Daily Moxifloxacin, Clarithromycin, Minocycline, and Clofazimine in Nonresponsiveness Leprosy Cases to Recommended Treatment Regimen

Francisco Bezerra de Almeida Neto, MD1,2,3*, Rebeca Daniele Buarque Feitosa, OT3 and Marqueline Soares da Silva, RN3

1Department of Dermatology, Mauricio de Nassau Recife University Center, Brazil
2Department of Tropical Medicine, Federal University of Pernambuco (UFPE), Brazil
3Cabo de Santo Agostinho Health Care Hansen’s Disease Specialized Center, Cabo de Santo Agostinho, Brazil

*Corresponding author: Francisco Bezerra de Almeida Neto, MD, Department of Dermatology, Mauricio de Nassau Recife University Center; Department of Tropical Medicine, Federal University of Pernambuco (UFPE); Cabo de Santo Agostinho Health Care Hansen’s Disease Specialized Center, Cabo de Santo Agostinho, R Jonathas de Vasconcelos, 316, Boa Viagem, Recife, PE, CEP 51021140, Brazil, Tel: +5581988484442

Abstract

Background: In hyperendemic countries for leprosy, there has been a growing increase in clinical multibacillary “nonresponsiveness” leprosy cases to the fixed-duration treatment recommended by World Health Organization (MDT-MB). There are no defined protocols to treat these patients.

Methods: A retrospective, observational case series study was conducted of 4 patients with multibacillary leprosy who presented to a specialized Leprosy health care. Between July 1, 2018, and July 31, 2020, they were treated with moxifloxacin, clarithromycin, minocycline, and clofazimine. A literature review was conducted on the clinical features and treatment of nonresponsiveness leprosy cases to regular MDT-MB.

Results: Four patients (3 male; mean [SD] age at treatment, 41 [22.6] years) with nonresponsiveness lepromatous leprosy to regular MDT-MB were successfully treated with daily moxifloxacin, clarithromycin, minocycline, and clofazimine for 24 months. There were reduction in the number and size of skin lesions and infiltrations, neural thickening (both visually and by palpation), bacteriological index, and leprosy reactions at 6 to 24 months after initiation of therapy.

Conclusions: This case series suggests that daily moxifloxacin, clarithromycin, minocycline and clofazimine association may hold promise as a treatment for refractory lepromatous leprosy to regular MDT-MB.

Keywords

Leprosy, Lepromatous Leprosy, Drug resistance

Introduction

Although leprosy is the first infectious disease attributed to a pathogen by Gerard Amäuer Hansen, it remains a relevant public health problem in countries considered hyper-endemic for the disease [1].

The main etiologic agents are Mycobacterium leprae and Mycobacterium lepratosis. The clinical manifestations of the disease are diverse, according to the pattern of genetically predetermined immune response [2,3].

The WHO recommended treatment (MDT-MB-12 doses) has contributed to the reduction of the prevalence coefficient of the disease over the years. However, there is an increase trend of pediatric cases with grade 2 disability, polar lepromatous (LL), and other multibacillary cases. LL or subpolar lepromatous leprosy patients (BL) remain a challenge for specialists who accompany them in specialized care centers [4].

There is no consensus in the literature to assess criteria for a cure in patients after regular MDT-MB. Patients with high initial bacteriological index (BI), who remain without improvement after the completion of treatment, should extend it by an additional 12 doses, according national guidelines in Brazil. This situation is
considered "therapeutic insufficiency". If after 24 doses of regular MDT-MB, patients remain without improvement, these cases are called "therapeutic failure" and drug resistance test (DR) should be performed. There is no recommendation on treatment after the DR test result [5].

Therefore, no well-established, effective cure criteria and treatment protocols exist for nonresponsiveness leprosy cases (NR-Lep). The main ways of assessing the clinical improvement of these patients are the appearance of the lesions at the end of treatment and the decline in BI in log-units per year, determined previously by internationally used Ridley’s logarithmic scale [6].

Antibody responses to specific Mycobacterium leprae antigens can be evaluated by levels of immunoglobulin M (IgM) against phenolic glycolipid-1 (PGL-1). The presence of high titles of anti-PGL-1 IgM reflects the total bacterial load in the body, however, test anti-PGL-1 are not widely available in field, being restricted to research institutions [7,8].

In fact, there is no consensus as to what the actual patients’ cure means. The established cure happens only because these patients complete the period recommended for treatment with MDT-MB, which is insufficient in several cases of relapse of the disease mainly in lepromatous leprosy [9].

Moxifloxacin, clarithromycin, minocycline, and clofazimine (MCMC) used for the treatment of this case series patients are mentioned in the WHO guidelines, still without retrieved evidence, based on the opinion of experts [10].

Methods

We conducted a retrospective case series of 4 patients with NR-lep presenting to Municipal Cabo de Santo Agostinho Health Care Hansen’s Disease Specialized Center (Cabo de Santo Agostinho - Pernambuco - Brazil) between July 1, 2018, and July 31, 2020. This project was not human participants’ research, and the Brazilian Ministry of Health received a report on the clinical situation and treatment of this case series. All patients provided written consent.

Four sites provided the collection of skin smears by an experienced laboratory technician: The two ear lobes, right elbow, and an active lesion, according to the protocol in Brazil. Slides were stained using Ziehl Neelsen Carbol Fushin and scored for BI, determined for an exclusive Biomedical professional specializing in mycobacteria using international Ridley’s logarithmic scale. The laboratory has a regular quality control service both for the technicians and for the field. The Oswaldo Cruz Institute (Fiocruz), Rio de Janeiro - RJ, Brazil performed the molecular biology tests for drug resistance.

Case Series

Patient 1 was a female patient, 29-years-old, presenting infiltrations in earlobes and external ears and multiple fibromatous lesions scattered over the skin, predominantly in the abdomen and lower limbs. In addition, there was involvement of the ulnar and posterior tibial nerves. She also had corneal anesthesia on the right. The diagnostic of LL was in May 2013. BI was 5.0. She started MDT-MB. After the first month of treatment, she developed intolerance to dapsone, and ofloxacin 400 mg/day replaced this drug, as recommended by the Brazilian Ministry of Health. After 24 doses of treatment, there was no clinical improvement, showing the lesions with the same initial appearance. There was infiltration of the earlobes and external ears in the same way and BI remained 5.0. Drug resistance test was negative. Additional twelve months extended treatment to 36th month. She concludes it and remained without clinical improvement, with the BI = 5.0 (Figure 1A and Figure 1B). Subsequent daily MCMC therapy led to improvement of infiltrations 6 months after treatment (Figure 2A and Figure 2B). At the 24-month follow-up, the BI is 0.75. Patient’s skin is free of initial lesions. During the entire follow-up period, this patient developed erythema nodosum leprosum (ENL). The first episode was at the fourth month of treatment and continued after the MCMC treatment.

Patient 2 was male, 30-years-old, diagnosed with LL...
MB were prescribed. The BI at the end of the 36th dose remains 4.25 and the patient continued to develop LR (Figure 1C). After starting with daily MCMC treatment, LR ceased at the sixth month of treatment. There was a significant improvement in dermatological lesions (Figure 2C). The final BI was 0.

Patient 4 was a 26-year-old man, diagnosed with LL in March 2013. He had infiltration of auricular pavilions and some cutaneous areas, in addition to thickening and palpation pain in ulnar, common fibular and posterior tibial nerves. Initial BI was 4.75. He performed 24 doses of regular MDT-MB. From the first month of treatment, he developed several episodes of ENL with associated neuritis, two of which are more severe and in need of hospital intervention. At the end of treatment, BI was 4.50, and drug resistance tests were negative. The treatment also was extended with an additional 12 months with regular MDT, with the BI at 36th dose = 4.75. Subsequent daily MCMC therapy led to improvement of infiltrations 1 year after treatment. He remained with less intense LR throughout the period and continued after the MCMC treatment. The final BI was 0.25.

All patients were treated for leprosy reactions with corticosteroids (prednisone, dexamethasone, and methylprednisolone), with dosages ranging from 20 to 60 mg/day and chosen according to the intensity and duration of the episode. Besides, all of them also used thalidomide, with different dosages, between 100 to 400 mg/day. Patient 4 required the addition of pentoxifylline, as leprosy reactions were not controlled with the maximum doses of corticosteroids and thalidomide allowed. Amitriptyline (50 mg/day to 75 mg/day) was an adjuvant for the control of neuropathic pain in all patients. The use of 100 mg/day of acetylsalicylic acid (ASA) minimized the risk of thromboembolic events. There were no predisposing factors for the occurrence of leprosy reactions (Table 1).

Discussion

This case series documents the efficacy of 24 months daily MCMC in the treatment of NR-lep in a small cohort.

Skin lesions, neural involvement and leprosy reactions episodes remained unchanged in all patients, after completing 36 doses of MDT-MB treatment. No patient achieved clinical improvement after the end of 36 doses of regular MDT-MB.

BI declined very little and below expectations in all cases. After completing the 24 doses of MDT-MB treatment, all patients underwent drug resistance testing. Due to the negative results, they have same treatment extended by 12 more doses, since the patients did not improve.

The BI decline after MDT-MB treatment averages 0.5 to 1 unit per year. At the end of the 36th month, the pa-

Figure 2: Patient 1 and patient 3 in the 6th month of MCMC treatment (A) Right earlobe patient 1; (B) Left earlobe patient 1; (C) Dorsal region patient 3.
| Patient No./Sex/Age (y) | LCP | Affected nerves/R or L or both | Associated diseases | Initial BI | Final BI After 24 regular-MDT-MB | Final BI After 36 regular-MDT-MB | Final BI after 24 months MCMC | Leprosy reaction before MCMC | Leprosy reaction types/Treatments | MCMC: drug/daily dosage | Side-Effects/Laboratorial Alterations’/Drug Resistance-test |
|------------------------|-----|--------------------------------|---------------------|------------|---------------------------------|---------------------------------|-------------------------------|-----------------------------|---------------------------------|-------------------------------|-----------------------------|
| 1/F/29                 | LL  | Ulnar/both posterior tibial/both | Systemic arterial hypertension | 5.0        | 5.0                             | 5.0                             | 0.75                          | yes                         | yes                             | Type 2 with neuritis/Prednisone, dexamethasone thalidomide, ASA, amitriptilin | None/none/negative          |
| 2/M30                  | LL  | Great auricular/L Ulnar/R       | None                | 4.75       | 4.25                            | 4.50                            | 0                             | yes                         | no                              | Type 2 with neuritis/Prednisone, thalidomide, pentoxifilin, ASA, amitriptilin | None/none/negative          |
| 3/M/67                 | BL  | Ulnar/both Posterior tibial/both | Systemic arterial hypertension and diabetes mellitus | 4.50       | 4.25                            | 4.25                            | 0                             | yes                         | no                              | Types 1 and 2 with neuritis/ Prednisone, thalidomide, ASA, amitriptilin | None/none/negative          |
| 4/M/26                 | LL  | Ulnar/both Posterior tibial/both Common fibular/both | Systemic arterial hypertension | 4.75       | 4.50                            | 4.75                            | 0.25                          | yes                         | yes                             | Type 2 with neuritis/Prednisone, dexamethasone, metil-prednisolone, methotrexate, thalidomide, pentoxifilin, ASA, amitriptilin | Vertigo, dizziness/none/negative |

**Abbreviations:** LCP: Leprosy Clinical Presentation; LL: Lepromatous Leprosy; BL: Borderline-Lepromatous Leprosy; R: Right; L: Left; BI: Bacteriological Index; MDM: Multibacillary Multidrug Therapy; MCMC: Moxifloxacin, Clarithromycin, Minocycline and Clofazimine; ASA: Acetylsalicylic Acid; *: Laboratory alterations are the follow tests performed monthly: blood count, blood glucose, urea, creatinine, triglycerides, total cholesterol, alanine aminotransferase, aspartate aminotransferase, lactic dehydrogenase, amylose and urinalysis.
tients also remained without improvement or changes in the BI, except for patient 4, who increased the BI to the initial level.

It is important to note that all patients were tested for drug resistance and were negative. For this reason, patients remained on regular treatment with MDT-MB until the 36th dose.

Current tests for drug resistance in leprosy target the genes of the drug resistance determinant region (DRDR) dihydropteroate synthase (Folp1), DNA-directed RNA polymerase subunit beta (RpoB), and DNA gyrase subunit A (Gyr-A). However, there are mutations identified in other regions, DRDR’s outside, such as bifunctional enzyme riboflavin biosynthesis protein (ribD), DNA-directed RNA polymerase subunit alpha (rpoA), DNA-directed RNA polymerase-beta chain (RpoC) and nonsense mutations in the nth excision repair gene. Single-nucleotide polymorphisms (SNPs) were associated with drug resistance. Besides that, genomic deletion mechanisms in Mycobacterium leprae-variable number tandem repeat (VNTR), called TTC repeats are described and might lead to microbiological nonresponse in treated cases of leprosy, which add a chapter to Mycobacterium leprae [11,12].

Daily MCMC treatment was well tolerated. All patients were monitored in the laboratory, with no changes in blood count, liver, kidney, and pancreatic function. One patient complained of dizziness and vertigo within the first month of treatment, which subsequently disappeared.

With the MCMC treatment, all patients improved skin lesions, infiltrations, and reduced neural thickening, with improved function of the affected nerves (both visually and by palpation). There was a rapid reduction in BI, higher than that usually described with the MDT-MB treatment [13].

In addition to the bactericidal effect, all drugs used have recognized anti-inflammatory activity. This fact may have contributed to the control of LR.

Moxifloxacin has important immunomodulatory properties, such as the inhibition of TNF-α and IL-6, which are involved in the pathogenesis of the LR, especially type 2, contributing to homeostasis in LL patients [14,15].

Clarithromycin is a macrolide with known anti-inflammatory activity, with immunomodulatory effects on inflammatory cells, fibroblasts, and epithelial cells, in addition to modulating the expression of cytokines and chemokines [16].

Small trials have already tested the efficacy of moxifloxacin and clarithromycin in LL patients, both used separately and for short periods [17,18].

Minocycline is a second-line drug in the treatment of leprosy. A robust trial demonstrated the effectiveness and tolerability for minocycline and clarithromycin long-term use in this disease. This drug is usefulness in cases of drug resistance also in the control of LR due to its immunomodulatory properties [19,20].

Clofazimine is a first-line drug in the treatment of leprosy, which has the advantage of not inducing drug resistance known through molecular biology techniques, except in a few studies showing resistance in inoculation in mice. It is a option to control LR, especially type 2 [21,22].

Interestingly, two patients who remained with a positive smear after MCMC treatment continued to express type 2 LR with the need to use drugs against leprosy reactions. These patients may have a longer period of disease evolution, between the beginning of the first symptoms and the beginning of treatment. A delay of fewer than 3 years between initial symptoms of leprosy and the start of treatment may lead to a significantly faster clearance of bacilli from the body [6].

It is also possible that these two patients are not yet cured. The relapse rate is not low and can increased to patients treated with MDT-MB 24 month fixed-dose in contrast to patients treated until smear-negative [21,23].

Type 1 LR has always been understood as a purely immunological phenomenon, however, its occurrence has already been demonstrated in the presence of bacillary multiplication due to the detection of Mycobacterium leprae-antigen 85 (Ag85) in lesions of type 1 LR. Ag85 is a 30 kDA mycolyl transferase secreted during growth and multiplication of Mycobacterium leprae [24].

On the other hand, ENL after the removal of MDT-MB is associated with disease’ relapse [25,26].

All the patients’ household contacts of this case series are healthy, ruling out the possibility of reinfection.

Limitations
This study has some limitations. It was a small, retrospective case series at a single institution, and there are no validated tools to measure treatment response in NR-Lep.

Conclusions
A subset of leprosy patients with high initial BI remains a serious public health problem in countries that are hyperendemic for leprosy. They may not respond satisfactorily to MDT-MB 12-24-month fixed-dose. These patients have genetically determined energy contributing to the perpetuation of the transmission chain. We present 4 patients with NR-Lep who have been successfully treated with the daily MCMC scheme presented in this case series. Large studies are needed to understand the unsatisfactory response to MDT-MB in order to make better therapeutic alternatives available, especially considering that there are already bet-
other drugs that can be used in this neglected disease. However, the long time required for large prospective cohorts can greatly influence the quality of life of these patients.

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**Authors' Contribution**

All authors contributed equally to the elaboration of this manuscript.

**References**

1. GETZ B (1958) Leprosy research in Norway. 1850-1900. Med Hist 2: 65-67.
2. Han XY, Aung FM, Choon SE, Werner B (2014) Analysis of the leprosy agents Mycobacterium leprae and Mycobacterium lepromatosis in four countries. Am J Clin Pathol 142: 524-532.
3. Schurr E, Alcaïs A, de Léséleuc L, Abel L (2006) Genetic predisposition to leprosy: A major gene reveals novel pathways of immunity to Mycobacterium leprae. Semin Immunol 18: 404-410.
4. Mushtaq S, Dogra N, Dogra D, Faizi N (2020) Trends and patterns of leprosy over a decade in a tertiary care hospital in Northern India: A retrospective analysis. Indian J Dermatol Venereol Leprol 86: 141-149.
5. MoHo (2016) Guidelines for Surveillance, Care and Elimination of Leprosy as a Public Health problem: Technical and operational manual. In: Diseases. SoHS-DoSoI, BVS - Biblioteca Virtual em Saúde, Brazil 1-51.
6. Amenu A, Saunderson P, Desta K, Byass P (2000) The pattern of decline in bacillary index after 2 years of WHO recommended multiple drug therapy: The AMFES cohort. Lepr Rev 71: 332-337.
7. Nicchio MVC, Araujo S, Martins LC, Pinheiro AV, Pereira DC, et al. (2016) Spatial and temporal epidemiology of Mycobacterium leprae infection among leprosy patients and household contacts of an endemic region in Southeast Brazil. Acta Trop 163: 38-45.
8. da Conceição Oliveira Coelho Fabri A, Carvalho APM, Araujo S, Goulart LR, de Mattos AMM, et al. (2015) Antigen-specific assessment of the immunological status of various groups in a leprosy endemic region. BMC Infect Dis 15: 218.
9. Shen J, Yan L, Sun P (2015) Clinical features of relapse after multidrug therapy for leprosy in China. Lepr Rev 86: 165-169.
10. (2018) Guidelines for the diagnosis, treatment and prevention of leprosy. World Health Organization, Regional Office for South-East Asia, New Delhi.
11. Benjak A, Avanzi C, Singh P, Loiseau C, Girma S, et al. (2018) Phylogenomics and antimicrobial resistance of the leprosy bacillus Mycobacterium leprae. Nat Commun 9: 352.
12. Reja AHH, De A, Patra PK, Biswas S, Duttagupta U, et al. (2018) Genomic reduction at TTC repeats in the bacterial genome of treated cases of Hansen's Disease: A possible survival mechanism of Mycobacterium leprae. Indian J Dermatol 63: 449-454.
13. Jesudasan K, Vijayakumaran P, Manimozhi N, Rao PS, Samuel P (1996) Effectiveness of MDT in multibacillary leprosy. Int J Lepr Other Mycobact Dis 64: 128-132.
14. Dalhoff A, Shalit I (2003) Immunomodulatory effects of quinolones. Lancet Infect Dis 3: 359-371.
15. Sales-Marques C, Cardoso CC, Alvarado-Arnez LE, Illaramendi X, Sales AM, et al. (2017) Genetic polymorphisms of the IL6 and NOD2 genes are risk factors for inflammatory reactions in leprosy. PLoS Negl Trop Dis 11: e0005754.
16. Arikata M, Itoh Y, Shichinohe S, Nakayama M, Ishigaki H, et al. (2019) Efficacy of clarithromycin against H5N1 and H7N9 avian influenza in cynomolgus monkeys. Antiviral Research 171: 104591.
17. Pardillo FEF, Burgos J, Fajardo TT, Dela Cruz E, Abalos RM, et al. (2008) Powerful bactericidal activity of moxifloxacin in human leprosy. Antimicrob Agents Chemother 52: 3113-3117.
18. Chan GP, Garcia-Ignacio BY, Chavez VE, Livelo JB, Jimenez CL, et al. (1994) Clinical trial of clarithromycin for lepromatous leprosy. Antimicrob Agents Chemother 38: 515-517.
19. Rea TH (2000) Trials of daily, long-term minocycline and rifampin or clarithromycin and rifampin in the treatment of borderline lepromatous and lepromatous leprosy. Int J Lepr Other Mycobact Dis 68: 129-135.
20. Narang T, Sawatkar GU, Kumaran MS, Dogra S (2015) Minocycline for recurrent and/or chronic erythema nodosum leprosum. JAMA Dermatol 151: 1026-1028.
21. Shetty VP, Wakade AV, Ghate S, PaiVV, Ganapati R, et al. (2003) Viability and drug susceptibility testing of M. leprae using mouse footpad in 37 relapse cases of leprosy. Int J Lepr Other Mycobact Dis 71: 210-217.
22. Roy K, Sil A, Das NK, Bandyopadhyay D (2015) Effectiveness and safety of clofazimine and rifampin in type 2 lepra reaction: A double-blind, randomized, controlled study. Int J Dermatol 54: 1326-1332.
23. Gelber RH, Balagon VF, Cellona RV (2004) The relapse rate in MB leprosy patients treated with 2-years of WHO-MDT is not low. Int J Lepr Other Mycobact Dis 72: 493-500.
24. Save MP, Dighe AR, Natrajan M, Shetty VP (2016) Association of viable Mycobacterium leprae with Type 1 reaction in leprosy. Lepr Rev 87: 78-92.
25. Guerrero-Guerrero MI, Muvdi-Arenas S, León-Franco CI (2008) Association between leprosy reactions after treatment and bacterial load evaluated using anti PGL-1 serology and bacilloscopy. Rev Soc Bras Med Trop 41: 67-72.