

MAJOR ARTICLE

Genomic Analyses of Longitudinal *Mycobacterium abscessus* Isolates in a Multi-Center Cohort Reveal Parallel Signatures of In-Host Adaptation

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Background: Nontuberculous mycobacteria (NTM) are ubiquitous in the environment and an increasingly frequent cause of opportunistic infections. Mycobacterium abscessus complex

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(MAB) is one of the major NTM lung pathogens which disproportionately colonize and infect the lungs of individuals with cystic fibrosis (CF). MAB infection can persist for years, and antimicrobial treatment is frequently ineffective.

Methods: We sequence the genomes of 175 isolates longitudinally collected from 30 patients with MAB lung infection. We contextualize our cohort amidst the broader MAB phylogeny and investigate genes undergoing parallel adaptation across patients. Finally, we test the phenotypic consequences of parallel mutations by conducting antimicrobial resistance and mercury resistance assays.

Results: We identify highly related isolate pairs across hospital centers with low likelihood of transmission. We further annotate non-random parallel mutations in 22 genes and demonstrate altered macrolide susceptibility co-occurring with a nonsynonymous *whiB1* mutation. Finally, we highlight a 23kb mercury resistance plasmid whose loss during chronic infection confers phenotypic susceptibility to organic and non-organic mercury compounds.

Conclusions: We characterize parallel genomic processes through which MAB is adapting to promote survival within the host. The within-lineage polymorphisms we observe have phenotypic effects, potentially benefiting fitness in the host at the putative detriment of environmental survival.

Keywords: Nontuberculous mycobacteria, *Mycobacterium abscessus* complex, in-host adaptation, comparative genomics

BACKGROUND

Nontuberculous *Mycobacterium* spp. (NTM) are a diverse group of mycobacteria outside the *Mycobacterium tuberculosis* and *Mycobacterium leprae* complexes¹. Commonly found in soil and water², NTM are mostly considered environmental saprophytes. However, NTM can cause opportunistic infection, particularly in humans who are immunocompromised or have preexisting lung conditions such as chronic obstructive pulmonary disease (COPD), cystic fibrosis (CF), or non-CF bronchiectasis³. Within the NTM, *Mycobacterium abscessus* complex (MAB) disproportionately colonize and infect patients with CF^{4–7}. Colonization and infection can persist for years, and even with prolonged multidrug therapy⁸ only ~30% of patients experience treatment success⁹. With infection rates increasing worldwide⁵, MAB pose an imminent public health challenge.

The M. abscessus complex comprises three subspecies: Mycobacterium subsp. bolletii, Mycobacterium subsp. massiliense, and Mycobacterium subsp. abscessus¹⁰. Of these, over 50% of global clinical isolates have been shown to belong to three clusters of dominant circulating clones (DCCs)^{11,12}. However, it is unclear how frequently infection is caused by transmission between patients relative to independent environmental acquisition. Recent studies

demonstrating the intercontinental presence of highly related pairs of clinical isolates (less than 20 single nucleotide polymorphisms; SNPs) suggest transmission between patients may occur more often than previously thought^{13–14}, with some studies demonstrating epidemiological linkage such as overlapping clinic visit dates or outpatient visits by the same team of clinicians¹⁵. DCCs have also been found to exhibit elevated drug resistance and intracellular survival¹¹, suggestive of adaptation to a pathogenic lifestyle. However, epidemiological evidence of transmission is often lacking^{16–19}. It remains unclear whether the prevalence of DCCs is a result of recent transmission, widespread environmental presence, sampling bias, a slower mutation rate among DCCs maintaining fitness benefits in the lung milieu– or a combination of these factors. Thus, understanding how this environmental species survives and persists in the host is critical for elucidating MAB transmission and evolution.

Despite their ability to cause chronic infection, little is known of the in-host adaptive behavior of MAB. MAB is intrinsically resistant to many antimicrobials^{20,21} but can also acquire antimicrobial resistance through SNPs in ribosomal rRNA genes *rrl* and *rrs*²⁰. A smooth-to-rough morphotype switch via loss of glycopeptidolipid (GPL) expression on the cell surface has been shown to correspond with heightened virulence²², but the mechanisms regulating this process are unknown²³. To date, most studies investigating in-host adaptation of MAB have been limited to isolates from a single patient^{24–26}. One study querying longitudinal samples from 18 patients found evidence for convergent evolution in 30 genes, including the GPL locus and virulence regulators¹³. Some of these mutations impaired survival on fomites¹³, suggesting fitness trade-offs between the environment and host. Thus, utilizing high-resolution sequencing to explicitly investigate MAB in-host adaptation may inform prevention strategies and effective treatment development.

Here, we leverage a multi-center cohort of 175 isolates, longitudinally collected from 30 patients with MAB infection who were treated at academic medical centers in the United States between 2002 and 2020. Through whole genome sequencing (WGS) we first contextualized the genomic relatedness of isolates, incorporating an additional 1,455 published MAB genomes for comparative genomic analyses. Next, we investigated within-lineage genomic diversity and characterized parallel in-host adaptation. Finally, we probed differences in antimicrobial resistance and mercury compound resistance correlated with specific genomic variants. This work provides high-resolution insight into MAB adaptive mechanisms and identifies novel candidate genes for parallel adaptation. We highlight how MAB adapts to the host milieu while shedding obsolete functions, furthering our understanding of MAB evolution during chronic infection.

METHODS

Isolate collection

122 isolates from 22 patients were recovered from clinical specimens collected as a part of routine clinical care at the Barnes-Jewish Hospital (BJH) microbiology laboratory between 2002 and 2019. Another 53 isolates from 8 patients were obtained from clinical samples at Michigan Medicine (University of Michigan (UM)) between 2014 and 2020 (Supplementary material, Supplemental Table 1). Culture collection dates, clinic appointment dates, and pulmonary provider were reviewed for isolates with high genomic relatedness to assess potential epidemiological links.

Characterization of within-lineage diversity

Lineage-specific whole genome alignments were generated and annotated for SNPs and insertions/deletions (Supplementary material). For the permutation analysis, the total number of observed mutations per lineage was noted (based on the lineage-specific whole genome alignment). Next, the same number of mutations per lineage were randomly generated across each lineage's initial isolate's genome, and then annotated for which genes they landed on by mapping back to the prokka-generated gff file. For each round of permutation, the total number of lineages randomly mutated in parallel for a given gene was noted. This process was repeated for a total of 10,000 rounds to generate a hypothetical distribution (script: s_randommuts.py). P-values were calculated by calculating the percentile of the actual number of lineages within the hypothetical distribution (s_pval.R). Bubble plots were visualized in R v3.6.3²⁷ using ggplot2²³. PE/PPE genes were removed from the final list of significant hits due to their highly repetitive nature²⁹. The presence of a 23kb mercury plasmid was validated through BLASTN³⁰ (100% identity, 100% coverage).

Antimicrobial resistance assays

Isolates were inoculated into 7H9 broth supplemented with OADC. After aerobic growth to midlog phase in 37 °C, suspensions were diluted to 0.05 OD_{600} in 7H9, approximately 10⁸ CFU. In a 96-well plate, serial dilutions of each drug were prepared by adding 100 µL antimicrobial solution to 100 µL 7H9 broth. 100 µL of the diluted sample was added to each well and mixed by pipetting up and down. The plate was placed in a sealed container to grow shaking for 4 days in 37 °C in air, after which 10uL 0.02% resazurin was added to each well. After 24 hours (shaking in 37 °C in air), the minimum inhibitory concentration (MIC) was recorded as the minimum concentration observed to inhibit cell growth (determined from a color change from blue to pink). ATCC19977 was used as a control strain, along with a positive control row (containing just sample and 7H9 broth), and negative control row (containing just 7H9 broth) for each sample.

Mercury resistance assays

Isolates were grown out to mid-log phase and diluted to 0.05 OD_{600} as described above. Mercury resistance was tested in the methodology described by Steingrube et al³¹: 600uL of each isolate suspension was spread on an 7H10 plate supplemented with ADC. 6mm discs were loaded with 20 µL HgCl₂ or PMA at tenfold diluted concentrations: $10^{-2}M$, $10^{-3}M$, $10^{-4}M$, $10^{-5}M$. A blank disc was added as a negative control. The plates were set to grow for 72 hours at 37°C in air, after which the zone of inhibition was measured.

RESULTS

Cohort comprises two major subspecies of MAB spanning the global phylogeny

To understand the phylogenetic relationships in the cohort, we annotated the genomes of all 175 clinical isolates, as well as 3 NCBI reference genomes for each MAB subspecies (subsp. *abscessus*, subsp. *massiliense*, subsp. *bolletii*). We aligned 2,693 core genes to generate a maximum likelihood phylogenetic tree (Figure 1A, Supplemental Table 2). Most isolates (74.9%, 131/175) belonged to subsp. *abscessus*, and a smaller group of subsp. *massiliense* genomes (25.1%, 44/175) was also identified. No subsp. *bolletii* were present in the cohort. Therefore, the tree was re-rooted with the subsp. *bolletii* reference genome as the outgroup.

We then calculated pairwise average nucleotide identity (ANI) for all isolate pairs (Figure S1). All isolate pairs showed at least 97% ANI, fulfilling the genomic gold standard for microbial species³². Again, two major clusters were identified corresponding to each of the major subspecies present in the cohort. Pairwise comparisons within the same subspecies were at least 98.49% ANI (Figure 1B, Supplemental Table 3).

To contextualize our cohort within the global phylogeny, we conducted another core genome alignment (1,423 core genes), incorporating 1,452 additional published MAB genomes downloaded from NCBI and the European Nucleotide Archive (ENA) for a total of 1,630 genomes included in the analysis (Figure S2A, Supplemental Table 2, genomes downloaded Dec. 28, 2021). Most genomes in this larger cohort belonged to subsp. *abscessus* (72,1%, 1175/1630), followed by subsp. *massiliense* (27.2%, 444/1630). There were 8 subsp. *bolletii* genomes (0.49%, 8/1630), all contributed from downloaded genomes. The 175 isolates in our study were broadly distributed throughout the species phylogeny, suggesting a large range of genomic diversity is represented in the study cohort. ANI was also measured against each of the 3 subspecies reference genomes, and again the same subspecies were determined to be at least 98.5% ANI against a reference genome (Figure S2B, Supplemental Table 3).

Genomic relatedness indicates within-patient diversity of subspecies and lineages

To define lineages, we measured the relatedness of intra-patient isolate pairs in the 175 isolate core genome alignment. When looking at the overall distribution of pairwise core genome SNP distances, isolate pairs > 40,000 SNPs apart belonged to different subspecies (Figure 1C, Figure S3). While same-patient isolate pairs generally exhibited low core genome SNP distance (median: 134; mean: 1,671; range: 1- 62,011), pairs of isolates belonging to the same subspecies but corresponding to large SNP distances (10,000-14,000 SNPs) were also identified, pointing to diverse MAB populations within some hosts (Figure 1C).

To further quantify within-host diversity, we applied pairwise ANI as an orthogonal approach and found that at a 99.99% ANI cutoff, we could distinguish clusters of highly related isolates (Figure 1C, Figure 2A). This cutoff corresponded to <400 whole genome SNPs and <150 cgMLST distance (Supplemental Table 4). While most clusters comprised isolates from a single patient, there were also groups of closely related isolates found across multiple patients (L1-L4, Figure 2A). Interestingly, these multi-patient groups were closely related (<200 core genome SNPs) to published DCC genomes (Supplemental Table 4). Whole genome alignment of multipatient clusters further revealed SNP distances ranging from 7 to 490 SNPs (mean: 177, median: 135, Supplemental Table 4, Supplementary material). Two patient pairs (MAB 04 and MAB_26, MAB_04 and MAB_30) exhibited DCC1 isolates less than 10 SNPs apart but were treated at different sites. Three other patient pairs exhibited DCC1 and DCC3 isolates less than 38 SNPs apart (suggested as an indicator of possible transmission¹¹), with one pair (MAB_25 and MAB_05) treated at different sites in different years, one pair (MAB_26 and MAB_30) treated at the same site in different years, and just one pair (MAB_06 and MAB_17) at the same site with overlapping treatment years, with first positive sputum occurring four months apart. A review of these patients' culture collection dates, clinic appointment dates, and pulmonary provider did not suggest an epidemiological link between the two. Overall, the observation of highly related isolate pairs from distinct locations and years suggests presence of nearly clonal DCCs on a broad geographic scale.

To contextualize the evolutionary history of MAB lineages, we generated maximum parsimony trees using PHYLIP³³. From these trees' ancestral nodes, we were able to infer the average distance from the Most Recent Common Ancestor (dMRCA) for each lineage (mean: 4.82 SNPs, 95% CI [2.41, 7.23]). We then applied the estimated molecular clock for each corresponding subspecies¹⁰ (1.8 SNPs/year for subsp. *abscessus*, 0.46 SNPs/year for subsp. *massiliense*) to each lineage and calculated the average time from Most Recent Common Ancestor (tMRCA, mean: 3.76 years, 95% CI [2.18, 5.34]). Comparison of these estimates with collection dates revealed that despite variation among lineages (Figure S4), average estimated tMRCA corresponded closely to the actual average time since initial positive NTM isolate culture of 3.57 years (95% CI [2.53, 4.62]), lending credence to the accuracy of our genomic measurements.

Antimicrobial Resistance Genes are prevalent and conserved within lineages

Next, we sought to characterize the antimicrobial resistance genes (ARGs) that may influence survival *in vivo* (Supplemental Table 5). We found that 100% (175/175) of isolates carry *bla*, *arr*, and *cmx_cmrA* genes, which confer resistance to beta lactams, rifamycin, and chloramphenicol, respectively. 92.6% (162/175) of isolates carried the aph(3") gene conferring resistance to streptomycin. 74.3% (130/175) of isolates, all subsp. *abscessus*, carried the *erm(41)* gene for macrolide (induced clarithromycin) resistance. We did not observe substantial within-lineage variance for ARGs, and in 97% of cases (33/34 lineages) whole lineages were identical in predicted ARG profile (Supplemental Table 5). The single instance of within-lineage ARG profile variability was observed in patient MAB_14, in which the initial isolate MAB_14_01 carried mercury resistance genes *merA* and *merB*, but subsequent isolates did not.

Lineages undergo parallel within-host adaptation in Mycobacterial virulence genes

We sought to identify adaptive mutations that may have conferred fitness advantages for longterm survival in the host. For each lineage, we identified polymorphisms by aligning isolate reads against the genome from the lineage's earliest collection time and annotating whole genome SNPs and insertions/deletions (indels). At this higher resolution, isolate pairs from the same lineage and patient were on average 10 SNPs apart (95% CI [8.74, 11.3], Supplemental Table 4).

In total we found 28 genes mutated in parallel across multiple patients and lineages (Figure 2B, Supplemental Table 6). We then applied a permutation test with 10,000 iterations to assess the significance (non-randomness) of these parallel findings (Figure 2B, Supplemental Table 6, Methods). 78.6% (22/28) of the genes were also significant by the permutation test (p-value < 0.05, Benjamini Hochberg). Some of these genes were also found to be variable in isolates collected just 9 days apart from multiple body sites (Figure 2C). Four genes (*crp*, *embC*, *whiB1*, and *espR*) were previously reported in the literature as evidence of within-host parallel evolution¹³, suggesting diverse populations of MAB undergo similar adaptive trajectories during chronic infection.

Within-lineage diversity affects phenotypic antimicrobial resistance

We sought to test the phenotypic effects of parallel mutated genes by examining a lineage from patient MAB_18, which featured nonsynonymous variation in the coding region of the *whiB1* gene (Figure 3A). *whiB1* (MAB_3539) encodes for a nitric oxide-sensitive transcriptional repressor in the WhiB family of proteins, and has been implicated in regulation of the ESX-1 secretion system in *M. tuberculosis*³⁴. Deletion of the *whiB7* gene has been reported to confer sensitivity to the ribosome targeting drugs amikacin, clarithromycin, erythromycin, tetracycline, and spectinomycin³⁵. In MAB_18, three isolates exhibited a nonsynonymous *whiB1* mutation resulting in a glycine (Gly) to alanine (Ala) switch at the Gly24 locus conserved between *M. tuberculosis* H37rV and *M. abscsessus* ATCC 19977. We hypothesized that the nonsynonymous

whiB1 mutation at this conserved locus would affect clinical isolates' antimicrobial susceptibility. Thus, we measured their MICs in amikacin, erythromycin, azithromycin, and clarithromycin using a resazurin microplate assay^{36,37} (Methods). We observed that the three isolates with a *whiB1* mutation were significantly more susceptible to erythromycin and clarithromycin, but not amikacin or azithromycin (Figure 3B, clarithromycin p=0.001053, erythromycin p=0.000789, Kruskal-Wallace rank sum test), demonstrating within-lineage variability in phenotypic resistance corresponding to *whiB1* mutation.

Loss of 23kb mercury resistance plasmid is linked to mercury susceptibility

We observed the loss of a 23kb contig carrying mercury resistance genes in isolates from patient MAB_14. This patient had seven isolates belonging to two distinct lineages (Figure 4A). The initial isolate MAB_14_01 genome contained a 23kb mercury resistance plasmid identical to one in ATCC19977, reported to have originated in *M. marinum*^{38,39} and predicted to encode a 472 amino acid (AA) MerA protein and 218 AA MerB protein. This plasmid was not present in subsequent isolate genomes of the same lineage (MAB_14_02 through MAB_14_05). Instead, these genomes contained a predicted 281 AA MerB in the chromosome, in a region encoding for cell wall components MmpL4 and PPE4 (Figure 4C). This region was highly conserved across all subsp. *abscessus* isolates (Supplemental Table 7). In contrast, the MAB_14_06 and MAB_14_07 genomes of a separate lineage contained a 7kb contig carrying both predicted 474 AA MerA and 219 AA MerB (Figure 4B).

MerA is a mercuric reductase which reduces inorganic mercury Hg(II) to Hg(0), while MerB is an organic mercury lysase which cleaves the Hg-C bond in organic mercury compounds to generate inorganic mercury Hg(II) (Supplementary material). We sought to validate the phenotypic differences between isolates in MAB_14 that had different combinations and alleles of *merA* and *merB*, by conducting disc diffusion assays with an inorganic mercury compound, mercury chloride (HgCl₂), and an organic mercury compound, phenylmercury acetate (PMA, Methods). We found striking differences in phenotypic resistance to both compounds between the two genotypes (Figure 4D). Isolates encoding *merA* had higher resistance to both PMA and HgCl₂, but this difference was significant only at the highest concentration of the inorganic compound, HgCl₂ (Kruskal-Wallis test, p=0.011). In contrast, resistance to the organic compound PMA was significantly higher in the *merA* isolates across three different concentrations (Kruskal-Wallis test, p=0.011, 0.0064, 0.0016), despite *merB* being present in all tested isolates.

DISCUSSION

The possibility of patient-to-patient or fomite-directed transmission of MAB has been debated^{10,15,16,18,19}. Here, we found instances of highly related isolate pairs (<20 whole genome SNPs) across different hospital centers as well as within the same center. We were unable to

substantiate transmission in the current study, without formal epidemiological investigation. Importantly, we did not conduct environmental sampling of the healthcare centers, which have previously been linked to MAB reservoirs and outbreaks^{14,40}. Future studies will benefit from conducting both genomic and epidemiological investigation. Barring the existence of an undiscovered clinical reservoir, these occurrences may indicate widely circulating lineages of MAB acquired through separate infection events as has been suggested by others^{16–19}. The high genomic relatedness of these isolates could be explained by a slow rate of mutation among highly successful host-adapted pathogens. Whether this is a species-wide trend towards obligate pathogenicity (as the case for *M. tuberculosis* and *M. leprae*)^{13,41}, or driving a divide between clinical and environmental populations of MAB, warrants exploration. Further research on the species' molecular clock and genomic comparisons with environmental isolates will provide more clarity on the transmission dynamics and evolutionary trajectory of MAB.

We identify candidate genes as hotspots of in-host adaptation, potentially conferring advantages for survival within the lung milieu. Though average tMRCA corresponded closely to average time difference between first and last isolate collection times, tMRCA values derived from SNP distances trended higher than actual collection times within each lineage, suggestive of great-than-expected genomic diversity. It is possible that even greater diversity of MAB was present but not captured due to limitations in study design. Here we only obtained one isolate per timepoint, but there may be additional co-existing lineages, or lineages that emerged prior to the dates to which we attributed them. Furthermore, CF patients exhibit polymicrobial lung infections⁴², and cross-species interactions such as horizonal gene transfer or competition may occur. Future studies may capture more diversity by picking multiple colonies⁴³, conducting plate sweeps^{11,16}, or conducting metagenomic sequencing of sputum samples^{44,45}.

We demonstrated *in vitro* that within-lineage variations are associated with diverse phenotypic susceptibility to drugs. We found that a nonsynonymous mutation at a conserved site in *whiB1* was associated with increased susceptibility to clarithromycin and erythromycin, but not amikacin or azithromycin. The patient was on azithromycin therapy for CF at earlier timepoints (MAB_18_01 through MAB_18_05), while latter isolates (MAB_18_07 through MAB_18_09) were exposed to azithromycin and cefoxitin, and amikacin (Figure S5). WhiB proteins are involved in regulation of pathogenesis, cell division, as well as oxidative stress response²⁰. It is possible that *whiB1*, similar to *whiB7*²⁸, regulates antimicrobial resistance. Thus, the observed *whiB1* mutation may pose a trade-off between increased antimicrobial susceptibility and greater in-host survival through differences in transcriptional regulatory pathways of *whiB1* in MAB is required to test this hypothesis. Further studies investigating treatment outcome in relation to prevalence of MAB parallel mutations may reveal additional fitness trade-offs which may be exploited for treatment development.

Finally, we demonstrate how the loss of a 23kb mercury resistance plasmid is correlated with increased susceptibility to both organic and inorganic mercury compounds. Mercury exists

naturally in the soil, water, and atmosphere, and cycles globally through processes such as industrial wastewater, landfills, burning fossil fuels, and processing through microorganisms⁴⁶. Bacteria use MerA and MerB to break down organomercury into Hg(II), and subsequently to elemental mercury (Hg(0)), which is highly volatile and rapidly diffused out of the bacterial cell^{47,48}. A study of *Arabidopsis thaliana* found that insertion of bacterial *merA* and *merB* genes together conferred tenfold higher resistance to organic methylmercury than insertion of *merB* alone⁴⁸. These findings reflect our own observations, and potentially signify that both genes are imperative to successfully break down and remove mercury from the bacterial cell. While we cannot rule out the possibility of plasmid loss during subcultivation, the consistent finding of the mercury plasmid among temporally initial isolates suggests loss of the mercury resistance plasmid is likely due to the fitness trade-off of maintaining the plasmid in the low-mercury pulmonary environment^{49,50}.

In this study we highlight genomic processes through which MAB adapts to promote its own survival within the host. Many of these events occur in parallel across patients and hospital sites and include DCCs circulating on a global scale. We suggest highly infectious strains of MAB may exhibit low rates of mutation to maintain a pathogen lifestyle. Further, the within-lineage polymorphisms we observed have phenotypic effects, potentially benefiting fitness in the host, at the putative detriment of environmental survival. This work thus contributes to our understanding of in-host survival of MAB and may inform development of treatment strategies against these chronic infections.

FIGURE TITLES AND LEGENDS

Figure 1. Genomic comparisons of entire cohort (A) Core genome alignment of 175 isolate genomes with 3 reference genomes for each *M. abscessus* subspecies. Aligned with roary, converted with FastTree, visualized with iTOL. 2,693 core genes. Tree is rerooted at the *M. bolletii* reference genome. Branch colors indicate subspecies determined by clade: green for subspecies *massiliense* (n=44), purple for subspecies *abscessus* (n=131). Outer rings (from inner to outer) denote patient ID, sample source, and study site (MO: Missouri, MI: Michigan, BAL: bronchoalveolar lavage). (B) ANI across isolate pairs. X-axis shows comparisons between different subspecies have pairwise ANI values above 98.5%. (C) Core genome SNP distance across isolate pairs. Each point represents a pairwise comparison. Comparisons are grouped as distances between isolates from different patients or the same patient. Points are colored by subspecies comparison as well as corresponding ANI: different subspecies pairs are purple, highly related pairs of at least 99.99% ANI are turquoise, and less related pairs less than 99.99% ANI are salmon.



Figure 2. Multiple subspecies or lineages coexist within four patients (A) Network visualization of isolate genomes at least 99.99% ANI. Nodes indicate genomes and edges indicate a pairwise ANI value of at least 99.99%. Colors indicate patient ID; nodes of the same color are isolates from the same patient. Clusters have been divided into two panels to distinguish clusters of isolates coming from a single patient from multi-patient clusters. Multipatient clusters are labeled L1-L4. Nodes highlighted with higher opacity are isolates from patients with multiple subspecies present (MAB 20, blue) or multiple lineages (MAB 07, green; MAB_14, pink; MAB_28, orange). Two isolates without any 99.99% ANI matches are not pictured: MAB 22 01 and MAB 27 02. ANI measured with dnadiff, clusters visualized on Cytoscape. (B) Bubble plot displaying results of permutation analysis. Mutations were randomly distributed across representative isolate genomes to generate a neutral (expected) distribution for parallel mutations across lineages. This distribution was then compared with the observed number of lineages with mutations in each gene. X-axis denotes position in the ATCC19977 reference genome, Y-axis denotes negative log p-value. Bubble size corresponds to number of lineages the gene was found to be mutated in. Genes mutated in at least 3 lineages are colored and named. Area with grey background indicates p-value < 0.05. (C) Unrooted tree showing whole genome SNP distances between isolates from MAB_21, which were collected within 9 days of each other. X-axis is SNP distance, tree nodes contain isolate IDs. Nodes are also annotated for site of collection and key mutations observed. SNPs annotated by aligning reads against initial isolate genome MAB 21 01.



Figure 3. MAB_18 isolates show varied macrolide susceptibility (A) Unrooted tree showing whole genome SNP distances between isolates from MAB_18. X-axis is SNP distance, tree nodes contain isolate IDs. Nodes are also annotated for key observed mutations. (B) Results of antimicrobial resistance assay. Each panel indicates a tested drug. Isolates (n=9) are grouped by observed mutation: MAB_18_03, MAB_18_04, MAB_18_05 are categorized as "whiB1" and the remaining MAB_18 isolates are "18-1". ATCC19977 was included as a control ("wt"). Significant differences among the groups were observed for clarithromycin (p=0.001053) and erythromycin (p=0.000789, Kruskal-Wallace rank sum test). Background colors in the panel represent clinical interpretation according to CLSI M24-A2 guidelines: resistant (red), intermediate (yellow) or susceptible (green). Interpretations for erythromycin or azithromycin are not available and thus left blank.



Figure 4. Loss of mercury resistance genes affects mercury susceptibility (A) Unrooted tree showing whole genome SNP distances between isolates from MAB_14. X-axis is SNP distance and tree nodes contain isolate IDs. Nodes are also annotated for presence of merA or merB. Isolates MAB_14_01 through MAB_14_05 belong to one lineage, while MAB_14_06 and MAB_14_07 belong to a second lineage. (B-C) Genomic context of merA and merB genes across isolate genomes. merA is colored grey, while other coding sequences are orange. Each row is a visualization of an assembled contig containing merA or merB. Grey regions between contig rows indicate regions of high percent identity according to BLASTn. Visualized using easyfig. (D) Results of mercury resistance assays. Each clinical isolate was exposed to inorganic (HgCl₂) or organic (PMA) mercury compounds via disc diffusion assay. ATCC19977 was included as a control (purple). Isolates are grouped by genotype: merA and merB (MAB_14_01, MAB_14_07 and MAB 14 08, turquoise) or merB only (MAB 14 02 through MAB 14 05, orange). ATCC19977 is "wt" (purple) and contains a 23kb plasmid identical to MAB 14 01. Significant differences were observed between groups in HgCl₂ 10⁻² M (p=0.011, Kruskal-Wallace rank sum test), PMA 10⁻² M (p=0.011), PMA 10⁻³ M (0.0064), and PMA 10⁻⁴ (, 0.0016). 6mm indicates disc size and no zone of inhibition.



NTM: Nontuberculous Mycobacterium spp.

OADC: Oleic albumin dextrose catalase

PHYLIP: PHYLogeny Inference Package

PMA: Phenylmercury acetate

SNP: Single nucleotide polymorphism

UM: University of Michigan

WGS: Whole genome sequencing

Supplementary Data: The sequencing data supporting the conclusions of this article is available in the NCBI repository under Bioproject PRJNA882917. Custom scripts are available on Github: https://github.com/jhchoi17/MAB_inhost.

Ethics approval: This research was conducted under Institutional Review Board (IRB) ID # 201904074.

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