LMG_4337\textsuperscript{f} (92.3\% identity). On the basis of Clinical Laboratory Standards Institute guidelines (2), this similarity indicated a possible novel species.

The DPHL Microbiology Department’s laboratory manager sent the organism to CDC’s Special Bacteriology Reference Laboratory. Results identified the organism as \textit{H. jordaniae}. Upon review of MicrobeNet at a later date, the organism was found to match \textit{H. jordaniae} H5569\_ con\textsuperscript{f} by 98.9\%.

\textit{H. jordaniae} is a common environmental microbe, but it was implicated in this clinical case in a man in Delaware. The patient had symptoms characteristic of other pathogenic bacterial illnesses. Concern exists that slow-growing, gram-negative rods identified in blood culture could be potential bioterrorism agents. Humrighouse et al. (1) described how \textit{Francisella tularensis} infection was suspected in 2 clinical cases that were actually \textit{H. jordaniae} infections.

Humrighouse et al. (1) proposed the name \textit{H. jordaniae} on the basis of an isolate received in 2010. Previously, 14 organisms identified at CDC were isolated from blood taken from men 39–78 years of age with symptoms including swelling of the lower extremities (2 patients), septicemia (3 patients), and bacteremia (1 patient). The symptoms of the patient we report mirrored those symptoms.

This discovery is important because it demonstrates that organisms conceived to be environmental in nature and suspected to have limited clinical implications are emerging as human pathogens. The ability to identify bacteria by sequencing (in this case, sequencing of the 16S rRNA gene) was necessary to identify \textit{H. jordaniae} because clinical information on this pathogen is still limited.

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### Molecular Typing and Antifungal Susceptibility of \textit{Candida viswanathii}, India

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We report invasive candidiasis caused by \textit{Candida viswanathii} over 2 time periods during 2013–2015 in a tertiary care hospital in Chandigarh, India. Molecular typing revealed multiple clusters of the isolates. We detected high MICs for fluconazole in the second time period.

Invasive candidiasis is a life-threatening infection caused by various \textit{Candida} species. Although \textit{C. albicans} has been the predominant species causing invasive candidiasis, non-albicans \textit{Candida} (NAC) species have emerged globally (1). \textit{C. viswanathii}, a pathogen first isolated from the cerebrospinal fluid of a patient in 1959 (2), is rarely encountered, and only 17 cases have been reported worldwide (3). This agent has been isolated sporadically from animal and environmental sources (4–6).

We report on 23 cases of invasive candidiasis caused by \textit{C. viswanathii} at a tertiary care center in Chandigarh, India, involving 7 case-patients during December 2013–April 2014.
and 16 case-patients during December 2014–April 2015. In the
first time period, all isolates were from blood, whereas
in the second time period, the agent was isolated from pus
(n = 5), blood (n = 5), cerebrospinal fluid (n = 3), and lung
nodule, lung aspirate, and ieliac fluid (n = 1 each).

Of the 23 patients, 16 were men and 7 were women. Six (26%)
patients had neutropenia, and 18 (90%) had tuberculous,
pancreatitis, or chronic kidney disease. Eight (34.7%)
patients acquired the infection after surgery. Twelve patients
used indwelling devices: 3 (15%) had a central venous catheter,
4 (20%) an endotracheal tube, 3 (15%) a drainage catheter,
and 2 (10%) a urinary catheter.

We screened the hospital environment and the hands of
healthcare workers for a possible source of C. viswanathii
infection during the second time period. We could not iso-
late C. viswanathii from any of those samples from a total
of 46 workers and 57 different environmental sites.

Conventional methods failed to differentiate C. viswa-
натii and C. tropicalis. C. viswanathii assimilated sucrose
and cellobiose but failed to assimilate trehalose and raffi-
nose. C. tropicalis has a variable assimilation pattern for
these sugars.

To identify the isolates, we performed matrix-assisted
laser desorption/ionization time-of-flight mass spectros-
copy using MALDI-TOF MS, version 3 (Bruker Daltonik
GmbH, Bremen, Germany) and sequenced the D1/D2 re-
gion of a large subunit of ribosomal DNA. We
identiﬁed the isolates with a log score of >1.8
by using the modiﬁed database. The rDNA sequence of
the isolates showed 100% similarity with the type strain
of C. viswanathii ATCC 22981 (GenBank accession no.
NG_054835.1) except for 1 isolate (99% similarity with
type strain, accession no. MF682371). The molecular phy-
logenetic analysis revealed that 1 isolate (B-30815) had
1 nucleotide substitution (T to C), which was 1 of the 5
substitutions we observed in C. pseudoviswanathii while
comparing it with C. viswanathii.

Amplified fragment-length polymorphism revealed a
similarity coefﬁcient of ≥90% of the isolates (online
Technical Appendix Figure, https://wwwnc.cdc.gov/EID/
article/24/10/18-0801-Techapp1.pdf). The isolates from the
first time period formed 2 clusters (clusters A and B); 1 isolate from the second period was also in cluster B.
Isolates of the second time period had 3 major clusters
(clusters C, D, and E) and had higher MICs for ﬂuconazole.

We performed antifungal susceptibility testing for
amphotericin B, ﬂuconazole, itraconazole, voriconazole,
posaconazole, caspofungin, anidulafungin, and micafun-

gin by the microbroth dilution method recommended by
the Clinical and Laboratory Standards Institute. After
incubating the plates for 24 h at 37°C, we took a visual
reading to determine the MICs. The isolates of the second
time period had higher MICs (MICs 64 μg/mL, MICs 0.25 μg/mL
for ﬂuconazole compared with the isolates of the
first period (MICs 1 μg/mL, MICs 0.125 μg/mL). We also
recorded higher MICs (MICs 1 μg/mL, MICs 2 μg/mL) for
voriconazole for the isolates of the second period (Table).

In conclusion, our study showed multiple clusters of
C. viswanathii causing invasive infections in patients with
neutropenia and chronic diseases at a single healthcare cen-
ter in India. We could not trace the source of the agent.
Conventional identiﬁcation methods could not differentiate
the isolates from those of C. tropicalis. The high MICs for
voriconazole among the isolates from the second time period
also raise concerns about possible antifungal resistance.

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Community-Acquired *Staphylococcus argenteus* Sequence Type 2250 Bone and Joint Infection, France, 2017

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We report a rare case of *Staphylococcus argenteus* bone and joint infection in a 9-year-old boy in France. His finger arthritis was complicated by osteitis 5 weeks later, which resulted in a secondary intervention. This case indicates the virulence of *S. argenteus*, an emerging pathogen whose clinical effects are poorly described.

*S. argenteus* (formerly *S. aureus* clonal complex 75) is an emerging species in the *S. aureus* complex (1). Several studies reported sporadic cases of *S. argenteus* infections mainly in Asia, Oceania, and the Pacific Islands (2) but rarely in Europe (3). We report the clinical characteristics of a community-acquired bone and joint infection with *S. argenteus* in a child living in France.

At the end of July 2017, a 9-year-old boy with no unusual medical history or previous local trauma was hospitalized because of acute signs of infection of the third finger on his right hand. He was first seen in a local hospital and given an initial diagnosis of cellulitis (arthritis). Two days later, he was admitted to the emergency pediatric ward of a tertiary care hospital where a surgical joint exploration was performed and confirmed the diagnosis of arthritis associated with an abscess of the extensor tendon sheath (Table).

Surgical microbiological samples cultured on blood agar plates (aerobic conditions at 37°C for 24 h) grew a strain that was identified by matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (Microflex LT, Bruker, France) as having log scores ranging from 1.39 to 1958 Emerging Infectious Diseases • www.cdc.gov/eid • Vol. 24, No. 10, October 2018