In clinical practice, unidentified *Mycobacterium* species are sometimes detected in respiratory specimens. We identified a novel *Tsukamurella* species (*Tsukamurella* sp. TY48, RIMD 2001001, CIP 111916<sup>T</sup>), *Tsukamurella toyonakaense*, from a patient given a misdiagnosis of nontuberculous mycobacterial pulmonary disease caused by unidentified mycobacteria. Genomic identification of this *Tsukamurella* species helped clarify its clinical characteristics and epidemiology.

In clinical practice, unidentified *Mycobacterium* species are sometimes detected in respiratory specimens. Few *Mycobacterium* species can be identified by using methods available in clinical practice, although there are ≈200 species of nontuberculous mycobacteria (NTM) (1). We reported a case of pulmonary disease caused by a novel *Tsukamurella* species identified by using multilocus sequence typing (MLST) and whole-genome sequencing (WGS) (2).

The Study


WGS analysis of preserved unidentified mycobacteria culture isolates was approved by the institutional research ethics board (TNH2019063–2). The requirement for informed consent was waived because of the retrospective nature of the analysis. The opt-out recruitment method was applied to provide an opportunity for all patients to decline participation. Results of WGS analysis of TY48 were deposited in BioProject (accession no. PRJDB10620) and BioSample (accession no. SAMD00250050).

We performed MLST and WGS of culture isolates from 8 patients given diagnoses of NTM pulmonary disease caused by unidentified mycobacteria. We identified *Mycobacterium shimoidei*, *M. shinjukuense*, *M. paragordonae*, *M. heckeshornense*, *M. lentiflavum* (3 isolates), and a novel *Tsukamurella* species (*Tsukamurella* sp. TY48, RIMD 2001001, CIP 111916<sup>T</sup>).

The patient with *Tsukamurella* infection was an 82-year-old woman who had received a diagnosis of NTM pulmonary disease 23 years earlier. Then a 59-year-old previously healthy woman, she was referred to our hospital because of abnormal chest radiographic findings. Although she had no symptoms, chest computed tomography findings showed centrilobular nodules and bronchiectasis. During follow-up, a cough and occasional hemoptysis developed. *M. chelonae* was repeatedly identified from her sputum. We started airway clearance therapy with erythromycin and expectorants. After 2 years of treatment, the *M. chelonae* disappeared from her sputum. However, her symptoms and radiologic findings slowly but steadily progressed (Figure 1), and rapidly growing acid-fast bacilli were repeatedly detected in her sputum for 8 years. The culture isolates were Ziehl-Neelsen stain positive. However, the species/subspecies could not be identified by using conventional methods. Therefore, she was given a diagnosis of NTM pulmonary disease caused by unidentified mycobacteria.

Unidentified *Mycobacterium* species are sometimes detected in respiratory specimens. We identified a novel *Tsukamurella* species (*Tsukamurella* sp. TY48, RIMD 2001001, CIP 111916<sup>T</sup>), *Tsukamurella toyonakaense*, from a patient given a misdiagnosis of nontuberculous mycobacterial pulmonary disease caused by unidentified mycobacteria. Genomic identification of this *Tsukamurella* species helped clarify its clinical characteristics and epidemiology.
We continued erythromycin treatment for >20 years on the basis of evidence regarding successful treatment of NTM pulmonary disease with erythromycin (3). However, her symptoms and radiologic findings of lung destruction and structural alterations slowly but steadily progressed.

Because of this progression, we performed WGS by using a MinION Sequencer and Flow Cell R94 (Oxford Nanopore Technologies, https://nanoporetech.com). We extracted genomic DNA from cultured isolates by using a NucleoSpin Microbial DNA Kit (Takara Bio, https://www.takarabio.com) and prepared a library by using the Rapid Barcoding Kit (Oxford Nanopore Technologies). Using MinION raw sequencing reads, we performed MLST analysis on the 184-gene accessory genome with mlstverse.

Figure 1. Comparison of chest computed tomography findings over time for patient who had chronic pulmonary disease caused by Tsukamurella toyonakaense. Findings are shown from before Tsukamurella species was detected (A, C, E, and G) and 6 years later (B, D, F, and H). A and B show that nodules in right segment 2 and left segment 6 were unchanged. C and D show that bronchiectasis in lingula had progressed. E and F show that bronchiectasis newly appeared in the middle lobe. G and H show that nodules newly appeared in left segments 8–10.
Pulmonary Disease Caused by *T. toyonakaense*

The unidentified mycobacterium was presumed to be *M. fallax* (MLST score 0.083). However, the low MLST score prompted a deeper analysis of the bacterial genome.

We conducted a 16S rRNA analysis by performing a homology search using blastn (https://blast.ncbi.nlm.nih.gov) and compared our data with that in the SILVA rRNA database (4). The phylogenetic tree constructed using full-length 16S rRNA genes showed that strain TY48 was closely related to other *Tsukamurella* species (>98.7%), whereas its homology to 2 type species belonging to the related bacteria *Gordonia bronchialis* and *Williamsia muralis* was only 94.0% (Figure 2). We next determined the complete genome sequence of TY48 as reported (1) and performed WGS by using MinION and HiSeq 2500 instruments (Illumina, https://www.illumina.com). We performed genome assembly for strain TY48 by using flye (https://www.flye.com) for long reads obtained from MinION and corrected sequencing error by using pilon (https://bio.tools/pilon).

A comparison of the TY48 genome sequence with those of other *Tsukamurella* species indicated that the nearest related species was *T. paurometabola* (average nucleotide identity of 86.2%) (Table 1). This finding suggested that *Tsukamurella* sp. TY48 (RMD 2001001; CIP 111916) was a novel *Tsukamurella* species.

We performed antimicrobial drug susceptibility tests for rapidly growing mycobacteria by using the broth microdilution method in accordance with Clinical and Laboratory Standards Institute M24-A2 guidelines (5). We transferred the culture to Middlebrook 7H9 broth and vortexed. We adjusted the culture medium to a 0.5 McFarland standard with sterile distilled water; we then added 60 µL of the 0.5 McFarland suspension to a Cation-Adjusted Mueller-Hinton Broth (Kyokuto Pharmaceutical Industrial Co. Ltd., https://www.kyokutoseiyaku.co.jp) and dispensed 100 µL of this solution into each well of the panel. After confirming adequate growth of the control over a 3-day incubation in a standard atmosphere at 30°C, we determined the MICs (µg/mL) for 15 drugs: clarithromycin, 0.25; azithromycin, ≤0.25; cefoxitin, ≤8; imipenem, ≤0.5; meropenem, ≤0.5; faropenem, ≤1; amikacin, ≤1; tobramycin, 2; minocycline, ≤0.25; doxycycline, ≤1; linezolid, ≤4; moxifloxacin, ≤0.25; ciprofloxacin ≤0.5; levofloxacin, ≤0.5; and trimethoprim/sulfamethoxazole, ≤2/38. *Tsukamurella* sp. TY48 was sensitive to all 15 drugs.

Figure 2. Maximum-likelihood phylogenetic tree constructed by using 16S rRNA sequences of *Tsukamurella* spp. and other bacterial species. Bold indicates strain isolated in this study. Reference sequences were obtained from SILVA database (4) release 138 as small subunit reference nonredundant 99 sequences, which showed >98.7% identity with strain TY48. GenBank accession numbers are provided for reference sequences. Scale bar indicates nucleotide substitutions per site.
We renamed TY48 as *T. toyonakaense* after the location of its discovery, Toyonaka, Japan. *T. toyonakaense* is an aerobic, nonmotile, gram-positive rod that grows at 30°C and 37°C, but not at 42°C, and produces catalase. After a 72-h incubation at 30°C on 7H11 agar, it forms white and creamy, rough, non-pigmented colonies (10 mm in diameter). According to the API 50 CH system (bioMérieux, https://www.biomerieux.com), this bacterium can assimilate fructose, glucose, starch, sucrose, and trehalose but not arabinose, mannitol, mannose, or xylose.

After diagnosis, we attempted combination drug therapy with clarithromycin (200 mg/d) and ethambutol (250 mg/d). The patient refused continuation of treatment after 2 weeks because of antimicrobial drug-induced fatigue. We then resumed treatment with erythromycin. Her symptoms and radiologic findings are slowly improving (Table 2, https://wwwnc.cdc.gov/EID/article/28/7/21-2320-T2.htm; Appendix, https://wwwnc.cdc.gov/EID/article/28/7/21-2320-App1.pdf).

**Conclusions**

*T. toyonakaense* commonly causes acute onset pneumonia with cavity and consolidation (Table 2) and fever, coughing, sputum, fatigue, and hemoptysis. Although appropriate drugs and treatment durations are unknown, combination medications of ≥2 drugs, including rifampin or quinolone, are widely used and presumed effective on the basis of case reports (6,8,12–14). These reports also indicated a good prognosis for *Tsukamurella* pulmonary disease (8,12,13). No relapses were reported, in contrast to NTM pulmonary disease. Although the Clinical and Laboratory Standards Institute has proposed breakpoints for aerobic actinomycetes (5), no definitive drug breakpoints for *Tsukamurella* spp. have been established. However, the strain we identified showed extensive antimicrobial drug susceptibility.

Because a clinically applicable identification technique is not available, *Tsukamurella* infections are probably underestimated and more prevalent than has been recognized. Misidentification as related genera, especially *Mycobacterium*, results in missed opportunities to properly treat *Tsukamurella* infections. Use of genomic sequencing to identify *Tsukamurella* species and more cases of *Tsukamurella* infections will help identify clinical characteristics and clarify epidemiology of *Tsukamurella* pulmonary disease.

**Acknowledgments**

We thank the patient for granting permission to publish her information and Traci Raley and Katie Oakley for editing drafts of this manuscript.

This study was supported in part by the Japan Agency for Medical Research and Development (grants JP20fk0108129, JP21fk0108129h0702, and JP21lm0203007), GlaxoSmithKline (grant A-32), the Japan Society for the Promotion of Science Grants-in-Aid for Scientific Research (grants JP21K16118 and JP21K08194), the Smoking Research Foundation, the Takeda Science Foundation, and the Japan Intractable Diseases (Nanbyo) Research Foundation (grant 2020B02).
K.F. designed the study; K.H., H.S., T.N., and A.K. performed mycobacterial culture and analysis of culture isolates; K.F. and T.K. performed clinical and laboratory data extraction and analysis; Y.A., T.S., E.A., Takahiro Kawasaki, T.M., K.T., M.M., K.M., and H.K. assisted with data extraction and analysis; Y.M., D.M., and S.N. performed multilocus typing analysis and whole-genome analysis; M.T., Y.H., and T.I. analyzed biochemical profiles of the strain; K.F. and S.K. wrote the manuscript; and A.K. and H.K. supervised the study. All authors read and approved the manuscript.

About the Author
Dr. Kuge is a clinical fellow in respiratory medicine in the respiratory medicine department of Osaka University Medical Hospital, Osaka, Japan. His primary research interest is respiratory infections.

References

Address for correspondence: Kiyoharu Fukushima, Department of Respiratory Medicine and Clinical Immunology, Osaka University Graduate School of Medicine, 2-2 Yamadaoka, Suita, Osaka, Japan; e-mail: fukushima@imed3.med.osaka-u.ac.jp
Chronic Pulmonary Disease Caused by 
*Tsukamurella toyonakaense*

Appendix

References

   https://doi.org/10.1128/jcm.16.4.604-607.1982

   https://doi.org/10.1016/S0163-4453(03)00139-7


   https://doi.org/10.7326/0003-4819-150-2-200901200-00022

   https://doi.org/10.1186/1476-0711-8-30

   https://doi.org/10.1016/j.rmedc.2009.02.004

   https://doi.org/10.1097/MAJ.0b013e3182129d02

https://doi.org/10.1080/00325481.2017.1344513