Letter to the Editor

*Mycobacterium riyadhense* overlooked: we can only find what we are looking for

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In 2009, we identified a novel mycobacterial species, *Mycobacterium riyadhense*, which was isolated from a patient in Riyadh, Saudi Arabia [1]. In 2012, there were two new reports on three additional cases in Korea, Bahrain and France, caused by the same bacteria, thus showing its emerging potential worldwide [2-3]. In these three cases, *M. riyadhense* was isolated from broncho-alveolar lavage and sputum, and associated with disease. All of the reported patients received regular anti-tuberculosis treatment until the causative agent was properly identified; for some, these results became available one year after presentation [3].

*M. riyadhense* is a slow-growing *Mycobacterium* which produces rough, white colonies after four weeks of incubation at 36°C. Biochemical properties include (i) nitrate reduction, Tween 80 hydrolysis, arylsulfatase, and urease activities; (ii) absence of niacin accumulation, heat stable catalase and tellurite reduction; (iii) no growth on MacConkey agar; (iv) negative tolerance to *p*-nitrobenzoic acid, hydroxylamine and oleic acid; and (v) tolerance to thiophene-2-carboxylic hydrazide, thioacetone and isoniazid.

Phylogenetically, *M. riyadhense* is very close to *M. szulcga* and contains the region of difference-1 (RD1) with virulent factors (*esat-6* and *cfp-10* genes), also present in *M. tuberculosis*. The 16S rRNA, *hsp65* and *rpoB* genes and the 16S-23S internal transcribed spacer sequences are unique [1]. The emerging reports on diseases mimicking tuberculosis (TB), but caused by *M. riyadhense*, emphasize the need for proper identification of mycobacteria in the diagnosis of mycobacterial diseases. It is conceivable that nontuberculous mycobacterial (NTM) diseases are more frequent than previously assumed and misdiagnosed as TB. This is especially a problem for bacteria such as *M. riyadhense* with its ability to cause pulmonary and extra pulmonary, TB-like disease in apparently immunocompetent individuals and which are actually identified as *M. tuberculosis* by some popular commercial identification assays such as GenoType MTBC/CM [1,3-4].

Proper distinction of TB and NTM disease is of the utmost importance, because this has profound consequences for treatment. Standardized TB treatment is insufficient for NTM, including *M. riyadhense*; currently reported cases followed at least nine months of first-line drug therapy to attain a cure [1-3]. The problem with the identification of newly recognized mycobacteria such as *M. riyadhense* is that there are no commercially available kits to facilitate a timely and accurate diagnosis; only DNA sequencing will offer definitive identification. The currently available identification assays focus on the most frequently isolated species and lack the flexibility to adjust to new and emerging species. If *M. riyadhense* infections are becoming more associated with serious disease, it would be important to include testing for these bacteria in commercial identification kits. Only frequent testing for *M. riyadhense* will provide an accurate assessment of its true prevalence. This
species may be of specific interest because it harbours an ESX-1 system, a well-known virulence factor of *M. tuberculosis* that is also functional in NTM, at least in *M. kansasii* and *M. szulgai* [5].

More cases of TB than currently assumed may actually represent NTM disease, caused by species such as *M. riyadhense*; therefore, more sophisticated identification tools, including DNA sequencing and the inclusion of newly emerging species into commercial identification kits, are imperative research goals.

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**References**


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