Case Report

Cutaneous Mycobacterium Gordonae Infection in A Patient with Myelodysplastic Syndrome

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Introduction
A 78 year-old Caucasian male patient, presented to the Dermatology Unit of San Marco Hospital in Catania following the appearance for about one year of both erythematous-papular lesions tending to confluence in plaque and nodular ones were observed, in correspondence of the back of the hands and fingers and on the left ear helix. These lesions showed red-purple colour, scaling sharp slightly raised borders, and dimensions between 0.5 and 5 cm diameter; the largest ones also showed central resolution and atrophy.
Figure 1: In correspondence of the back of the hands and fingers of a 78-year old male patient, we observe erythematous-papular lesions tending to confluence in plaque and nodular lesions of red-purple colour, scaling sharply slightly raised borders, and dimensions between 0.5 and 5 cm diameter; the largest plaques also showed central resolution and atrophy.

During the previous months he had undergone topical steroid and systemic antibiotic therapy, without any benefit. His medical history was notable for myelodysplastic syndrome.

2. Methods
A cutaneous 5 mm punch biopsy of a nodular lesion from the back of the left hand and routine blood chemistry tests were performed. Histological exam of the hematoxylin and eosin stain revealed hyperkeratotic hyperplastic epidermis with scaly-crust and foci of spongiosis, angiectasias in the dermis, intense inflammatory lymphoplasmacytic and abscessualizing granulocytic infiltrate, with plurinuclear giant cells and foci of collagen necrosis. Outbreaks of granulocyte exocytosis on the follicular epithelium and epidermis were also noticed. In conclusion, the histological finding was compatible with the diagnosis of abscessualizing granulomatous dermatitis with foci of collagen necrosis.
Figure 2: H&E stained longitudinal section obtained from a 5 mm punch biopsy of a nodular lesion from the back of the left hand: hyperkeratotic hyperplastic epidermis with scaly-crust and foci of spongiosis, angiectasias in the dermis, intense inflammatory lymphoplasmacytic and abscessualizing granulocytic, with plurinuclear giant cells and foci of collagen necrosis, outbreaks of granulocyte exocytosis on the follicular epithelium and epidermis.

Anemia, leukopenia and thrombocytopenia in accordance with the underlying hematological disorder were detected. Liver enzymes were normal (GOT 12 U/I and GPT 10 U/I) and creatinine values were 1,03 mg/dl. An HIV test was performed, and the result was negative. Chest X-ray showed a mild interstitial reinforcement of the bilateral ilo-paracardiac broncho-vasal weft, a normal cardiac shadow and free cost-phrenic breasts.

3. Results
Biochemical analysis of tissue samples, performed through the extraction of the genomic material with the QIAamp DSP DNA FFPE Tissue kit, the amplification of the genomic material with the Anuplex plus method MTB / NTM Real-Time PCR detection, and the genotyping of mycobacteria with the Genotype Mycobacterium CM VER 2.0 test, highlighted the presence of M. gordonae. Then, in the light of clinical, laboratory and instrumental data, the patient was diagnosed with M. gordonae infection.

After diagnosis, the patient was treated with a combined therapy of Clarithromycin (500 mg PO daily), Ethambutol (1200 mg PO daily), Rifabutin (600 mg PO daily), Fusidic acid + betamethasone valerate cream (twice daily). The skin lesions were completely resolved after 6 months of treatment.
Mycobacterium gordonae, isolated from sputum of patients, is a rare Nontubercular Mycobacterium named after Ruth E. Gordon, who isolated it in 1962. In the past, it was simply considered a non-pathogenic water contaminant [1, 2], while nowadays it is commonly associated with environmental contamination of organic samples. In fact, M. gordonae has been isolated from gastric fluid, urine and mucous membrane swabs collected from healthy patients [3]. Over the years, it has been found that Mycobacterium is responsible for cutaneous and non-cutaneous infections in both immunocompromised and immunocompetent patients [4].

Furthermore, the diagnosis has been certified from the patient’s testimony that affirmed to have had the first cutaneous events after working in the fields. As the literature shows, it has been hypothesized that the lesion on the helix was due to autoinoculation, as the patient was accustomed to touch his ear very often. To the best of our knowledge, the case reported is the third documented case of cutaneous infection from M. gordonae in Italy [5, 6].

4. Conclusions
In conclusion, the infection with M. gordonae can be particularly difficult to diagnose, as it is first of all a very rare disease and furthermore, in-depth and very detailed histological and microbiological investigations are necessary; finally long periods are needed for growth and identification of the isolates.

Conflicts of Interest
The authors certify that there is no conflict of interest with any financial organization regarding the material discussed in the manuscript.

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Authors’ Contributions
All authors have participated to drafting the manuscript, Prof. Rocco De Pasquale revised it critically. All authors read and approved the final version of the manuscript.

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