genome sequence of strain C7 (beta) (GenBank accession no. NC_016801), which belongs to the same sequence type, ST26, as ISS 3319 (7). Automatic annotation and identification of rRNAs and tRNAs were accomplished using the Prokka version 1.11 software tool (12).

The new sequence was compared to the previously published assembly of ISS 3319, which was also reannoated using Prokka to obtain an equivalence of annotation for comparative analysis. The annotated genomes were compared using Roary (13, 14); 2,195 genes are conserved between the assemblies, and only four single nucleotide polymorphisms (SNPs) were observed in the core genomic alignment. Most of the 26 genes reported to be specific to the previous assembly (GenBank accession no. JAQN00000000) showed significant similarities to several genes in the new assembly and may not be specific. The proteins encoded by the majority of these genes are short and fall within or near the identified genomic islands that were identified using IslandViewer 4 (15).

The closed genome of 2,404,936 bp contained 9 rRNAs, 53 tRNAs, 1 transfermessenger RNA, 1 repeat region, 2,257 protein-coding sequences, and 53.46% GC **Received** 15 December 2017 **Accepted** 19 December 2017 **Published** 1 February 2018

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content. Moreover, a circular plasmid of 1,960 bp with four coding sequences was also identified. A discontinuous mega-BLAST search of this plasmid sequence in GenBank revealed partial similarities (67% identity in 612-bp alignment) with plasmid pTA144 in *Moraxella* sp. (GenBank accession no. NC_001316) and the metamobilome of an uncultured strain (65% identity in 833-bp alignment; GenBank accession no. LN853682). A nonredundant translation alignment of the plasmid sequence showed a conserved domain with homology to the phage replication protein Cri, described as essential for DNA replication in *Vibrio cholerae* serotype O1 (16), and a sequence with 88% identity to a gene encoding an $\alpha\beta$ -hydrolase from a *Streptococcus* sp. (GenBank accession no. WP_002883881). Coinfections of *C. diphtheriae* with other pathogens, such as *Staphylococcus* spp., *Streptococcus* spp., *Arcanobacterium* spp., and *Pseudomonas* spp., were reported previously (8, 17, 18), and this close contact might support horizontal DNA transfer.

Accession number(s). The *C. diphtheriae* bv. mitis ISS 3319 whole-genome shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession no. CP025209 and CP025210. The versions described in this paper are the second versions.

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REFERENCES

- Burkovski A. 2014. Diphtheria and its etiological agents, p 1–14. In Burkovski A (ed), Corynebacterium diphtheriae and related toxigenic species. Springer, Dordrecht, The Netherlands.
- Belchior E, Henry S, Badell E, Collet L, Benoit-Cattin T, de Montera AM, Guiso N, Patey O, Levy-Bruhl D, Filleul L, Chieze F, Olivier S. 2017. Diphtheria in Mayotte, 2007–2015. Emerg Infect Dis 23:1218–1220. https://doi.org/10 .3201/eid2307.170262.
- Kolios AGA, Cozzio A, Zinkernagel AS, French LE, Kündig TM. 2017. Cutaneous Corynebacterium infection presenting with disseminated skin nodules and ulceration. Case Rep Dermatol 9:8–12. https://doi.org/10.1159/000476054
- Santos LS, Sant'Anna LO, Ramos JN, Ladeira EM, Stavracakis-Peixoto R, Borges LL, Santos CS, Napoleão F, Camello TCF, Pereira GA, Hirata R, Vieira VV, Cosme LMSS, Sabbadini PS, Mattos-Guaraldi AL. 2015. Diphtheria outbreak in Maranhão, Brazil: microbiological, clinical and epidemiological aspects. Epidemiol Infect 143:791–798. https://doi.org/10.1017/ S0950268814001241.
- Gomes DL, Martins CA, Faria LM, Santos LS, Santos CS, Sabbadini PS, Souza MC, Alves GB, Rosa AC, Nagao PE, Pereira GA, Hirata R, Mattos-Guaraldi AL. 2009. Corynebacterium diphtheriae as an emerging pathogen in nephrostomy catheter-related infection: evaluation of traits associated with bacterial virulence. J Med Microbiol 58:1419–1427. https:// doi.org/10.1099/jmm.0.012161-0.
- Sangal V, Hoskisson PA. 2016. Evolution, epidemiology and diversity of Corynebacterium diphtheriae: new perspectives on an old foe. Infect Genet Evol 43:364–370. https://doi.org/10.1016/j.meegid.2016.06.024.
- Sangal V, Blom J, Sutcliffe IC, von Hunolstein C, Burkovski A, Hoskisson PA. 2015. Adherence and invasive properties of *Corynebacterium diphtheriae* strains correlates with the predicted membrane-associated and secreted proteome. BMC Genomics 16:765. https://doi.org/10.1186/s12864-015-1980-8.
- von Hunolstein C, Alfarone G, Scopetti F, Pataracchia M, La Valle R, Franchi F, Pacciani L, Manera A, Giammanco A, Farinelli S, Engler K, De Zoysa A, Efstratiou A. 2003. Molecular epidemiology and characteristics of Corynebacterium diphtheriae and Corynebacterium ulcerans strains isolated in Italy during the 1990s. J Med Microbiol 52:181–188. https:// doi.org/10.1099/jmm.0.04864-0.
- 9. Ott L, Höller M, Gerlach RG, Hensel M, Rheinlaender J, Schäffer TE,

- Burkovski A. 2010. *Corynebacterium diphtheriae* invasion-associated protein (DIP1281) is involved in cell surface organization, adhesion and internalization in epithelial cells. BMC Microbiol 10:2. https://doi.org/10.1186/1471-2180-10-2.
- Ott L, Scholz B, Höller M, Hasselt K, Ensser A, Burkovski A. 2013. Induction of the NFκ-B signal transduction pathway in response to *Corynebacte-rium diphtheriae* infection. Microbiology 159:126–135. https://doi.org/10.1099/mic.0.061879-0.
- Puliti M, Von Hunolstein C, Marangi M, Bistoni F, Tissi L. 2006. Experimental model of infection with non-toxigenic strains of *Corynebacterium diphtheriae* and development of septic arthritis. J Med Microbiol 55:229–235. https://doi.org/10.1099/jmm.0.46135-0.
- Seemann T. 2014. Prokka: rapid prokaryotic genome annotation. Bioinformatics 30:2068–2069. https://doi.org/10.1093/bioinformatics/btu153.
- 13. Tange O. 2011. GNU parallel—the command-line power tool. Login: the USENIX magazine 36:42–47.
- Page AJ, Cummins CA, Hunt M, Wong VK, Reuter S, Holden MT, Fookes M, Falush D, Keane JA, Parkhill J. 2015. Roary: rapid large-scale prokaryote pan genome analysis. Bioinformatics 31:3691–3693. https://doi.org/10.1093/bioinformatics/btv421.
- Bertelli C, Laird MR, Williams KP, Simon Fraser University Research, Lau BY, Hoad G, Winsor GL, Brinkman FS. 2017. IslandViewer 4: expanded prediction of genomic islands for larger-scale datasets. Nucleic Acids Res 45:W30–W35. https://doi.org/10.1093/nar/gkx343.
- Heidelberg JF, Eisen JA, Nelson WC, Clayton RA, Gwinn ML, Dodson RJ, Haft DH, Hickey EK, Peterson JD, Umayam L, Gill SR, Nelson KE, Read TD, Tettelin H, Richardson D, Ermolaeva MD, Vamathevan J, Bass S, Qin H, Dragoi I, Sellers P, McDonald L, Utterback T, Fleishmann RD, Nierman WC, White O, Salzberg SL, Smith HO, Colwell RR, Mekalanos JJ, Venter JC, Fraser CM. 2000. DNA sequence of both chromosomes of the cholera pathogen Vibrio cholerae. Nature 406:477–483. https://doi.org/10.1038/35020000.
- Gordon CL, Fagan P, Hennessy J, Baird R. 2011. Characterization of Corynebacterium diphtheriae isolates from infected skin lesions in the Northern Territory of Australia. J Clin Microbiol 49:3960–3962. https://doi.org/10.1128/JCM.05038-11.
- May ML, McDougall RJ, Robson JM. 2014. Corynebacterium diphtheriae and the returned tropical traveler. J Travel Med 21:39–44. https://doi.org/10.1111/jtm.12074.

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