Environmental opportunistic mycobacteria, including *Mycobacterium avium*, *M. terrae*, and the new species *M. immunogenum*, have been implicated in outbreaks of hypersensitivity pneumonitis or respiratory problems in a wide variety of settings. One common feature of the outbreaks has been exposure to aerosols. Aerosols have been generated from metalworking fluid during machining and grinding operations as well as from indoor swimming pools, hot tubs, and water-damaged buildings. Environmental opportunistic mycobacteria are present in drinking water, resistant to disinfection, able to provoke inflammatory reactions, and readily aerosolized. In all outbreaks, the water sources of the aerosols were disinfected. Disinfection may select for the predominance and growth of mycobacteria. Therefore, mycobacteria may be responsible, in part, for many outbreaks of hypersensitivity pneumonitis and other respiratory problems in the workplace and home.

Hypersensitivity pneumonitis is an occupational hazard of workers in two different industries, automobile manufacturing (e.g., metal working) and leisure (e.g., indoor swimming pools). Pulmonary illness and infection have also been a consequence of exposure to aerosols generated by hot tubs, spas, and coolant baths. Respiratory problems have also been associated with exposure to water-damaged buildings during reconstruction, and mycobacteria isolated from materials from such buildings have been shown to provoke inflammatory reactions. The outbreaks share the common feature of aerosol exposure and respiratory illness. I propose that exposure to aerosols containing mycobacteria is a common feature of the outbreaks and that mycobacteria or their products could be responsible for the respiratory symptoms.

Epidemiologic studies have established that the workers in such outbreaks were exposed to aerosols generated in the workplace from water that was either a work tool (e.g., metalworking fluid) or an integral part of the workplace or household (e.g., swimming pools and hot tubs) (1–7). Outbreaks of respiratory disease occurred in spite of disinfectant treatment of the waters or fluids to reduce the number of microorganisms. Living or working in water-damaged buildings or as a consequence of reconstruction of water-damaged buildings has also been associated with outbreaks of respiratory problems (8,9). Respiratory disease has been associated with mycobacteria in reservoirs, aerosols, or structural material in a number of cases (2,3,6,7,9).

Hypersensitivity Pneumonitis in Workers Exposed to Metalworking Fluid

An estimated 1.2 million workers in the United States are exposed to aerosols generated by metal grinding (10). Metalworking fluids are widely used in a variety of common industrial metal-grinding operations to lubricate and cool the tool and the working surface. Metalworking fluids are oil-water emulsions that contain paraffins, pine oils, polycyclic aromatic hydrocarbons, and heavy metals (10,11). Exposure to metalworking fluid aerosols can lead to hypersensitivity pneumonitis and chronic obstructive pulmonary disease (1,6,12–14). Mycobacteria were recovered significantly more frequently from metalworking fluid samples collected from facilities where hypersensitivity pneumonitis was found; compared to facilities that did not have hypersensitivity pneumonitis (6). In one study, exposure to metalworking fluid mist resulted in hypersensitivity pneumonitis in 10 workers (7). Acid-fast microorganisms identified as mycobacteria were present in the reservoir at 10⁷ CFU/mL (7). A mycobacteria in the reservoir was considered to be a likely cause of the hypersensitivity pneumonitis because one patient was infected by a *Mycobacterium* sp. and had antibodies against the reservoir fluid (7).

Hypersensitivity pneumonitis appeared in spite of disinfection of the metalworking fluid with morpholine, formaldehyde, or quaternary ammonium-based disinfectants (1,6,12,13), and mycobacteria were recovered from the metal working fluid (6,14,15). Mycobacteria are resistant to formaldehyde and quaternary ammonium disinfectants (16) and the heavy metals in metalworking fluids (17). Further, mycobacteria can grow on the organic compounds in metalworking fluid, including the paraffins, pine

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oils, and polycyclic aromatic hydrocarbons (18,19) and can degrade the disinfectant morpholine (20). Mycobacteria present in the water (21) can likely grow on the organic compounds in metalworking fluids in the absence of competitors after disinfection. Cleaning would not be expected to eradicate mycobacteria because of their ability to form biofilms (21,22). Adding disinfectant and cleaning the reservoir in one facility did not prevent the reappearance of mycobacteria (7 x 10^5 CFU/mL by 2 weeks [7]). Further, disinfectant treatment would likely result in selection of mycobacteria remaining after the cleaning.

**Hypersensitivity Pneumonitis in Swimming Pool Attendants**

Granulomatous pneumonitis has been reported in lifeguards (“lifeguard lung”) who worked at an indoor swimming pool that featured waterfalls and sprays (5). Affected lifeguards with symptoms worked longer hours than unaffected lifeguards (5), which demonstrated a dose-response effect. The waterfalls and sprays increased the number of respirable particles fivefold and the levels of endotoxin eightfold (5). Based on the presence of endotoxin in the aerosol samples, endotoxin exposure was suggested as the cause of the pneumonitis in lifeguards (5). However, subsequent data provided evidence of a possible second factor resulting in hypersensitivity pneumonitis; aerosols containing mycobacteria were shown to cause granulomatous lung disease (4). Others have reported high numbers of mycobacteria in swimming pools and whirlpools (23) and in hot tubs (2,3,24). Further, amoebae were reported in the indoor swimming pool where lifeguards reported pneumonitis (5). Mycobacteria, including *M. avium* and *M. intracellulare*, can survive and grow in phagocytic amoebae (25) and protozoa (26). In fact, *M. avium* grown in amoebae or protozoa are more virulent (25; Falkinham JO, unpub. data). Mycobacteria are resistant to chlorine (27) and preferentially aerosolized from water (28).

**Mycobacterial Disease after Exposure to Aerosols Generated by Hot Tubs**

Hypersensitivity pneumonitis and mycobacterial pulmonary disease has been reported after exposure to hot tubs (2,3,24). The mycobacteria isolated (e.g., *M. avium*) were likely responsible for the infections based on the identity of patient and hot tub mycobacterial isolates by either restriction fragment length polymorphism analysis (24) or multilocus enzyme electrophoresis (2,3). Further, exposure was followed closely by the onset of symptoms, and the extent of symptoms was related to the length of exposure (i.e., time spent in the hot tub) (2,24). Although these reports do not document the use of disinfectants in the hot tubs, the waters had been heated. Mycobacteria are relatively resistant to high temperature (29) and concentrated in hospital hot water systems (30).

**Hypersensitivity Pneumonitis in Occupants of Water-Damaged Buildings**

Inflammatory reactions—including eye irritation, respiratory infections, wheeze, bronchitis, and asthma—in workers in water-damaged or “moldy” buildings have been associated with the presence of high numbers of microorganisms (8). Mycobacteria were recovered from materials collected from water-damaged buildings, as well as from microorganisms normally associated with building materials (9). During reconstruction, those mycobacteria could be aerosolized in the dust. Although other microorganisms could be responsible for the respiratory problems, both saprophytic (e.g., *M. terrae*) and pathogenic (e.g., *M. avium*) strains isolated from moldy buildings were capable of inducing inflammatory responses in a mouse macrophage cell line (31). The mycobacteria elicited dose-dependent production of cytokines interleukin-6 and tumor necrosis factor-α, nitric oxide, and reactive oxygen species from the murine macrophage (31). Because whole mycobacterial cells were used in the assays (31), whether cell metabolites, which are likely easily aerosolized, were responsible for the induction of inflammatory reactions is not known. Heat-shock proteins from a number of mycobacterial species have been shown to generate Th1-type responses, airway inflammation, and airway hyperresponsiveness (32). This evidence suggests that mycobacteria or their metabolites are possible causes of respiratory disease in persons exposed to water-damaged buildings.

**Ecology of Mycobacteria**

The unique combination of physiologic characteristics that distinguish the environmental opportunistic mycobacteria make them likely agents for causing respiratory disease in these diverse settings. Mycobacteria are found in a great variety of natural and human-influenced aquatic environments, including treated drinking water (21) and aerosols (33). Mycobacteria in drinking water are associated with the presence of particulates (21). Although these microbes are grown in rich media in the laboratory, they are oligotrophic and capable of substantial growth in low concentrations of organic matter. For example, *M. avium* and *M. intracellulare* can grow in natural and drinking water over a temperature range of 10°C to 45°C (34). Mycobacteria are relatively resistant to high temperatures. For example, 10% of cells of a strain of *M. avium* survived after 1 h at 55°C (29). Mycobacteria are slow growing as a consequence of their fatty acid- and wax-rich impermeable cell wall (35). The resulting cell surface hydrophobicity permits adherence to solid substrates (e.g., pipes and leaves) in aquatic environments, which results in
mycobacteria’s persistence and resistance to being washed away at high flow rates (21,22). Further, hydrophobicity is undoubtedly associated with the ability of these bacteria to metabolize a wide variety of nonpolar organic compounds (18–20) that are constituents of metal working fluids (15,16).

Resistance of Mycobacteria to Disinfection
Mycobacteria are very resistant to the disinfectants used in water treatment, including chlorine and ozone (27). For example, M. avium is almost 500 times more resistant to chlorine than is Escherichia coli (27). Mycobacteria are also quite resistant to agents used for surface and instrument disinfection, including quaternary ammonium compounds, phenolics, iodophors, and glutaraldehyde (16,22,23,36) and can degrade the disinfectant morpholine (20). Hydrophobicity and impermeability are undoubtedly factors contributing to the disinfection resistance of mycobacteria (35). Chemical or enzymatic removal of surface lipid, while not reducing viability, reduces surface hydrophobicity and alters cell charge (37). Because of their inherent impermeability, mycobacteria grow relatively slowly compared to other bacteria. The slow growth is not necessarily a disadvantage because it correlates with increased resistance to antimicrobial agents (35), including chlorine (Falkinham JO, unpub data).

Exposure of a mixed microbial population to disinfectants results in selection of a disinfectant-resistant or tolerant population (38). The persistence and growth of mycobacteria in drinking water systems (21) are due, in part, to their disinfectant-resistance (27) and ability to grow under oligotrophic conditions (21). Disinfection of swimming pools, therapy pools, and spas or hot tubs with chlorine is expected to kill nonmycobacterial flora and to permit the growth of even the slowly growing mycobacteria in the absence of competitors for nutrients. High temperature would also be expected to result in enrichment of mycobacteria (29,30). Resistance to disinfectants could also lead to the proliferation of mycobacterial populations in metal working fluid and coolants after disinfection (6,12,13).

Aerosolization of Mycobacteria
Although M. tuberculosis is transmitted between patients through aerosols, little information exists on aerosolization of the environmental opportunistic mycobacteria (e.g., M. avium and M. intracellulare). Patient-to-patient transmission of environmental opportunistic mycobacteria does not occur (39). M. avium and M. intracellulare are readily aerosolized from aqueous suspension (28,33). Transfer of mycobacteria occurs as a result of binding of mycobacterial cells to air bubbles and ejection of water droplets after the air bubbles reach the liquid surface (28). Aerosolization can result in >1,000-fold increase in numbers of viable mycobacterial cells per milliliter of water droplets ejected from water (28). Mycobacteria in natural aerosols are found in particles and droplets (i.e., <5 µm) that can enter the alveoli of the human lung (28,33). Cell surface hydrophobicity, not surface charge, is a major determinant of enrichment in ejected droplets (28). Transfer of mycobacteria from water to air is subject to prevailing physiochemical conditions and can be manipulated. Salts (e.g., NaCl) or detergents reduce the rate of transfer of mycobacteria from water to air by ejected droplets (28). The influence of the components of metal-working fluid or of chlorine or other disinfectants in water upon aerosolization mycobacteria is unknown.

Mycobacteria and Immune Responses and Airway Inflammation
Mycobacterial cells and cellular components provoke inflammatory responses. Cells of mycobacterial strains isolated from material collected from water-damaged buildings provoke inflammatory responses in macrophages (31). Mycobacterial heat-shock proteins generate Th1-type responses, airway inflammation, and hyper-responsiveness (32). The mycolic acid-containing glycolipids, mannose-containing phospholipids, glycopeptidolipid mycosides, phenolglycolipid mycosides, and sulfatides that are unique to mycobacteria have all been reported to stimulate immune responses in animals (40). Further, mycobacteria produce a variety of extracellular primary and secondary metabolites (19) that could be aerosolized and trigger immune responses, including hypersensitivity pneumonitis. Some of these immunostimulatory compounds are produced in response to growth on polycyclic aromatic hydrocarbons (18). Unfortunately, the studies of inflammatory responses provoked by mycobacteria have been limited to whole cells grown under a single condition (31) or single proteins (32). The influence of growth conditions (e.g., growth in metalworking fluid or chlorinated water) or cell fractions (e.g., membranes) or metabolites to stimulate inflammatory responses has not been measured.

Conclusion
Contemporary reviews of airway dysfunction all describe the need for information concerning microbial agents of workplace and household exposure (41). Although many more studies are needed, the evidence points to a role of environmental opportunistic mycobacteria in provoking hypersensitivity pneumonitis, respiratory disease, and respiratory infection in both the workplace and home. In addition to the recovery of identical species and types of mycobacteria from reservoirs and patients, physiologic characteristics of mycobacteria are consistent
with their presence in the sources, transmission by means of aerosols, and illnesses. Identifying the factors that influence the presence of mycobacteria in aerosols in these workplaces would have an impact on workers in a variety of occupational settings.

On the basis of several physiologic and ecologic characteristics of mycobacteria, several approaches to reduce the impact of mycobacteria in these settings are possible. Because mycobacteria are associated with particulates (21), their numbers in reservoirs can be reduced by removal of particular matter (e.g., filtration). UV light can be used to reduce mycobacterial numbers. Disinfection of mycobacteria at high temperatures (e.g., 40°C) is more effective at reducing numbers, especially if cells were grown at lower temperatures (e.g., 30°C). Agents or combinations with surfactant or detergent-like and disinfectant activity would increase permeation in cells and biofilms and kill more mycobacteria. Finally, aerosolized or waterborne mycobacteria may be trapped in filters coated with hydrophobic compounds (e.g., paraffin) and thereby intercepted before inhalation or ingestion.

Dr. Falkinham is a professor of microbiology in the Department of Biology at Virginia Polytechnic Institute and State University. His research interests include identifying the genes and physiologic characteristics of Mycobacterium avium that are responsible for its ecology, transmission, and virulence.

References


Address for correspondence: J.O. Falkinham, Fralin Biotechnology Center, West Campus Drive, Virginia Tech, Blacksburg, VA 24061-0346, USA; fax: (540) 231-7126; email:jofiii@vt.edu