Diagnosis and Genotyping of *Coxiella burnetii* Endocarditis in a Patient with Prosthetic Pulmonary Valve Replacement Using Next-Generation Sequencing of Plasma Microbial Cell-Free DNA

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Determining the causative etiology of culture-negative endocarditis can be challenging. We performed next-generation sequencing of plasma microbial cell-free DNA to facilitate rapid diagnosis and genotyping of *Coxiella burnetii* in a patient with culture-negative endocarditis of a prosthetic pulmonary valve, enabling early targeted treatment prior to valve replacement surgery.

**Keywords.** *Coxiella burnetii*; endocarditis; next-generation sequencing; plasma microbial cell-free DNA; Q fever.

Culture-negative endocarditis (CNE) comprises approximately 5%–55% of all cases of infective endocarditis and can be diagnostically challenging [1]. When severe valvular disease is present, immediate valve replacement surgery may be necessary prior to definitive microbiologic diagnosis or initiation of appropriate antimicrobial therapy. This increases the risk of re-infecting new prosthetic material. Next-generation sequencing (NGS) of microbial cell-free DNA (mcfDNA) has demonstrated clinical utility in identifying a broad range of pathogens with high sensitivity and short turnaround time [2–4]. We present a case of CNE where NGS of mcfDNA in a patient’s plasma facilitated rapid identification and genotyping of *Coxiella burnetii*, which was subsequently confirmed by serologic testing and detection of *C. burnetii* DNA in the explanted cardiac tissue.

**CASE PRESENTATION**

A 29-year-old male with 18 months of intermittent fevers, night sweats, and 6 kg weight loss presented to outpatient cardiology. He had a history of Tetralogy of Fallot with multiple cardiac surgeries, including Blalock shunt placement at age 7 days followed by repair at 3 years of age, homograft pulmonary valve replacement (PVR) in 2006, and bioprosthetic PVR in 2014. Other relevant history included travel to Pakistan, Thailand, Laos, and Myanmar after PVR in 2014, use of a LivaNova 3T Heater-Cooler device during PVR surgery in 2014, and consumption of unpasteurized milk in the Midwestern United States. He denied animal contact.

Vital signs were normal. Physical examination was notable for a holosystolic murmur and hepatosplenomegaly. Initial blood cultures and a fourth-generation HIV screening test were negative. Transesophageal echocardiography demonstrated severely elevated pulmonary artery (PA) pressure and an erratically moving echodensity on the pulmonary valve suggestive of vegetation. The patient was subsequently admitted to the hospital for further management of presumed CNE.

Initial concerns included *Mycobacterium chimaera* prosthetic valve endocarditis given a documented outbreak associated with contaminated LivaNova 3T Heater-Cooler devices [5]. Furthermore, the patient brought a letter from the hospital where the PVR was performed in 2014 warning of possible exposure to *M. chimaera*. Other potential etiologies included *Bartonella henselae*, *Brucella* species, and *C. burnetii*.

Although the patient was clinically stable, his cardiologist had concern for decompensation and sudden cardiac death due to severely elevated PA pressure. To prevent reinfection of new prosthetic material, the cardiologist consulted the infectious diseases service for empiric treatment recommendations before surgery. Upon our recommendation, serologic tests for *B. henselae*, *Brucella* species, *C. burnetii*, and *Legionella pneumophila*, as well as acid-fast bacilli blood cultures were sent. However, empiric therapy for the multiple etiologic agents under consideration, including *M. chimaera*, was not recommended given that it could reduce diagnostic yield from cardiac tissue culture and commit the patient to long-term empiric therapy with toxic antimicrobials (eg, amikacin). Based on advice from a physician with *M. chimaera* expertise, we sent the patient’s plasma for NGS of mcfDNA to facilitate rapid and comprehensive diagnosis including evaluation for *M. chimaera* infection (Figure 1A) [6].

**METHODS**

Plasma was collected from the patient at NewYork-Presbyterian Hospital–Weill Cornell Medical Center and sent to a Clinical Laboratory Improvement Amendments–certified and College of American Pathologists–accredited laboratory for NGS (Karius,
Inc., Redwood City, CA). Plasma mcfDNA was extracted, and NGS was performed according to previously described methods [2]. Human sequence reads were removed, and the remaining sequences were aligned to a curated database of >1000 pathogens including bacteria, mycobacteria, DNA viruses, fungi, and protozoa. Organisms above a predefined significance threshold were reported [2]. For C. burnetii strain typing, alignments to different Coxiella strains in the pathogen database were compared by BLAST bit score to determine the most closely related strain to the offending organism. Coxiella burnetii serology was performed by ARUP Laboratories (Salt Lake City, UT); IgG Phase I and Phase II titers were obtained by serial dilution.

**RESULTS**

Within 48 hours of sample receipt, NGS detected C. burnetii mcfDNA. Strain typing using BLAST alignments of C. burnetii
mcfDNA fragments identified a best match against sequence type (ST) 8 (including strains CbuK_Q154 and MSU Goat Q177) (Supplementary Figure 1), which belongs to genomic Group IV (Figure 1B) as determined by multiplexar sequence typing (MST) [7].

Based on the mcfDNA sequencing results, the patient was diagnosed with presumed C. burnetii endocarditis. Upon further discussion, the patient recalled witnessing the slaughtering of a goat at close distance in rural Pakistan, a more likely risk factor than the consumption of unpasteurized milk in the United States. Three days later, serologic testing revealed a C. burnetii Phase I IgG titer of 1:1 048 576 and a Phase II IgG titer of 1:2 097 152 (Figure 1A). After treatment with hydroxychloroquine (200 mg orally every 8 hours) and doxycycline (100 mg orally twice daily), the patient was able to completely resolve symptoms after 4 weeks of therapy and ultimately underwent successful PVR and graft exchange.

Hematoxylin and eosin staining of the explanted cardiac tissue demonstrated fibrosis and a chronic inflammatory infiltrate composed of lymphocytes, plasma cells, and macrophages, including scattered giant cells. Cardiac tissue was also sent to the Centers for Disease Control and Prevention (CDC) for further investigation. Coxiella burnetii DNA was successfully extracted from formalin-fixed, paraffin-embedded tissue, and strain typing was performed. Genotyping at the CDC using single nucleotide polymorphism (SNP) markers compatible with MST also placed the C. burnetii strain in genomic Group IV. SNP typing revealed that the C. burnetii strain was closely related to a group of STs including 9, 10, 27, 28, and 31 [8] but was unable to distinguish between these sequence types (Figure 1B).

Six months after the initial diagnosis, serologic titers were repeated to monitor response to antimicrobial and surgical therapy, demonstrating an 8-fold decrease in Phase I and 32-fold decrease in Phase II titers (1:131 072 and 1:65 536, respectively). At the time of this publication, the patient is doing well without recurrence of symptoms and will remain on hydroxychloroquine and doxycycline for at least 24 months [10].

DISCUSSION

Coxiella burnetii, the etiologic agent of Q (Query) fever, is transmitted via zoonosis and is an environmentally ubiquitous, pleomorphic gram-negative coccobacillus that can cause acute systemic illness or chronic infections such as endocarditis [10, 11]. Coxiella burnetii does not grow in culture using routine laboratory culture techniques, and thus diagnosis of C. burnetii endocarditis typically requires compatible clinical symptoms and serology [1, 10].

We present a case of CNE due to C. burnetii that was initially diagnosed by NGS of plasma mcfDNA and subsequently confirmed by serology and polymerase chain reaction amplification of C. burnetii DNA from cardiac tissue. Sequencing of mcfDNA facilitated noninvasive, rapid diagnosis of C. burnetii endocarditis within 48 hours of sample collection, decreasing the likelihood of M. chimaera infection. The significance of the NGS result was further supported by high Phase I and Phase II IgG titers and the patient's compatible exposure history.

There are several advantages of sequencing plasma mcfDNA. First, it can detect bacterial, fungal, parasitic, and DNA viral pathogens in a single assay, reducing the need for numerous pathogen-specific assays. The turnaround time is rapid (24–48 hours upon receipt of the specimen at the testing laboratory), facilitating early initiation of targeted therapy. The specimen collection is minimally invasive (venipuncture), which may decrease the requirement for invasive procedures such as tissue biopsy. Therefore, plasma NGS can permit earlier diagnosis than nucleic acid testing or culture of surgically obtained cardiac tissue [6]. In addition, strain-level information may be obtained by leveraging the plasma mcfDNA sequence data, which would not be available with serology alone. In this case, genotyping elucidated the nature and location of C. burnetii acquisition, which are important for epidemiologic investigations given the organism's classification as a Category B bioterrorism agent [12]. Strain typing of mcfDNA isolated from our patient was most consistent with ST8 among all publicly available whole-genome sequences. SNP typing of DNA isolated from cardiac tissue at the CDC placed the sample in the closely related ST groups 9, 10, 27, 28, and 31 based on an internal database. Reconciling strain typing information (Figure 1B) and exposure history, the patient likely acquired Coxiella infection during international travel, possibly while observing a goat slaughter in rural Pakistan, rather than through consumption of unpasteurized cow's milk in the United States.

There are several shortcomings of sequencing plasma mcfDNA: they are generally more expensive than culture-based methods, serology, or nucleic acid amplification tests, and at present do not offer some advantages of culture such as phenotypic antimicrobial susceptibility testing. Finally, it is important to recognize that detection of a wide range of organisms may lead to the identification of atypical organisms of unclear clinical significance, and the false-positive rate of NGS remains uncertain. In such cases, interpretation should be cautious, and involvement of infectious diseases and clinical microbiology specialists is essential.

CONCLUSIONS

To the best of our knowledge, this is the first report describing the use of NGS of mcfDNA obtained from plasma to diagnose C. burnetii CNE. Plasma NGS aided in rapid diagnosis, facilitated exclusion of alternative infectious agents in this complicated case, and allowed inference of strain-level information,
supporting further investigations regarding the patient’s acquisition of *C. burnetii*. Future clinical studies are needed to validate mcfDNA sequencing as a useful, robust, and informative diagnostic test in CNE [2].

**Supplementary Data**

Supplementary materials are available at *Open Forum Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

**Notes**

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**Author contributions.** M.K., K.M.M., and N.E. cared for the case patient and provided all clinical data. M.K. (with S.D.) wrote the manuscript. S.V. performed NGS analysis, performed genotyping of mcfDNA, and created the figures (with M.K.). B.D.R. contributed pathology slides. L.F.W. and K.M.M. coordinated the study and supervised the overall process. All authors reviewed and revised the manuscript.

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