High-level Relatedness among *Mycobacterium abscessus* subsp. *massiliense* Strains from Widely Separated Outbreaks

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Three recently sequenced strains isolated from patients during an outbreak of *Mycobacterium abscessus* subsp. *massiliense* infections at a cystic fibrosis center in the United States were compared with 6 strains from an outbreak at a cystic fibrosis center in the United Kingdom and worldwide strains. Strains from the 2 cystic fibrosis outbreaks showed high-level relatedness with each other and major-level relatedness with strains that caused soft tissue infections during an epidemic in Brazil. We identified unique single-nucleotide polymorphisms in cystic fibrosis and soft tissue outbreak strains, separate single-nucleotide polymorphisms only in cystic fibrosus outbreak strains, and unique genomic traits for each subset of isolates. Our findings highlight the necessity of identifying *M. abscessus* to the subspecies level and screening all cystic fibrosis isolates for relatedness to these outbreak strains. We propose 2 diagnostic strategies that use partial sequencing of *rpoB* and *secA1* genes and a multilocus sequence typing protocol.

Nontuberculous mycobacteria (NTM) and, in particular, the *Mycobacterium abscessus* group are recognized as emerging respiratory pathogens among patients with cystic fibrosis. Reports from the United States, France, and Israel have shown that the *M. abscessus* group accounts for a major proportion of NTM infections in patients with cystic fibrosis; prevalence rates range from 16% to 48% (1–3).

Previous studies have indicated great diversity within *M. abscessus* group strains among cystic fibrosis patients, suggesting independent acquisitions of NTM from the environment (2,4). However, suspicion of patient-to-patient transmission arose with the recent report of an outbreak of respiratory infection with *M. abscessus* subsp. *massiliense* at a cystic fibrosis center in Seattle, Washington, USA (5). The index case-patient and 4 additional patients all had multidrug-resistant isolates with resistance to amikacin and clarithromycin. All 5 strains were indistinguishable by repetitive unit sequence–based PCR patterns and pulsed-field gel electrophoresis analysis, which led to initiation of whole-genome sequencing. In a separate, recent study, whole-genome sequencing and epidemiologic analysis provided strong support for patient-to-patient transmission in 2 clustered outbreaks of *M. abscessus* subsp. *massiliense* at the Papworth Hospital Cystic Fibrosis Centre (Cambridge, UK) (6). Isolates from both clusters showed resistance to clarithromycin, and isolates from one of the clusters also had mutations conferring resistance to amikacin.

The availability of whole-genome sequences from different *M. abscessus* subsp. *massiliense* outbreaks, as well as unrelated strains, provides an unprecedented opportunity for multigenome comparisons. We conducted a genomic study of 3 recently sequenced strains from the Seattle cystic...
fibrosis outbreak, including the index strain, and compared them with representative strains from the Papworth cystic fibrosis outbreak, as well as with available strains from the United Kingdom, the United States, Brazil, South Korea, France, and Malaysia (Table 1). We found high-level relatedness among strains from the 2 geographically distant outbreaks in Seattle and Papworth. We also identified shared and unique genomic traits for strains from both cystic fibrosis outbreaks and for those from an outbreak of soft tissue infections in Brazil.

Materials and Methods

Sequence Analysis of Outbreak Strains

A subset of 6 isolates (2u, 12c, 14h, 19f, 20h, and 28c) representing the breadth of genomic diversity observed within the Papworth cystic fibrosis outbreak clusters 1 and 2 (6) were selected. Illumina sequencing reads from each of these isolates were assembled into sets of contigs by using Velvet software (21). These contigs were combined with draft genome sequences of the Seattle cystic fibrosis outbreaks and available whole-genome sequences of *M. abscessus* subsp. *massiliense* (Table 1) and subjected to whole-genome multiple sequence alignments by using Mugsy software (22). Core segments of the alignment that are shared among all isolates included in the analysis were identified and concatenated by using Phylomark software (23). Concatenated nucleotide sequences, including single-nucleotide polymorphisms (SNPs), were then used for construction of a neighbor-joining phylogenetic tree by using MEGA software (24). The use of microbial samples and data was approved by the ethics committees at each of the institutions involved.

To replicate data from the Papworth cystic fibrosis outbreak clusters 1 and 2 (6) by using a similar approach, we mapped sequencing reads from the subset of 6 Papworth isolates, together with reads from the 3 Seattle cystic fibrosis isolates and soft tissue strain CRM-0020 from Brazil (Table 1), onto the *M. abscessus* type strain ATCC 19977T reference genome by using BWA software (25). Variants, including SNPs, were called by using GATK software (26) and filtered for quality. The SNP panel was used for construction of a neighbor-joining phylogenetic tree by using MEGA software. The resulting tree replicated the topology of clusters 1 and 2 and showed that the Seattle isolates are most closely related to cluster 2.

PCR and In Silico PCR

Standard PCR and sequencing strategies were used to amplify and analyze partial sequences of the rpoB (723 bp) (27,28) and secA1 (465 bp) (29) genes. In addition, a multilocus sequence typing (MLST) scheme (29,30), including primers to 13 housekeeping genes (*cya, gdhA, argH, gipK, gnd, murC, pgm, pkaA, pta, pur, rpoB, hsp65*, and *secA1*) was used to conduct electronic PCR on the panel of 20 *M. abscessus* subsp. *massiliense* genomes (Table 1). Published forward primers for *cya* and *gdhA* (30) did not amplify in silico for some *M. abscessus* subsp. *massiliense* strains; therefore, the following new primers conserved across the *M. abscessus* group were used: *cya* _F_ new 5′-GCC TGC GTA AGG GTG ATG-3′ and *gdhA* _F_ new 5′-GTG AAG GTA AGG GTG ATG-3′. Alleles from each gene were

| Table 1. Twenty-four *Mycobacterium abscessus* group strain genomes analyzed for genetic relatedness* |
|---------------------------------------------------------------|
| **Subspecies/strain** | **Country** | **Outbreak** | **GenBank accession no.** | **Reference** |
|----------------------|-------------|--------------|--------------------------|--------------|
| Mm/2u                | UK          | Papworth     | NA                       | (6)          |
| Mm/12c               | UK          | Papworth     | NA                       | (6)          |
| Mm/14h               | UK          | Papworth     | NA                       | (6)          |
| Mm/19f               | UK          | Papworth     | NA                       | (6)          |
| Mm/20h               | UK          | Papworth     | NA                       | (6)          |
| Mm/28c               | UK          | Papworth     | NA                       | (6)          |
| Mm/2B-0107           | USA         | Seattle      | AKUN000000000             | This study   |
| Mm/MAB_082312_2258   | USA         | Seattle      | AYTA000000000             | This study   |
| Mm/MAB_091912_2446   | USA         | Seattle      | AYTF000000000             | This study   |
| Mm/CRM-0020          | Brazil      | Rio de Janeiro | ATFC000000000           | (7)          |
| Mm/GO-06             | Brazil      | Goiás        | CP003699                 | (8)          |
| Mm/47J26             | UK          | Not applicable | AGQU010000000           | (9)          |
| Mm/M18               | Malaysia    | Not applicable | AJSC010000000           | (10)         |
| Mm/M115              | Malaysia    | Not applicable | AJL200000000            | (11)         |
| Mm/M139              | Malaysia    | Not applicable | AKVR010000000           | (12)         |
| Mm/M154              | Malaysia    | Not applicable | AJMA010000000           | (13)         |
| Mm/Asan 50594        | South Korea | Not applicable | CP004374–CP004376       | (14)         |
| Mm/15-151–930        | USA         | Not applicable | AKUI000000000           | This study   |
| Mm/55-0817           | USA         | Not applicable | AKUB000000000           | This study   |
| Mm/CCUG 48898T       | France      | Not applicable | AKVF010000000           | (15,16)      |
| Mm/CF                | France      | Not applicable | CAHZ000000000           | (17)         |
| Mm/ATCC 19977T       | USA         | Not applicable | CU458896,CU458745       | (18)         |
| Mm/BD                | France      | Not applicable | AHAS000000000           | (19)         |
| Mm/M24               | Malaysia    | Not applicable | AJLY000000000           | (20)         |

*M, *M. abscessus* subsp. *massiliense*; NA, not available; Ma, *M. abscessus* subsp. *abscessus*; Mb, *M. abscessus* subsp. *boleti*. 
extracted and concatenated for each genome, the panel of concatenated sequences was aligned by using ClustalW software (31), and the core segments of the alignment were used for construction of a neighbor-joining phylogenetic tree by using MEGA software.

Results

Phylogenetic Characteristics of Outbreak Strains

A core genome phylogenetic tree (Figure 1) showed a tight cluster of the 3 Seattle cystic fibrosis outbreak strains. The Seattle cystic fibrosis cluster was closely related to the 2 cystic fibrosis clusters described for the Papworth outbreak (6) and the Birmingham, UK, cystic fibrosis isolate 47J26 (9). Furthermore, the Seattle and Papworth cystic fibrosis outbreak strains showed some relatedness to strains CRM-0020 and GO-06 derived strains (known collectively as BRA-100) isolated during an epidemic of soft tissue infections in Brazil (32) and the M. abscessus subsp. massiliense M18 strain from Malaysia (10).

The cumulative size of core segments of Mugsy alignments provides information on relatedness among groups of strains compared. The core genome reduces in size as more genomes are added; an expected major decrease occurs after addition of more distant strains to the group. The average genome size of cystic fibrosis outbreak strains was 4.81 Mb for Seattle (n = 3) and 4.97 Mb for Papworth (n = 6). The Seattle and Papworth cystic fibrosis outbreak strains (n = 9) shared a core genome of 4,264,844 nt, which is almost unchanged by including the Birmingham cystic fibrosis strain 47J26 (n = 10; 4,264,127 nt). Addition of the soft tissue outbreak strain CRM-0020 from Brazil (n = 11) (32) decreased the core to 4,231,390 nt, and adding the related outbreak strain GO 06 from Brazil (n = 12) (8,33), led to an additional decrease in the core genome to 4,043,718 nt. As expected, including unrelated available clinical M. abscessus subsp. massiliense strains (n = 20, including M139 with ambiguous subspecies taxonomic assignment (12), reduced the core genome size to 3,869,950 nt. Further addition of M. abscessus subsp. abscessus (n = 2) and M. abscessus subsp. bolletii (n = 2) genomes (Table 1) reduced the core to 3,828,656 nt.

Genomes of Strains from Cystic Fibrosis and Soft Tissue Infection Outbreaks

The core genome of 10 strains representing the Papworth cystic fibrosis (n = 6), the Seattle cystic fibrosis (n = 3), and soft tissue CRM-0020 (n = 1) outbreaks comprised 4,231,390 nt. Strain GO 06 was excluded from the analysis because its genome harbors a large number of ambiguous nucleotides and an unusual hybrid appearance with fragments of M. abscessus subsp. massiliense and M. abscessus subsp. abscessus sequences. Strains 47J26 and M18, isolated from the sputum of a cystic fibrosis patient in Birmingham, UK, and a lymph node sample from a patient in Malaysia, respectively, were related to the outbreak strains...
(Figure 1). However, no information was available about any epidemiologic link between cystic fibrosis strain 47J26 to reported or unpublished outbreaks, and no clinical information was available about the patient from whom strain M18 was isolated. Therefore, both strains were excluded from the SNP analysis. Nevertheless, SNPs for these 3 strains at positions relevant to the outbreak strains are shown in the Technical Appendix (wwwnc.cdc.gov/EID/article/20/3/13-1106-Techapp1.xlsx).

A total of 293 identical SNPs in the core segments of Mugsy alignments were shared by the 10 outbreak strains but were different in available M. abscessus subsp. massiliense strains not related to outbreaks (Figure 2; online Technical Appendix). Of the 293 SNPs, 95 gave rise to nonsynonymous mutations in several genes, including virulence factors (mammalian cell entry and yrbE proteins), transcriptional regulators (TetR family), and lipid metabolism genes (online Technical Appendix). Eleven SNPs were shared only by Papworth and Seattle cystic fibrosis outbreak strains (n = 9), including nonsynonymous mutations in the prepprotein translocase secA1 and a putative lyase (Figure 2; online Technical Appendix). Sixteen SNPs were shared only by the 3 Seattle cystic fibrosis outbreak strains, including nonsynonymous mutations in a mycobacterial large membrane protein (MmpL) family involved in lipid transport and virulence (34) and genes involved in amino acid and energy metabolism (Figure 2; online Technical Appendix). Eighty-six SNPs were present only in strain CRM-0020 (soft tissue outbreak) from Brazil, including nonsynonymous mutations in an MmpL family protein; transcriptional regulators; and lipid, amino acid, and energy metabolism genes (Figure 2; online Technical Appendix).

Having shown high-level relatedness among Papworth and Seattle cystic fibrosis outbreak strains and their relatedness to the soft tissue outbreak strains from Brazil, we also searched for genomic regions ≥200 nt outside the core genome that were specific to subsets of isolates. A single region of ≈11.5 kb was unique to the Papworth cystic fibrosis isolates (n = 6) and encompassed 2 conserved hypothetical proteins, 2 phage integrase family proteins, and an MmpL family protein. Alignment of the MmpL family protein with distinct MmpL proteins described above for the Seattle cystic fibrosis outbreak and the Brazil soft tissue outbreak showed diversity at several amino acid residues in all 3 proteins.

No region was unique to the Seattle cystic fibrosis isolates (n = 3). The soft tissue isolate CRM-0020 from Brazil harbored several large unique regions, including a previously described broad-host-range IncP-1β plasmid (35) and 3 regions (contigs) of 5 kb, 10.6 kb, and 79 kb of unknown origin encoding almost exclusively hypothetical proteins.

We also searched for polymorphisms associated with macrolide and aminoglycoside resistance. The Papworth cystic fibrosis and Seattle cystic fibrosis outbreak set of strains showed an A2058C/G mutation in 23S rRNA, which conferred macrolide resistance (36) (A2058G in strains 2u and 28c representative of Papworth cluster 2 and the Seattle strains). Strains 19f, 14h, 12c, and 28a, representative of Papworth cluster 1, and Seattle strains shared the A1408G mutations in 16S rRNA, which conferred aminoglycoside resistance (37). None of these mutations were found in the soft tissue outbreak strains CRM-0020 and GO 06 from Brazil or the M18 strain.

**Diagnostic Tools for Identification of Outbreak Strains**

In light of the possibility of a common ancestor and/or intercontinental transmission of strains, we identified SNPs in genes commonly used for identification of mycobacteria and an MLST scheme that could be used by clinical laboratories to assess relatedness of newly isolated strains to this global cluster. In the first approach, we retrieved rpoB sequences from the 6 genomes of representative strains of the Papworth cystic fibrosis outbreak and performed partial sequencing of the rpoB gene for selected isolates from the Seattle cystic fibrosis outbreak. We then compared these sequences with those of isolates from the outbreak in Brazil and unrelated clinical isolates comprising M. abscessus subsp. abscessus, massiliense, and bolletii, as well as other rapidly growing mycobacteria.
By using the \textit{rpoB} gene MAB\_3869c from the \textit{M. abscessus} subsp. \textit{abscessus} type strain as a reference (Table 2) described in the BRA-00 outbreak isolates from Brazil (32,33), we showed that Seattle (n = 4) and Papworth (n = 6) cystic fibrosis isolates carried the 2 \textit{rpoB} SNPs (C→T at position 2569 and T→C at position 2760). However, none of the \textit{M. abscessus} subsp. \textit{abscessus} or subsp. \textit{bolletii} or other rapidly growing mycobacterial isolates outside the \textit{M. abscessus} group harbored this 2-SNP \textit{rpoB} signature (Table 2). The second SNP (T→C substitution at position 2760) was present in several strains, but the combination of both \textit{rpoB} SNPs (C→T at position 2569 and T→C at position 2760) was not present. Most of the 26 \textit{M. abscessus} subsp. \textit{massiliense} strains not related to outbreaks tested did not harbor this 2-SNP \textit{rpoB} signature. However, 4 strains harbored this signature (Table 2) (29,38).

Multiple alignment of \textit{rpoB} sequences among available \textit{M. abscessus} subsp. \textit{massiliense} genomes showed the absence of the 2-SNP \textit{rpoB} signature in most strains. However, both SNPs were present in 1 strain not related to an outbreak (1S-151–0930) (Table 2).

Multiple alignment of \textit{secA1} sequences among available \textit{M. abscessus} subsp. \textit{massiliense} genomes showed a G→T substitution at position 820 (by using the \textit{secA1} gene MAB\_3869c from the \textit{M. abscessus} subsp. \textit{abscessus} type strain) shared by the Papworth and Seattle cystic fibrosis outbreak strains but not by the soft tissue outbreak strains from Brazil or additional unrelated strains. Further analysis of \textit{secA1} sequences from 12 \textit{M. abscessus} subsp. \textit{massiliense} identified by multitarget sequencing and PCR-based typing (29,38) showed a G→T substitution at position 820 in 2 strains unrelated to the outbreak (Table 2). Those 2 strains were included among the 4 strains that had the 2-SNP \textit{rpoB} signature. Although the SNPs described for \textit{rpoB} and \textit{secA1} were not 100% specific markers for the outbreak strains, these SNPs could be used for first-level identification of newly isolated strains as possibly being related to cystic fibrosis clusters or soft tissue outbreak strains from Brazil to be confirmed by a second assay.

We also developed a simple MLST protocol that could be used as a second confirmatory assay. Alleles for each of 13 housekeeping genes (\textit{cytA, gdhA, argH, glpK, gnd, murC, pgm, pknA, pta, pur, rpoB, hsp65, and secA1}) were extracted and concatenated for each \textit{M. abscessus} subsp. \textit{massiliense} genome (Table 1), and the panel of concatenated sequences was used for construction of a neighbor-joining phylogenetic tree by using MEGA software. The Seattle and Papworth cystic fibrosis outbreak strains grouped together in the tree with cystic fibrosis strain 47J26 and isolate M18 from Malaysia (Figure 3). Thus, partial sequencing of \textit{rpoB} and \textit{secA1} gens, followed by 13-target MLST analysis, could be used to rule out isolates as belonging to these 2 cystic fibrosis clusters.

### Table 2. Detection of \textit{rpoB} and \textit{secA1} SNP signature in the \textit{Mycobacterium abscessus} group and rapidly growing mycobacteria*

| Strains                    | \textit{rpoB} T 2569 | \textit{rpoB} C 2760 | \textit{rpoB} T 2569 and \textit{rpoB} C 2760 | \textit{secA1} T 820 |
|----------------------------|----------------------|----------------------|-------------------------------------------|----------------------|
| MAB (CSU)                  | 0/44 (0)             | 44/44 (100)          | 0/44 (0)                                  | NT                   |
| MAB (NIH)                  | 0/29 (0)             | 29/29 (100)          | 0/29 (0)                                  | NT                   |
| MMA non-outbreak strain (CSU) | 1/14 (7)             | 1/14 (7)             | 0/14 (0)                                  | NT                   |
| MMA non-outbreak strain (NIH) | 4/12 (33)            | 10/12 (83)           | 4/12 (33)                                 | 2/12 (17)            |
| MMA Seattle                | 4/4 (100)            | 4/4 (100)            | 4/4 (100)                                 | 3/3 (100)            |
| MMA Brazil                 | 9/9 (100)            | 9/9 (100)            | 9/9 (100)                                 | NT                   |
| MBO (CSU)                  | 0/11 (0)             | 11/11 (100)          | 0/11 (0)                                  | NT                   |
| MBO (NIH)                  | 0/2 (0)              | 2/2 (100)            | 0/2 (0)                                   | 0/2 (0)              |
| Other RGM (CSU)            | 0/42 (0)             | 0/42 (0)             | 0/42 (0)                                  | NT                   |
| MMA type strain            | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |

**MMA in silico data†**

| Strains | \textit{rpoB} T 2569 | \textit{rpoB} C 2760 | \textit{rpoB} T 2569 and \textit{rpoB} C 2760 | \textit{secA1} T 820 |
|----------|----------------------|----------------------|-------------------------------------------|----------------------|
| MMA UK   | 6/6 (100)            | 6/6 (100)            | 6/6 (100)                                 | 6/6 (100)            |
| MMA Seattle | 3/3 (100)           | 3/3 (100)            | 3/3 (100)                                 | 3/3 (100)            |
| MMA Brazil | 2/2 (100)            | 2/2 (100)            | 2/2 (100)                                 | 0/2 (0)              |
| 47J26    | 1/1 (100)            | 1/1 (100)            | 1/1 (100)                                 | 1/1 (100)            |
| M18      | 1/1 (100)            | 1/1 (100)            | 1/1 (100)                                 | 1/1 (100)            |
| 1S-151–0930 | 1/1 (100)         | 1/1 (100)            | 1/1 (100)                                 | 0/1 (0)              |
| 5S-0817  | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |
| M115     | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |
| M139     | 0/1 (0)              | 1/1 (100)            | 0/1 (0)                                   | 0/1 (0)              |
| M148     | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |
| M154     | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |
| M156     | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |
| M159     | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |
| M172     | 0/1 (0)              | 0/1 (0)              | 0/1 (0)                                   | 0/1 (0)              |
| Asan 50594 | 0/1 (0)            | 1/1 (100)            | 0/1 (0)                                   | 0/1 (0)              |

*SNP, single-nucleotide polymorphism; MAB, \textit{M. abscessus} subsp. \textit{abscessus}; NT, not tested. CSU, Colorado State University; NIH, National Institutes of Health; MMA, \textit{M. abscessus} subsp. \textit{massiliense}; MBO, \textit{M. abscessus} subsp. \textit{bolletii}; RGM, rapidly growing mycobacteria.

†Information collected from available whole genome sequencing data.
Discussion

The implications of this study are extensive. Currently, most experts recommend identifying isolates of *M. abscessus* to subspecies level (39). This report further corroborates these recommendations and places even greater pressure on clinical laboratories to fully identify *M. abscessus* subspecies *massiliense*.

Strains from the 2 cystic fibrosis outbreaks showed high-level relatedness (4,264,844 nt core genome alignment size, 11 shared unique SNPs) with each other and major-level relatedness (4,231,390 nt core genome alignment size) with soft tissue epidemic strains from Brazil. Genomic features shared between strains from all 3 breaks might make them more transmissible, whether from patient to patient (directly or indirectly as in cystic fibrosis outbreaks) or from a common source, as in soft tissue infections. However, the soft tissue strain from Brazil had the largest number of unique SNPs (86) not shared with either of the cystic fibrosis outbreak strains, harbored an IncP-1β plasmid, and did not show mutational resistance to amikacin or clarithromycin. We speculate that some of these specific genomic traits may be favorable for the successful establishment of epidemic soft tissue infections.

A previous study did not detect a common source or person-to-person transmission of the *M. abscessus* group among cystic fibrosis patients and suggested that it may not be necessary to segregate persons infected or colonized with *M. abscessus* from those who are not infected or colonized (40). Our findings emphasize the necessity of screening all isolates of *M. abscessus* subsp. *massiliense* recovered from patients with cystic fibrosis to outbreak strains in an effort to prevent future outbreaks. Because of evidence supporting patient-to-patient transmission of multiple different respiratory tract organisms, the Infection Control Guidelines (currently in draft form for public comment) of the United States Cystic Fibrosis Foundation (CFF) (www.cff.org/LivingWithCF/Webcasts/ArchivedWebcasts/Germs/#Infection_Prevention_and_Control_Policy_Update) have been recently changed. Patients with cystic fibrosis are advised not to attend indoor meetings with other cystic fibrosis patients (CFF and Infection Prevention and Control Guidelines 2013). In addition, screening of all cystic fibrosis patients in the United States...
at least annually for mycobacteria is now recommended (CFF and Infection Prevention and Control Guidelines 2013) to enable early treatment if the organism is detected.

It remains unclear why intercontinental organisms are so closely related. One hypothesis is that direct patient contact led to transmission. The Seattle index case-patient traveled to British Columbia, Canada, before and after acquiring mycobacterial infection, to Oregon before mycobacterial infection, and to Atlanta, Georgia, and Bethesda, Maryland, after mycobacterial infection. However, the patient did not report any contact with other cysitic fibrosis patients at these destinations. A second hypothesis is that the mycobacterial strain could have been carried by persons with cysitic fibrosis who were clinically well. A third hypothesis is that there was an independent selection of *M. abscessus* subsp. *massiliense* clones in the cystic fibrosis airway milieu on both sides of the Atlantic Ocean toward potentially more transmissible lineages. Availability of additional whole-genome sequencing data tracking the global epidemiology of the *M. abscessus* group may help differentiate between these scenarios. In addition, this data will help delineate global clusters of *M. abscessus* subsp. *massiliense* strains with potentially higher transmissibility.

**Addendum**

Recent whole-genome data show deep genetic separation of 3 subspecies, ruling against grouping *M. massiliense* and *M. bolletii* under *M. abscessus* subsp. *bolletii*.

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