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## Draft Genome Sequence of *Clostridium difficile* Belonging to Ribotype 018 and Sequence Type 17

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Clostridium difficile, belonging to ribotype 018 (RT018), is one of the most prevalent genotypes circulating in hospital settings in Italy. Here, we report the draft genome of C. difficile CD8-15 belonging to RT018, isolated from a patient with fatal C. difficileassociated infection.

Received 5 July 2016 Accepted 6 July 2016 Published 1 September 2016

Citation Riccobono E, Di Pilato V, Della Malva N, Meini S, Ciraolo F, Torricelli F, Rossolini GM. 2016. Draft genome sequence of Clostridium difficile belonging to ribotype 018 and sequence type 17. Genome Announc 4(5):e00907-16. doi:10.1128/genomeA.00907-16.

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"lostridium difficile infection (CDI) is one of the most common hospital-acquired infections and the leading cause of antibiotic-associated diarrhea and pseudomembranous colitis (1). Increasing CDI incidence rates have been mainly attributed to successful clones belonging to a few predominant ribotypes (RTs), with different distribution in different geographic regions (2, 3).

RT018 has been reported as one of the most prevalent genotypes circulating in hospital settings in Italy (4, 5) and in the Far East (i.e., South Korea and Japan) (6, 7) but has been rarely described in other countries (2, 8). RT018 is highly transmissible and generally shows a multidrug-resistant phenotype. It produces both toxin A and toxin B, but is negative for binary toxin, and is associated with complicated outcomes (4, 7, 9). In this report, we announce the draft genome of C. difficile CD8-15 belonging to RT018, which was isolated from a patient admitted to a hospital in Florence (central Italy) in 2015 with fatal C. difficile-associated infection evolved as toxic megacolon.

C. difficile CD8-15 was grown in 5 ml of thioglicollate broth and incubated under anaerobic conditions for 48 h at 37°C. Genomic DNA was subjected to whole-genome sequencing with the MiSeq platform (Illumina Inc., San Diego, CA, USA) using a 2×300-bp paired-end approach. A total of 2,118,990 reads were generated and then de novo assembled using SPAdes version 3.6.1 (10) into 44 scaffolds (largest scaffold = 831,956 bp;  $N_{50}$  = 235,991 bp;  $L_{50} = 5$ ; average GC = 28.63%), with an estimated genome size of 4,249,791 bp and an average coverage of  $50\times$ .

The draft genome was then subjected to the following in silico analyses: (i) multilocus sequence typing (MLST; http://pubmlst .org/cdifficile); (ii) toxinotyping (http://www.mf.uni-mb.si /mikro/tox); (iii) detection of antibiotic resistance mechanisms, using the C. difficile 630 strain (11). The presence of phage-related sequences and CRISPR-Cas systems were investigated using PHAST (http://phast.wishartlab.com) and CRISPRFinder (http: //crispr.u-psud.fr), respectively.

MLST analysis assigned C. difficile CD8-15 to sequence type

(ST) 17, as previously reported for other C. difficile strains belonging to RT018 (12, 13). ST17 and other STs of C. difficile belong to clade I (12). Toxinotyping classified C. difficile CD8-15 as a nonvariant strain (toxinotype 0) producing toxin A and toxin B but not binary toxin CDT.

C. difficile CD8-15 was resistant to rifampin (MIC > 32 mg/L), levofloxacin (MIC >32 mg/L), and erythromycin (MIC >256 mg/L) but remained susceptible to clindamycin (MIC 2 mg/L). Sequence analysis revealed the presence of point mutations in the rpoB gene (R505K and I548M), which is known to be associated with high-level resistance to rifamycins (14). Mutations were detected also in the quinolone resistance-determining region of gyrA (T82I), which is consistent with the fluoroquinolones resistance phenotype. Resistance to erythromycin was likely attributable to the presence of a multidrug efflux system encoded by the cme gene (15), while erm genes and other genes involved in macrolides resistance (i.e., mef, msr, cfr) were not detected (9, 16). PHAST identified 3 intact and 5 incomplete prophage regions. CRISPRFinder identified two class I CRISPR-Cas systems.

Accession number(s). This whole-genome shotgun project has been deposited at DDBJ/EMBL/GenBank under the accession number LYDP00000000.

## **ACKNOWLEDGMENTS**

We thank Raffaele Laureano, Christian Adamo, and Chiara Barchielli for their contribution.

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