The Role of Molecular Methods in Identification of Nocardiosis

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Dear Editor,

The genus *Nocardia* one of the important pathogenic group of aerobic *actinomycetes*, which includes high rate of GC, presence of mycolic acid (60-40 carbons), and the type IV of peptidoglycan (consisting of meso-diaminopimelic acid, arabinose and galactose) in the cell wall structure (1-2). This group of bacteria was introduced by Edmond Nocard in 1888 from bovine farcy; the term "Nocardia" was established by Trevisan in 1889, and the first report of human infection of nocardiosis was described by Epinger in 1890 (1-3).

*Nocardia* spp. are gram positive, branching, relatively slow-growing and partially acid-fast bacteria, which can enter into the human body using inhalation of aerosol or traumatic inclusion causing various infections. Nocardiosis that can be the localized cutaneous infection to pulmonary, brain abscess, keratitis, as well as the disseminated and catheter-related infection in immune-compromised patients and healthy individuals. According to the literatures, the infection rate of nocardiosis in Iran is estimated to be 1.88% (2-4). The selection of appropriate *Nocardia* isolation method is one of the important steps for diagnosing and the treatment of nocardiosis. Isolation of *Nocardia* species from clinical specimens are difficult, as *Nocardia* spp. have slow growth (between 7-30 days), are sensitive to decontaminating materials, and misidentified as fungi and contamination in medical laboratories. The paraffin baiting technique is the reliable and suitable method for the diagnosis of nocardiosis infections, which is popular and frequently used by various researchers (1-2,5-6). This method (the paraffin baiting method) was introduced by Mishra et al., which includesthe carbon free media, such as McClung carbon-free broth and a paraffin coated rod glass. Unlike the other bacteria, genus *Nocardia* produced enzymes, which metabolized the paraffin, and the colonies appear between 1-4 weeks (7).

*Nocardia* species are identified and classified using conventional methods, such as the colony morphology, growth rate, Kinyoun staining, aerial hyphae, growth to lysozyme broth, growth in different temperatures, pigment production, hydrolysis of tyrosine, xanthine, hypoxanthine, casein, esculin, urease production, consumption of citrate, gelatinase activity, nitrate reductase, and utilization of various carbohydrates as the sole source of carbon. These methods are time-consuming, labor-intensive, expensive, need skilled microbiologists, and are difficult to be standardized, and confusing. Moreover, the presence of complexes such as *Nocardia asteroides* complex (N. asteroides, *Nocardia cyriacigeorgica*, N. farcinica), *Nocardia nova* complex (Nocardia nova, Nocardiaveterea, Nocardia-africana, Nocardiakruczakiae), *Nocardiatransvalensis* complex (*Nocardiatransvalensis*, *Nocardiablacklockiae*, *Nocardia wallacei*) in the genus *Nocardia* provided extensive difficulties. However, molecular tests are inexpensive, accurate, reliable, fast in emergency cases, being appropriate identifying methods, which include sequencing, polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) and DNA hybridization technique using housekeeping genes such as 16S rRNA, ITS (16S-23S rRNA internal transcribed spacer), *rpoB*, *hsps6*, *secA*, *gyrB* and *sodA* genes (1-2). Among the mentioned methods, 16S rRNA direct-sequencing is the gold standard method for identification and differentiation of nocardial species, especially in emerging *Nocardia* spp. 16S rRNA has made from hyper-variable and conserved regions that according to Laurent et al., *Nocardia* spp. can easily be identified using a 596-bp region in 16S rRNA gene, which can be amplified by NG1 (5′-ACGACCAAGGGG-3′) and NG2 (5′-GTGTGTAACCTTTCGA-3′) paired primers. However, some reports have proved otherwise; e.g., the presence of two different copies of 16S rRNA in *N. nova*, *Nocardiaconcava*, *Nocardiaignorata* or *Nocardiaamanashiensis*, and similarities between closely related species of *N. nova* complex and diversity in the 16S rRNA gene sequence of *N. cyriacigeorgica*can be observed (1-2,8). Thus, it is better to survey and study the two or more housekeeping genes using MLSA (Multi-locus sequence analysis) technique (9). In addition, the...
REA (Restriction Endonuclease Analysis) method was also an inexpensive and rapid method for identification of Nocardia spp., achieved via restriction of the enzymes that cleavage the unique palindromic sequences; e.g. hsps65 sequence via BstII, HinfI, HaeIII, BsaFI and MspI or 16S rRNA using BstII, DpnII, HindIII, Hhal, NdeI, SphI and HinPII restricted the enzymes and subsequently compared with the available reported REA patterns (1-2,10).

Nowadays, the prevalence of nocardiosis infection is increasing, and this increase can be due to enhancing the number of patients affected to immune system disorders and also improving the diagnosing methods, especially the development of molecular methods (11-12).

The genus Nocardia has six drug-susceptibility patterns consisting of N. abscessus as type I, Nocardia brevicatenata-paucivorans complex as type II, N. nova complex as type III, N. transvalensis complex as type IV, N. farcinica type V and N. cyriacigeorgicae type VI; but due to spreading of antibiotic-resistant gene elements among community of bacteria and emerging of novel Nocardia spp., the drug susceptibility patterns did not cover all the species (1-2).

In brief, the identification and differentiation of Nocardia species are essential for the final diagnosis, patient management, appropriate treatment, and epidemiological studies. Moreover, this group of bacteria can be efficient in the biodegradation process and production of secondary metabolites such as antibiotic. Molecular methods are the best options for rapid and accurate identification and differentiation of Nocardia species.

**Conflicts of interest**

None to declare.

**References**


