## **UC** Irvine

## **UC Irvine Previously Published Works**

#### **Title**

Genome Sequence of Erysipelothrix sp. Strain Poltava, Isolated from Acute Septic Erysipelas of Swine in Ukraine.

#### **Permalink**

https://escholarship.org/uc/item/11g428q2

### Journal

Microbiology Resource Announcements, 11(8)

#### **Authors**

Tarasov, Oleksandr Kovalenko, Ganna Muzykina, Larysa et al.

#### **Publication Date**

2022-08-18

#### DOI

10.1128/mra.00438-22

Peer reviewed







# Genome Sequence of *Erysipelothrix* sp. Strain Poltava, Isolated from Acute Septic Erysipelas of Swine in Ukraine

Oleksandr Tarasov, a Ganna Kovalenko, a,b,c Larysa Muzykina, a Maksym Bezymennyi, a Eric Bortz, a,b Devin M. Drown<sup>d,e</sup>

<sup>a</sup>National Academy of Agrarian Sciences of Ukraine, Institute for Veterinary Medicine (IVM), Kyiv, Ukraine

**ABSTRACT** The complete genome of *Erysipelothrix* sp. strain Poltava, isolated from fatal acute septic erysipelas of swine in Ukraine, was assembled using Nanopore sequences. One circular chromosome of 1,794,858 bp ( $N_{50}$ , 1,794,858 bp) encodes 16 putative antibiotic resistance genes and secreted virulence factors, highlighting the risk of cross-species livestock and human infection.

s part of a scientific initiative to understand differential swine infections that cocirculate with African swine fever, in Ukraine (1), we sequenced the genome of an *Erysipelothrix* sp. strain isolated in 2006 from domestic swine with an acute septicemic form of swine erysipelas from a backyard farm in Ukraine (Poltava Oblast, Shyshaky Raion; 49°52′40.5948″N, 34°3′51.5262″E) (2). The sequencing of this strain using an Oxford Nanopore Technologies (ONT) MinION platform in veterinary laboratories in Ukraine represents a genomic exploration of a collection of archived bacterial isolates, providing historical and contemporary insight into circulating livestock pathogens (3–5).

This isolate was collected from tissue samples from swine mesenteric lymphatic nodes and spleen in 2006 and cultured in brain heart infusion (BHI) agar (M1611; HiMedia) and in selective medium, modified blood-azide medium (CM0259; Oxoid). The culture was incubated under aerobic conditions at 37°C for 24 h. Biochemical tests were performed using the API Coryne test (BioMeriex, France) (6), and PCR was performed as a confirmatory test using the primers ER1 and ER2 (7), identifying this isolate as *Erysipelothrix rhusiopathiae* (class *Erysipelotrichia*, phylum *Firmicutes*), a Gram-positive rod bacterium that often presents as erysipelas and in severe cases causes acute septicemia, or chronic endocarditis with polyarthritis, leading to severe wasting disease in swine (8). The stock culture was stored lyophilized at  $-20^{\circ}$ C, subcultured, and relyophilized every 24 months. After growth for 24 h in BHI agar, a colony was resuspended in 200  $\mu$ L PBS, incubated for 1.5 h at 37°C, followed by washing, and resuspended in demineralized water. Three sample vials from the same isolate were pooled under biosafety level 2 (BSL2) for DNA isolation, using a DNeasy UltraClean microbial kit (Qiagen).

We used 495 ng of DNA as input for a rapid sequencing library (SQK-RAD004; ONT) and sequenced it on an R9.4.1 flow cell (FLO-MIN106; flow cell ID FAL31485) for 16 h using a MinION Mk1B device. We base called the raw data using Guppy v6.1.3 (ONT) in super-accuracy mode (-c dna\_r9.4.1\_450bps\_sup.cfg), filtering reads with a quality below 10 (-min\_qscore 10) for an output of 1,080,139,073 bp in 1,047,659 reads with a read length  $N_{50}$  value of 2,654 bp. When accounting for only reads passing the quality filter, the run generated 770,677,927 bp in 524,171 reads with a read length  $N_{50}$  value of 3,130 bp and a median Q score of 12.4. We used Filtlong v0.2.0 (https://github.com/rrwick/Filtlong) to subset 50% of the reads (-keep\_percent 50) and prioritize them by read quality (-mean\_q\_weight

**Editor** Catherine Putonti, Loyola University Chicago

Copyright © 2022 Tarasov et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

Address correspondence to Oleksandr Tarasov, tarasovaleksandr003@gmail.com, or Devin M. Drown, dmdrown@alaska.edu.

The authors declare no conflict of interest.

Received 3 May 2022 Accepted 20 July 2022 Published 2 August 2022

<sup>&</sup>lt;sup>b</sup>Department of Biological Sciences, University of Alaska Anchorage, Anchorage, Alaska, USA

<sup>&</sup>lt;sup>c</sup>Division of Virology, Department of Pathology, University of Cambridge, Cambridge, United Kingdom

<sup>&</sup>lt;sup>d</sup>Department of Biology and Wildlife, University of Alaska Fairbanks, Fairbanks, Alaska, USA

elnstitute of Arctic Biology, University of Alaska Fairbanks, Fairbanks, Alaska, USA

**TABLE 1** Antimicrobial resistance genes

AMR mechanism	Gene(s)
Antibiotic target in susceptible species	Alr, Ddl, EF-G, EF-Tu, folA, Dfr, gyrA, gyrB, Iso-tRNA, MurA, rpoB, rpoC, S10p, S12p
Gene conferring resistance via absence	gidB
Protein altering cell wall charge, conferring antibiotic resistance	PgsA

10). After filtering, we had 385,339,219 bp in 132,040 reads with a read length  $N_{50}$  value of 4,664 bp and a median Q score of 14.5.

We assembled the genome *de novo* using Flye v2.9 (9) with the 385-Mb quality-controlled data set (coverage, 213×), specifying high-quality Nanopore reads (–nano-hq). Our initial assembly contained one contig, identified as circular using Flye. We corrected the assembly using Medaka v1.6.0 (https://github.com/nanoporetech/medaka), specifying base-caller model (-m r941\_min\_sup\_g507). Our 1,794,858-bp ( $N_{50}$ , 1,794,858 bp) polished assembly consists of a single circular contig (GC content, 36.39%). We rotated the circular genome start position to *dnaA* using Circlator v1.5.5 (10). Default parameters were used for all software unless otherwise specified.

During the data submission pipeline, the genome deposited at GenBank was annotated using PGAP v6.1 (11–13) and contained 53 tRNAs, 12 rRNAs, and 2,436 coding DNA sequences (CDS). CheckM (14) reported 86.94% completeness with 0.96% contamination. PATRIC v3.6.12 (15, 16) identified 16 antibiotic resistance genes (Table 1). Using the Comprehensive Genome Analysis service in PATRIC, a phylogenetic analysis found the isolate to be similar to members of the genus *Erysipelothrix*. We used FastANI (17) to calculate an average nucleotide identity of 98.86% to the most similar genome, *Erysipelothrix rhusiopathiae* strain Fujisawa (GenBank accession no. NC\_015601) (8). We extracted a consensus sequence of the four full-length 16S rRNA gene copies to use as a query with blastn (18) against the NCBI 16S rRNA database. We found 99% identity to both *Erysipelothrix piscisicarius* strain 15TAL0474 (NR\_170394) and *Erysipelothrix rhusiopathiae* strain ATCC 19414 (NR\_040837). Therefore, we designated this isolate *Erysipelothrix* sp. strain Poltava.

**Data availability.** This whole-genome sequencing project has been deposited at GenBank under the accession no. CP096542.1. The raw data for this project can be found under SRA accession no. SRR18770851 and BioProject accession no. PRJNA827134.

#### **ACKNOWLEDGMENTS**

We thank Natalia Gudz, Serhii Nychyk, and Ihor Halka at NAAS IVM (Kyiv).

The research reported herein was supported in part by the U.S. Defense Threat Reduction Agency (DTRA) Biological Threat Reduction Program in Ukraine (BTRP), through a subaward to the University of Alaska, and by an NIH NIGMS Institutional Development Award (IDeA), grant no. P20GM103395 (Alaska INBRE). We also acknowledge the generous support of the Institute of Arctic Biology at UAF and the UAA College of Arts & Sciences.

The contents of this publication are the responsibility of the authors and do not necessarily reflect the views of the DoD, NIH, or the U.S. Government.

#### **REFERENCES**

- Kovalenko G, Ducluzeau A-L, Ishchenko L, Sushko M, Sapachova M, Rudova N, Solodiankin O, Gerilovych A, Dagdag R, Redlinger M, Bezymennyi M, Frant M, Lange CE, Dubchak I, Mezhenskyi AA, Nychyk S, Bortz E, Drown DM. 2019. Complete genome sequence of a virulent African swine fever virus from a domestic pig in Ukraine. Microbiol Resour Announc 8:e00883-19. https://doi.org/10.1128/ MRA.00883-19.
- Pinchuk NG, Golovko AN, Garkavenko TA. 2019. Analysis of the epizootic situation of the swine erysipelas on the territory of Ukraine for 2006–2017. Vet Biotechnol 34:108–118. https://doi.org/10.31073/vet\_biotech34-13.
- Arefiev V, Kovalenko G, Frant M, Chumachenko T, Polyvianna Y, Pivnenko S, Bolotin V, Mayboroda O, Solodiankin O, Tarasov O, Bezyemenni M, Lyon C, Redlinger M, Sapachova M, Mezhenskyi AA, Ducluzeau A-L, Bortz E, Gerilovych A, Drown DM. 2020. Complete genome sequence of Salmonella enterica subsp. enterica serovar Kottbus strain Kharkiv, isolated from a commercial pork
- production facility in Ukraine. Microbiol Resour Announc 9:e01171-20. https://doi.org/10.1128/MRA.01171-20.
- Bolotin V, Kovalenko G, Marchenko N, Solodiankin O, Rudova N, Kutsenko V, Bortz E, Gerilovych A, Drown DM. 2021. Complete genome sequence of Brucella abortus 68, isolated from aborted fetal sheep in Ukraine. Microbiol Resour Announc 10:e01436-20. https://doi.org/10.1128/MRA.01436-20.
- Rudova N, Buttler J, Kovalenko G, Sushko M, Bolotin V, Muzykina L, Zinenko O, Stegniy B, Dunaiev Y, Sytiuk M, Gerilovych A, Drown DM, Bortz E, Solodiankin O. 2022. Genetic diversity of porcine circovirus 2 in wild boar and domestic pigs in Ukraine. Viruses 14:924. https://doi.org/ 10.3390/v14050924.
- Soto A, Zapardiel J, Soriano F. 1994. Evaluation of API Coryne system for identifying coryneform bacteria. J Clin Pathol 47:756–759. https://doi.org/ 10.1136/jcp.47.8.756.

- Shimoji Y, Mori Y, Hyakutake K, Sekizaki T, Yokomizo Y. 1998. Use of an enrichment broth cultivation-PCR combination assay for rapid diagnosis of swine erysipelas. J Clin Microbiol 36:86–89. https://doi.org/10.1128/JCM.36 186-89 1998
- Ogawa Y, Ooka T, Shi F, Ogura Y, Nakayama K, Hayashi T, Shimoji Y. 2011.
   The genome of Erysipelothrix rhusiopathiae, the causative agent of swine erysipelas, reveals new insights into the evolution of Firmicutes and the organism's intracellular adaptations. J Bacteriol 193:2959–2971. https://doi.org/10.1128/JB.01500-10.
- Kolmogorov M, Yuan J, Lin Y, Pevzner PA. 2019. Assembly of long, errorprone reads using repeat graphs. Nat Biotechnol 37:540–546. https://doi .org/10.1038/s41587-019-0072-8.
- Hunt M, Silva ND, Otto TD, Parkhill J, Keane JA, Harris SR. 2015. Circlator: automated circularization of genome assemblies using long sequencing reads. Genome Biol 16:294. https://doi.org/10.1186/s13059-015-0849-0.
- Tatusova T, DiCuccio M, Badretdin A, Chetvernin V, Nawrocki EP, Zaslavsky L, Lomsadze A, Pruitt KD, Borodovsky M, Ostell J. 2016. NCBI prokaryotic genome annotation pipeline. Nucleic Acids Res 44:6614–6624. https://doi.org/10.1093/ nar/gkw569.
- Haft DH, DiCuccio M, Badretdin A, Brover V, Chetvernin V, O'Neill K, Li W, Chitsaz F, Derbyshire MK, Gonzales NR, Gwadz M, Lu F, Marchler GH, Song JS, Thanki N, Yamashita RA, Zheng C, Thibaud-Nissen F, Geer LY, Marchler-Bauer A, Pruitt KD. 2018. RefSeq: an update on prokaryotic genome annotation and curation. Nucleic Acids Res 46:D851–D860. https://doi.org/10.1093/nar/gkx1068.
- Li W, O'Neill KR, Haft DH, DiCuccio M, Chetvernin V, Badretdin A, Coulouris G, Chitsaz F, Derbyshire MK, Durkin AS, Gonzales NR, Gwadz M, Lanczycki CJ,

- Song JS, Thanki N, Wang J, Yamashita RA, Yang M, Zheng C, Marchler-Bauer A, Thibaud-Nissen F. 2021. RefSeq: expanding the prokaryotic genome annotation pipeline reach with protein family model curation. Nucleic Acids Res 49: D1020–D1028. https://doi.org/10.1093/nar/qkaa1105.
- Parks DH, Imelfort M, Skennerton CT, Hugenholtz P, Tyson GW. 2015. CheckM: assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes. Genome Res 25:1043–1055. https://doi.org/10.1101/gr.186072.114.
- 15. Davis JJ, Wattam AR, Aziz RK, Brettin T, Butler R, Butler RM, Chlenski P, Conrad N, Dickerman A, Dietrich EM, Gabbard JL, Gerdes S, Guard A, Kenyon RW, Machi D, Mao C, Murphy-Olson D, Nguyen M, Nordberg EK, Olsen GJ, Olson RD, Overbeek JC, Overbeek R, Parrello B, Pusch GD, Shukla M, Thomas C, VanOeffelen M, Vonstein V, Warren AS, Xia F, Xie D, Yoo H, Stevens R. 2020. The PATRIC Bioinformatics Resource Center: expanding data and analysis capabilities. Nucleic Acids Res 48:D606–D612. https://doi.org/10.1093/nar/gkz943.
- Brettin T, Davis JJ, Disz T, Edwards RA, Gerdes S, Olsen GJ, Olson R, Overbeek R, Parrello B, Pusch GD, Shukla M, Thomason JA, Ill, Stevens R, Vonstein V, Wattam AR, Xia F. 2015. RASTtk: a modular and extensible implementation of the RAST algorithm for building custom annotation pipelines and annotating batches of genomes. Sci Rep 5:8365. https://doi.org/10.1038/srep08365.
- Jain C, Rodriguez-R LM, Phillippy AM, Konstantinidis KT, Aluru S. 2018. High throughput ANI analysis of 90K prokaryotic genomes reveals clear species boundaries. Nat Commun 9:5114. https://doi.org/10.1038/s41467-018-07641-9.
- Zhang Z, Schwartz S, Wagner L, Miller W. 2000. A greedy algorithm for aligning DNA sequences. J Comput Biol 7:203–214. https://doi.org/10.1089/ 10665270050081478.