

## Complete Genome Sequence of *Chlamydia avium* PV 4360/2, Isolated from a Feral Pigeon in Italy

Anna Maria Floriano,<sup>a</sup> Sara Rigamonti,<sup>b</sup> Francesco Comandatore,<sup>c</sup> Erika Scaltriti,<sup>d</sup> David Longbottom,<sup>e</sup> Morag Livingstone,<sup>e</sup> Karine Laroucau,<sup>f</sup> Alessandra Gaffuri,<sup>g</sup> <sup>D</sup>Stefano Pongolini,<sup>d</sup> Simone Magnino,<sup>b</sup> Nadia Vicari<sup>b</sup>

<sup>a</sup>Department of Biology and Biotechnology Lazzaro Spallanzani, University of Pavia, Pavia, Italy

<sup>b</sup>National Reference Laboratory for Animal Chlamydioses, Istituto Zooprofilattico Sperimentale della Lombardia e dell'Emilia Romagna (IZSLER), Pavia, Italy

CSky Net UNIMI Platform–Pediatric Clinical Research Center Romeo ed Enrica Invernizzi, Dipartimento di Scienze Biomediche e Cliniche Luigi Sacco, Università di Milano, Milan, Italy

<sup>d</sup>Risk Analysis and Genomic Epidemiology Unit, Istituto Zooprofilattico Sperimentale della Lombardia e dell'Emilia Romagna (IZSLER), Parma, Italy <sup>e</sup>Moredun Research Institute, Pentlands Science Park, Bush Loan, Edinburgh, United Kingdom

<sup>f</sup>Bacterial Zoonoses Unit, French Agency for Food, Environmental & Occupational Health Safety (Anses), Maisons-Alfort, France <sup>g</sup>Istituto Zooprofilattico Sperimentale della Lombardia e dell'Emilia Romagna (IZSLER), Bergamo, Italy

Anna Maria Floriano and Sara Rigamonti contributed equally to this work. Author order was determined on the basis of draft writing.

**ABSTRACT** We report here the whole-genome sequence of a *Chlamydia avium* isolate recovered from a feral pigeon in 1999 in Italy. Only one complete genome of a *C. avium* strain has been published so far. Future comparative analyses could provide valuable insights on the genomic evolution of the pathogen.

The *Chlamydia* genus encompasses host-restricted bacteria characterized by a twophase developmental cycle consisting of the alternation of an extracellular form, the elementary body, and an intracellular form known as the reticulate body (1). The genus currently includes 12 validly published species (2), and among them, *C. psittaci*, *C. gallinacea*, and *C. avium* have been isolated from birds. *C. avium* was originally isolated in Germany and Italy from pigeons and parrots (3). Only two *C. avium* genome assemblies are currently available (*C. avium* strain 10DC88 [3]), and one of them is part of an unpublished comparative genomic project (GenBank accession no. ASM41773.2).

We sequenced and investigated the genome of the C. avium field isolate PV 4360/2, isolated in 1999 from a pool of intestinal and liver tissue from a pigeon found dead but with no other signs of disease in the city of Bergamo, Italy, with no evidence of lesions at necropsy (4). The isolate was stored at the Italian National Reference Laboratory for Chlamydioses and cultured on LLC-MK2 cells. DNA was extracted from purified elementary bodies (5) using a NucleoSpin tissue kit (Macherey-Nagel), and a genomic DNA library was prepared using an Illumina Nextera XT kit following the manufacturer's instructions. Wholegenome sequencing (WGS) was then performed on an Illumina MiSeg platform with a paired-end 250-bp run producing 2,036,824 paired-end reads with an average length of 235.9 bp; the read quality was checked using FastQC v. 0.11.4 (6). The Chlamydia reads were separated from host/contaminant reads using the Blobology bioinformatic pipeline (7), selecting reads mapping to contigs with a GC content of 0.3 to 0.4 and log<sub>10</sub> (coverage) of ≥2.5× (Bowtie 2 v. 2.2.6) (8). Reads were then assembled using SPAdes v. 3.10 (9). Genome finishing was then performed with PCR validation on putative joins detected with Bandage software v. 0.8.1 (10). Thus, we obtained the complete genome assembly of the isolate, comprising a circular chromosome of 1,040,639 bp and a circular plasmid of 7,640 bp. The genome and the plasmid assemblies were then annotated with RAST v. 2.0 (11). Default parameters were used unless otherwise specified. A total of 947 genes in the chromosome plus 8 in the plasmid were predicted.

Citation Floriano AM, Rigamonti S, Comandatore F, Scaltriti E, Longbottom D, Livingstone M, Laroucau K, Gaffuri A, Pongolini S, Magnino S, Vicari N. 2020. Complete genome sequence of *Chlamydia avium* PV 4360/2, isolated from a feral pigeon in Italy. Microbiol Resour Announc 9:e01509-19. https://doi.org/ 10.1128/MRA.01509-19.

Editor David A. Baltrus, University of Arizona Copyright © 2020 Floriano et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

Address correspondence to Nadia Vicari, nadia.vicari@izsler.it.

Received 12 December 2019 Accepted 25 March 2020 Published 16 April 2020



The genome of strain PV 4360/2 (size, 1,041,169 bp; GC content, 36.9%) presents a total of 3,696 single-nucleotide polymorphisms (SNPs), aligned to the only other *C. avium* complete genome available (reference strain 10DC88, GenBank assembly no. ASM58387.1) by Mauve v. snapshot version 2015-02-13 (12). No evidence of any genomic rearrangements or indels was detected. Seven Pmps (type V autotransporter proteins that play a key role in virulence and are involved in immune evasion) (13, 14) were identified. The Trp system, an important virulence factor in *Chlamydia* species, was not detected, whereas the *bioBFDA* system (15) is identical to that in the strain 10DC88.

Interestingly, alignment of the plasmid sequences identified a genomic region of 414 nucleotides (nt) present in the PV4360/2 plasmid that was absent from the 10DC88 plasmid. This region encodes two virulence genes, pGP4-D and pGP3-D. The former is truncated, and the latter is absent from the plasmid of the 10DC88 strain. This could explain the difference in length between the two plasmids. Furthermore, we found that the virulence gene pGP8-D was annotated as two separate genes in the 10DC88 plasmid.

These differences in virulence gene content show that further genomic studies could provide important insights on the evolution of the virulence of *C. avium*.

**Data availability.** The raw reads and assemblies have been deposited in ENA under the accession no. PRJEB25740. The strain is available at www.ibvr.org.

## ACKNOWLEDGMENTS

This project was funded by the Italian Ministry of Health, Directorate-General for animal health and veterinary medicinal products (DGSAF), grant no. IZS LER 2015/003 RC.

We thank Manuela Donati (University of Bologna, Italy) for collaborating in the propagation of *C. avium* in cell culture.

## REFERENCES

- Moulder JW. 1991. Interaction of chlamydiae and host cells in vitro. Microbiol Rev 55:143–190. https://doi.org/10.1128/MMBR.55.1.143-190 .1991.
- Laroucau K, Vorimore F, Aaziz R, Solmonson L, Hsia RC, Bavoil PM, Fach P, Hölzer M, Wuenschmann A, Sachse K. 2019. Chlamydia buteonis, a new Chlamydia species isolated from a red-shouldered hawk. Syst Appl Microbiol 42:125997. https://doi.org/10.1016/j.syapm.2019.06.002.
- Sachse K, Laroucau K, Riege K, Wehner S, Dilcher M, Creasy HH, Weidmann M, Myers G, Vorimore F, Vicari N, Magnino S, Liebler-Tenorio E, Ruettger A, Bavoil PM, Hufert FT, Rosselló-Móra R, Marz M. 2014. Evidence for the existence of two new members of the family *Chlamydiaceae* and proposal of *Chlamydia avium* sp. nov. and *Chlamydia gallinacea* sp. nov. Syst Appl Microbiol 37:79–88. https://doi.org/10.1016/j .syapm.2013.12.004.
- Gaffuri A, Garbarino C, Consoli T, Carrara M, Broussard D, Pietra V, Magnino S, Paterlini F. 2000. Indagine sullo stato sanitario dei Piccioni della Provincia di Bergamo. Selezione Vet 41:827–830.
- Fukushi H, Hirai K. 1988. Immunochemical diversity of the major outer membrane protein of avian and mammalian Chlamydia psittaci. J Clin Microbiol 26:675–680. https://doi.org/10.1128/JCM.26.4.675-680.1988.
- Andrews S. 2010. FastQC: a quality control tool for high throughput sequence data. http://www.bioinformatics.babraham.ac.uk/projects/ fastqc.
- Kumar S, Jones M, Koutsovoulos G, Clarke M, Blaxter M. 2013. Blobology: exploring raw genome data for contaminants, symbionts and parasites using taxon-annotated GC-coverage plots. Front Genet 4:237. https:// doi.org/10.3389/fgene.2013.00237.
- Langmead B, Salzberg SL. 2012. Fast gapped-read alignment with Bowtie 2. Nat Methods 9:357–359. https://doi.org/10.1038/nmeth.1923.

- Bankevich A, Nurk S, Antipov D, Gurevich AA, Dvorkin M, Kulikov AS, Lesin VM, Nikolenko SI, Pham S, Prjibelski AD, Pyshkin AV, Sirotkin AV, Vyahhi N, Tesler G, Alekseyev MA, Pevzner PA. 2012. SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J Comput Biol 19:455–477. https://doi.org/10.1089/cmb.2012.0021.
- Wick RR, Schultz MB, Zobel J, Holt KE. 2015. Bandage: interactive visualization of de novo genome assemblies. Bioinformatics 31:3350–3352. https://doi.org/10.1093/bioinformatics/btv383.
- Aziz RK, Bartels D, Best AA, DeJongh M, Disz T, Edwards RA, Formsma K, Gerdes S, Glass EM, Kubal M, Meyer F, Olsen GJ, Olson R, Osterman AL, Overbeek RA, McNeil LK, Paarmann D, Paczian T, Parrello B, Pusch GD, Reich C, Stevens R, Vassieva O, Vonstein V, Wilke A, Zagnitko O. 2008. The RAST server: Rapid Annotations using Subsystems Technology. BMC Genomics 9:75. https://doi.org/10.1186/1471-2164-9-75.
- Darling ACE, Mau B, Blattner FR, Perna NT. 2004. Mauve: multiple alignment of conserved genomic sequence with rearrangements. Genome Res 14:1394–1403. https://doi.org/10.1101/gr.2289704.
- 13. Vasilevsky S, Stojanov M, Greub G, Baud D. 2016. Chlamydial polymorphic membrane proteins: regulation, function and potential vaccine candidates. Virulence 7:11–22. https://doi.org/10.1080/21505594.2015 .1111509.
- Becker E, Hegemann JH. 2014. All subtypes of the Pmp adhesin family are implicated in chlamydial virulence and show species-specific function. MicrobiologyOpen 3:544–556. https://doi.org/10.1002/mbo3.186.
- Sait M, Livingstone M, Clark EM, Wheelhouse N, Spalding L, Markey B, Magnino S, Lainson FA, Myers GSA, Longbottom D. 2014. Genome sequencing and comparative analysis of three *Chlamydia pecorum* strains associated with different pathogenic outcomes. BMC Genomics 15:23. https://doi.org/10.1186/1471-2164-15-23.