**Janibacter species with evidence of genomic polymorphism isolated from resected heart valve in a patient with aortic stenosis**

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**Abstract**

The authors report isolation and identification of two strains of bacteria belonging to the genus *Janibacter* from a human patient with aortic stenosis from a rural area of the country of Georgia. The microorganisms were isolated from aortic heart valve. Two isolates with slightly distinct colony morphologies were harvested after sub-culturing from an original agar plate. Preliminary identification of the isolates is based on amplification and sequencing of a fragment of 16S rRNA. Whole genome sequencing was performed using the Illumina MiSeq instrument. Both isolates were identified as undistinguished strains of the genus *Janibacter*. Characterization of whole genome sequences of each culture has revealed a 15% difference in gene profile between the cultures and confirmed that both strains belong to the genus *Janibacter* with the closest match to *J. terrae*. Genomic comparison of cultures of *Janibacter* obtained from human cases and from environmental sources presents a promising direction for evaluating a role of these bacteria as human pathogens.

**Materials and Methods**

**Bacterial culture**

Surgically removed cardiac valve material and whole blood were collected from a patient with suspected endocarditis. The clinical part of the work was approved by the Ethical Committee of the High Technology Medical Center, Tbilisi, Georgia. The material was frozen, transferred to the National Center for Diseases Control and Public Health, Tbilisi, Georgia, and stored at -80°C until it was plated. The homogenized cardiac tissue and whole blood were used to inoculate chocolate agar plates. The inoculated plates were incubated at 35°C with 5% carbon dioxide for 30 days. Each individual colony from agar was separately streaked onto secondary agar plates. The subcultures were harvested separately by scraping bacterial growth after sufficient growth was observed. DNA was heat extracted at 95°C for 10 min from whole bacterial cells. The prepared DNA was stored at -80°C.

**PCR amplification and sequencing**

Amplification of the 16S rRNA fragments from isolated DNA of both cultures
was performed by conventional broad-range PCR assay using 63F and 1187R primers in a C1000 Touch Thermal Cycler (Bio-Rad, Hercules, CA). The PCR products were analyzed for the presence of amplicons of the appropriate size by electrophoresis in 1.5% agar gels containing GelGreen stain (Biotium, Hayward, CA). Positive and negative controls were included in each PCR assay. PCR products with appropriate sizes of amplicons were purified using a QIAquick PCR Purification Kit (Qiagen, Valencia, CA) according to manufacturer’s instructions, then sequenced in both directions using ABI 3130 Genetic Analyzer (Applied Biosystems, Foster City, CA).

**Genome analysis**

Whole genome sequencing (WGS) was performed using the Illumina MiSeq instrument. Sequencing of the two samples resulted in 8 million 51 nucleotide long sequencing reads. The sequencing reads were filtered to exclude: (a) nucleotides below the quality threshold of 0.05 (using modified Richard Mott algorithm); (b) unknown nucleotides; and (c) sequencing library adapters. The reads were then assembled allowing for up to 5% mismatches using CLC Genomics Workbench De Novo Assembly tool (http://www.clcbio.com) and the original sequenced reads were mapped back to the assembled contigs (Table 1). Rapid Annotation using Subsystem Technology tool kit (RASTk) was used to cut open reading frames from the assembled contigs on the PATRIC web interface. Annotated features were clustered together in protein space using CD-HIT with the following parameters: 70% global identity and 70% alignment coverage. A Venn diagram of the feature clustering between the two sequenced isolates and the closest reference, *Janibacter terrae NBRC 107854 = JCM 12887*. Blastn was used to compare the features annotated with the closest reference, *Janibacter terrae NBRC 107854 = JCM 12887* (GCA_001598955.1). The results were visualized using Blast Ring Image Generator (Figure 2).

### Results

A 60-year-old previously healthy female experienced symptoms of headache, dizziness and syncope episodes, general weakness, exertional fatigue, and shortness of breath for 2 years. She also reported low-grade episodic fevers in the preceding 6 months. One month prior to hospital admission, the patient’s fevers became more persistent and she was evaluated in an outpatient clinic. She was prescribed empiric antibiotics of unknown type, but did not improve. She was admitted to the intensive care unit in May 2014 due to periodic squeezing pain in the chest that radiated under the left shoulder blade with the original diagnosis of endocarditis. The patient lived in a rural area near the town of Karelí in the region of Shida Kartlì, Georgia (South Caucasus). She was engaged in agricultural work and reported frequent contacts with farm animals (pigs and cattle) and pets at home.

Clinical signs upon hospital admission included fever of 38°C, ischemic disease, angina, and arterial hypertension. No rash or lymphadenopathy was reported. Coronary angiography showed unaltered coronary vessels. Echocardiography detected critical stenosis of the aortic valve with high gradient and critical opening. On auscultation there was rough systolic murmur at every auditory point, with epicenter on the aorta. Heart rate – 76 beats/min, pulse – arrhythmic, average filling and tension. T/A – 130/60 mmHg. The patient’s WBC was 5.60 10^3/uL (normal range: 3.4 – 9.6 10^3/uL), neutrophil count 42% (normal range: 40% – 60%), and lymphocyte count 48% (normal range: 20% – 40%), platelets – 193 10^3/uL(normal range: 157 – 371 10^3/uL for females); and ESR – 10 mm/h (normal range: 1 – 20 mm/h for females). Chest X-ray and abdominal ultrasound did not show any abnormalities. The case was diagnosed as aortic stenosis, and the patient underwent heart valve replacement. As standard post-surgical prophylaxis, the patient was treated with Ceftriaxone 1 gram two times per day after the surgery.

No bacterial growth was observed on agar plates inoculated with blood of the patient. On the seventh day after inoculation of agar plates with homogenized cardiac tissues two small bacterial colonies were observed. Two slow-growing colonies with slightly different morphology were observed on the agar plate inoculated with heart tissues. Both colonies had circular shape and light cream color with a slight difference in size. No hemolysis on chocolate agar was observed. These two colonies were separately sub-cultured onto new agar plates. Microscopy indicated Gram-positive cocccobacilli. When bacterial mass on both plates was sufficient, colonies were harvested on brain heart infusion medium and stored at –70°C until the strains were shipped to CDC-Fort Collins for additional characterization.

By Mega BLAST search, the ribosomal amplicons from both isolates were identified as a *Janibacter* sp. based on the fragment amplification and sequence analysis with the highest match to *Janibacter hoylei*. The 16S rRNA sequences from both isolates were undistinguished.

Sequencing of each isolate genome resulted in 2 and 6 million sequencing reads 51 nucleotides long in size trimmed to 32 bases long non-overlapping subsequences with a minimum threshold quality of 10. To determine the presence and identity of microbial contaminants, sequencing data was tested against a collection of 13,774 complete bacterial, viral, plasmid, phage, and fungal reference genome sequences, which included all reference genomes available from the NCBI Reference Sequence (collection as of November 2013).

*Intrasporangiaceae* family had the highest number of identified genes from the database. A more detailed mapping to every genome entry in NCBI under the *Intrasporangiaceae* family was performed. For references, which did not contain a fully assembled genome, the contigs were concatenated into one long sequence that was treated as a genome. Lowest Common Ancestry analysis of full length 16S sequences assembled from whole genome sequencing reads showed a 99.9% match to the genus *Janibacter*. The closest match by genome analysis was to *Janibacter ter-****

### Table 1. Assembly statistics genomes of two strains of *Janibacter* (TB1 and TB2)

| Feature               | Strain TB1 | Strain TB2 |
|-----------------------|------------|------------|
| N75                   | 20,309     | 22,383     |
| N50                   | 35,574     | 48,911     |
| N25                   | 56,767     | 84,095     |
| Minimum Contig Length | 205        | 210        |
| Maximum Contig Length | 118,653    | 165,607    |
| Average Contig Length | 15,154     | 22,752     |
| Number of Contigs     | 219        | 142        |
| Total Bases Assembled | 3,318,728  | 3,230,833  |
Discussion

Two discoveries can be highlighted from the reported results: 1) both bacterial cultures isolated from a human patient with aortic stenosis were identified as a species of genus *Janibacter* that has never been previously reported as a source of a cardiac disease in humans or animals; and 2) complete genome comparison indicated genomic difference between bacterial cultures obtained from the same cardiac tissue.

The report of the isolation of two closely related bacterial strains belonging to the genus of *Janibacter* from the case of aortic stenosis is novel not only for this region, but presents the first known case worldwide. The bacteria of genus *Janibacter* were originally found in the environment,15 but since 2005 at least nine cases of human infections associated with these microorganisms have been reported. Among these cases, one case was associated with *J. melonis*,16 one case with *J. hoyeli*,17 two cases with an undescribed species of *Janibacter*,18,19 and five cases with *J. terrae*.20,21 The cases associated with *J. terrae* are especially interesting since the isolates that we report here from the cardiac tissues were closely related by the genomic analysis to this species of *Janibacter*. Fernandez-Natal et al.20 reported cases of infection by *J. terrae* in four febrile patients with severe underlying conditions in Spain. All four patients were treated with antibiotics, two of them with favorable outcome, and two severely immunocompromised patients died. More recently, Wan et al.21 reported a case of bilateral psoas abscess caused by *J. terrae*.

Although this is the first report of bacteria of the genus *Janibacter* isolated from a cardiac valve, these microorganisms were broadly described as “coryneform” bacteria with some representatives being reported as sources of endocarditis.22 Coryneform bacteria, defined as non-sporing Gram-positive rods with an irregular outline, were historically dismissed as a part of the normal human skin flora when recovered from human patients, but have increasingly been implemented as causes of significant infections in clinical practice.23

Identification of a *Janibacter* strain in cardiac tissue can lead to evaluation of this bacteria as a potential source of infection; however, we wish to keep a conservative approach for a validation of this claim. Cardiac valve tissue is considered sterile; nevertheless, though a contamination of cardiac tissue during the surgery or culturing procedure is very unlikely, we cannot rule out such a possibility. More importantly, we cannot prove a causal connection between this bacterium and pathogenic manifestations in the patient’s heart because of very limited information available about these bacteria. Therefore, an etiological role of this bacterium as a source of aortic stenosis requires further investigations. Nevertheless, the presented findings suggest that clinicians should be aware of a possible risk of infection by bacteria of the genus *Janibacter* with the ability to cause invasive cardiac disease.

Having limited data on potential risk factors, it is hard to speculate about routes of transmission of the bacteria to the patient. The patient’s report of frequent exposure to farm animals, however, may suggest a possible zoonotic source of the infection. In Germany, *Janibacter* DNA was identified from a biofilter treating air emissions from a livestock facility.24

The study has demonstrated additional perspectives based on the ability of the computational analysis of the whole bacterial genomes to identify and characterize fastidious microorganisms discovered in clinical specimens. Although 16SrRNA analysis
identified Janibacter species in the culture colonies, the whole genome sequencing allowed to more precisely differentiate these strains from other Janibacter species and to evaluate differences between colonies. The latter finding highlighted the fact that the bacterial population occupying cardiac tissues from one patient was polymorphic. Genomic comparison of cultures of Janibacter obtained from human cases and from environmental sources presents a promising direction for evaluating a role of these bacteria as human pathogens.

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