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Recommended Citation  
Kersulyte, Dangeruta; Bertoli, M. Teresita; Tamma, Sravya; Keelan, Monika; Munday, Rachel; Geary, Janis; Veldhuyzen van Zanten, Sander; Goodman, Karen J.; and Berg, Douglas E., "Complete genome sequences of two Helicobacter pylori strains from a Canadian Arctic Aboriginal community." *Genome Announcements*. 3,2. e00209-15. (2015).  
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Complete Genome Sequences of Two Helicobacter pylori Strains from a Canadian Arctic Aboriginal Community

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We report here the complete genome sequences of two Amerind Helicobacter pylori strains from Aklavik, Northwest Territories, Canada. One strain contains extra iron-cofactored urease genes and ~140 rearrangements in its chromosome relative to other described strains (typically differing from one another by <10 rearrangements), suggesting that it represents a novel lineage of \textit{H. pylori}.

\textbf{Helicobacter pylori} is a genetically diverse gastric pathogen that chronically infects billions of people worldwide (1–3), including many residents of indigenous Arctic communities (4). Different sets of genotypes tend to predominate in different geographic regions or with different ethnic groups. Although strains from diverse peoples, including South American Amerindians, have been sequenced, no Arctic strains have been sequenced to date. Here, we report the complete genome sequences of two \textit{H. pylori} strains from Aklavik, Northwest Territories, a Canadian Aboriginal community (~600 residents; 93% of Inuvialuit or Gwich’in ethnicity) with a high prevalence of severe \textit{H. pylori}-associated gastritis (5–7).

Preliminary random amplified polymorphic DNA (RAPD) fingerprinting (8) placed 52 of 57 isolates into just six distinct groups, a homogeneity typical of isolated communities (9). Multilocus sequencing typing (MLST) (9) identified four RAPD groups and the five unique strains (43 isolates) as Amerind (more related to Amazonian than even East Asian strains). The other two RAPD groups exhibited European and hybrid Amerind/European MLSTs (4 and 10 isolates, respectively).

We completely sequenced (454 FLX with PCR-capillary to fill all gaps) (9, 10) two representative strains: Aklavik86, from the most abundant RAPD/MLST group (17 members), and Aklavik117, a unique RAPD/MLST Amerind type, distinguished by an ftsZ-linked IS606 fragment PCR marker, which is common in Peruvian Amerind strains but rare in Aklavik (although present in Alaskan Native) strains and absent from non-Amerind \textit{H. pylori} (9) (D. Kersulyte and D. E. Berg, unpublished data). Each strain contains a typical \textit{H. pylori} chromosome size (1.5 kb or 1.6 kb) and two unrelated plasmids (<3 kb and 12.1 kb or 18.8 kb). BLASTn analyses of sequential 1-kb chromosomal segments confirmed that these two strains are of the Amerind type genome-wide.

PCR tests showed that Aklavik86 and Aklavik117 lack \textit{cag} pathogenicity islands, as do all Aklavik Amerind strains, whereas most South American Amerind strains contain these islands (9). Aklavik117 does contain two \textit{Tn} virulence-associated transposons, (11), one type I (fragmented) and one type II (intact), whereas Aklavik86 lacked \textit{Tn} PZs. Only two insertion sequence (IS) elements were present, IS607 in Aklavik117 and IS605 (deletion mutant) in Aklavik86, and no prophages were present in these two genomes.

Two Aklavik86 features merit special mention: (i) iron-dependent urease genes (previously observed only in helicobacters from carnivores [10, 12, 13]), next to \textit{cheW}, as in \textit{Helicobacter aconyctis} and \textit{Helicobacter cetorum}, and (ii) an unprecedented ~140 rearrangements in its chromosome (identified using Mauve software) relative to Aklavik117 and all other sequenced \textit{H. pylori} strains. This contrasts with ≤10 such rearrangements between most other \textit{H. pylori} strain pairs (10, 14). Follow-up PCR tests for six Aklavik86-specific rearrangements identified each in most of the Aklavik Amerind strains. In contrast, no rearrangements were found in Aklavik117 or in one of the Amerind or one European RAPD group of Aklavik strains (11 and 10 isolates, respectively), or in any of 80 other strains variously from Peru, Europe, or Asia.

Aklavik86’s extraordinary iron-dependent urease genes and many chromosomal rearrangements suggest that it might represent a novel lineage, possibly derived from a jump many millennia ago from some ancient animal host. More generally, these findings should stimulate analyses of \textit{H. pylori} population structure and evolution in human health and disease, especially in Arctic Aboriginal communities.

**Nucleotide sequence accession numbers.** The complete \textit{H. pylori} chromosome and plasmid sequences reported here have been deposited in GenBank under the accession numbers CP003476, CP003477, and CP003478 (Aklavik86) and CP003483, CP003484, and CP003485 (Aklavik117). The versions described in this paper are the first versions (CP003476.1, CP003477.1, and
ACKNOWLEDGMENTS
The \textit{H. pylori} isolates studied here were obtained from Aklavik residents who provided written informed consented for endoscopy with gastric biopsy and characterization of the \textit{H. pylori} strains obtained. These residents were participants in the Aklavik \textit{H. pylori} Project, made possible through partnerships with territorial health authorities (NWT Health and Social Services, Stanton Territorial Health Authority, and the Beaufort-Delta Regional Health and Social Services Authority), the Hamlet of Aklavik mayor and council, the Aklavik Indian Band (Ehdidiat Gwich’in Council), the Aklavik Community Corporation (local Inuvialuit governance), and the Inuvialuit Regional Corporation. This project, part of the Canadian North Helicobacter pylori (CANHelp) Working Group research program, was guided by a committee comprising the Aklavik Health Committee and Rachel Munday, Nurse-in-Charge, Susie Husky Health Centre, and approved by the University of Alberta Health Research Ethics Board and the Aurora Research Institute (NWT research licensing program, was guided by a committee comprising the Aklavik Health Committee and Rachel Munday, Nurse-in-Charge, Susie Husky Health Centre, and approved by the University of Alberta Health Research Ethics Board and the Aurora Research Institute (NWT research licensing program). The DNA sequencing reported here was carried out with the approval of the CANHelp Working Group and the Aklavik \textit{H. pylori} Project planning committee. Prior to publication, our results were shared with members of the Aklavik \textit{H. pylori} Project planning committee, who provided helpful feedback from a community perspective.

This work was supported by the Canadian Institutes of Health Research (FRN 90386), the Alberta Heritage Foundation for Medical Research, the Aklavik Community Corporation, the Inuvialuit Regional Corporation, NWT Health and Social Services, Alberta Health Services, Canadian North Airlines, and Olympic Canada, as part of the CANHelp Working Group research program, and the U.S. National Institutes of Health (grants R21 AI078237 and R21 AI088337).

We thank the staff at the Susie Husky Health Centre, members of the project planning committee, and the Aklavik community for their participation, as well as MOgene, Inc., St. Louis, MO, for quality 454 FLX Titanium DNA sequencing and contig assembly.

REFERENCES
1. Cover TL, Blaser MJ. 2009. \textit{Helicobacter pylori} in health and disease. Gastroenterology 136:1863–1873. http://dx.doi.org/10.1053/j.gastro.2009.01.073.
2. Suerbaum S, Michetti P. 2002. \textit{Helicobacter pylori} infection. N Engl J Med 347:1175–1186. http://dx.doi.org/10.1056/NEJMra020542.
3. French RW, Jr, Clemens J. 2003. \textit{Helicobacter} in the developing world. Microbes Infect 5:705–713. http://dx.doi.org/10.1016/S1286-4579(03)00112-6.
4. Goodman KJ, Jacobson K, Veldhuyzen van Zanten S. 2008. \textit{Helicobacter pylori} infection in Canadian and related Arctic aboriginal populations. Can J Gastroenterol 22:289–295.
5. Cheung J, Goodman K, Munday R, Heavner K, Huntington J, Morse J, Veldhuyzen van Zanten S, Fedorak RN, Corriveau A, Bailey RJ, CANHelp Working Group. 2008. \textit{Helicobacter pylori} infection in Canada’s Arctic: searching for the solutions. Can J Gastroenterol 22:912–916.
6. Cheung J, Goodman KJ, Girgis S, Bailey R, Morse J, Fedorak RN, Geary J, Fagan-Garcia K, Veldhuyzen van Zanten SV, CANHelp Working Group. 2014. Disease manifestations of \textit{Helicobacter pylori} infection in Arctic Canada: using epidemiology to address community concerns. BMJ Open 4:e003689. http://dx.doi.org/10.1136/bmjopen-2013-003689.
7. Carraher S, Chang HJ, Munday R, Goodman KJ, CAN Help Working Group. 2013. \textit{Helicobacter pylori} incidence and re-infection in the Aklavik \textit{H. pylori} project. Int J Circumpolar Health 72:651–657. http://dx.doi.org/10.3402/ijch.v72i0.21594.
8. Herrera PM, Mendez M, Velapatino B, Santivanez L, Balqui J, Finger SA, Sherman J, Zimic M, Cabrera L, Watanabe J, Rodriguez C, Gilman RH, Berg DE. 2008. DNA-level diversity and relatedness of \textit{Helicobacter pylori} strains in shantytown families in Peru and transmission in a developing-country setting. J Clin Microbiol 46:3912–3918. http://dx.doi.org/10.1128/JCM.01453-08.
9. Kersulyte D, Kalia A, Gilman RH, Mendez M, Herrera P, Cabrera L, Velapatino B, Balqui J, Paredes Puente de la Vega F, Rodriguez Ulloa CA, Cok J, Hooper CC, Dailide G, Tamma S, Berg DE. 2010. Helicobacter pylori from Peruvian Amerindians: traces of human migrations in strains from remote Amazon, and genome sequence of an Amerind strain. PLoS One 5:e83177. http://dx.doi.org/10.1371/journal.pone.0083177.
10. Kersulyte D, Lee W, Subramaniam D, Anant S, Herrera P, Cabrera L, Balqui J, Barabas O, Kalia A, Gilman RH, Berg DE. 2009. \textit{Helicobacter pylori} strain’s plasticity zones are novel transposable elements. PLoS One 4:e6859. http://dx.doi.org/10.1371/journal.pone.0006859.
11. Stoof J, Breijer S, Pot RG, van der Neut D, Kuipers EJ, Kusters JG, van Vliet AH. 2008. Inverse nickel-responsive regulation of two urease enzymes in the gastric pathogen \textit{Helicobacter mustelae}. Environ Microbiol 10:2586–2597. http://dx.doi.org/10.1111/j.1462-2920.2008.01681.x.
12. Carter EL, Tronrud DE, Taber SR, Karpplus PA, Hausinger RP. 2011. Iron-containing urease in a pathogenic bacterium. Proc Natl Acad Sci U S A 108:13095–13099. http://dx.doi.org/10.1073/pnas.1106915108.
13. Duncan SS, Bertoli MT, Kersulyte D, Valk PL, Tamma S, Segal I, McClain MS, Cover TL, Berg DE. 2013. Genome sequences of three \textit{hpAAfrica2} strains of \textit{Helicobacter pylori}. Genome Announc 1(5):e00729-13. http://dx.doi.org/10.1128/genomea.00729-13.