INTRODUCTION

Leishmania donovani in Sudan causes different clinical forms ranging from cutaneous ulcer (cutaneous leishmaniasis [CL]), mucosal leishmaniasis (ML), to visceral leishmaniasis (VL). Visceral leishmaniasis can be complicated by development of a skin rash (post kala-azar dermal leishmaniasis [PKDL]) during or following treatment of VL patients with pentavalent antimonials.1,2

According to the guidelines of the Federal Ministry of Health, Sudan, VL patients are treated with pentavalent antimonials as the first line of treatment, given at a dose of 20 mg/kg for 28 days. The course of treatment is extended for treatment of ML and PKDL patients. Patients, who do not respond to Antimony treatment, are treated with 10 doses of Liposomal Amphotericin B (Ambisome) given at a dose of 2–3 mg/kg every other day.

Ambisome has been shown to be effective against different clinical forms of leishmaniasis and it is strictly used as a second line of treatment according to the guidelines of the Federal Ministry of Health because of its unaffordable high cost. In addition to its effectiveness, Ambisome has a wide range of safety with minimal side effects.3 Nevertheless, infusion reactions to liposomal amphotericin B were previously reported.4,5 In this work, we report the development of allergic reactions by a PKDL and a ML Sudanese patient to Ambisome. The findings warrant future close supervision of patients to be treated with the drug.

Case Report: First Report on Ambisome-Associated Allergic Reaction in Two Sudanese Leishmaniasis Patients

Maowia Mukhtar,* Mona Aboud, Musa Kheir, Sahar Bakheet, Nazik Abdullah, Ahmed Ali, Nadia Hassan, Elwaleed Elamin, and Atif Elagib

Institute of Endemic Diseases, University of Khartoum, Khartoum, Sudan; Umdurman Tropical Disease Hospital, Khartoum, Sudan; Faculty of Medicine, University of Khartoum, Khartoum, Sudan; Faculty of Medicine, University of Khartoum, Khartoum, Sudan; Faculty of Medicine, Al Nilian University, Khartoum, Sudan; Faculty of Medicine, Al Nilian University, Khartoum, Sudan; Faculty of Medical Laboratory Sciences, AL Zait Al Azhary University, Khartoum, Sudan; Tropical Diseases Research Institute, National Center for Research, Ministry of Science and Technology, Khartoum, Sudan

Abstract. Post kala-azar dermal leishmaniasis (PKDL) and mucosal leishmaniasis (ML) are serious clinical forms of leishmaniasis caused by Leishmania donovani parasites in Sudan. Although pentavalent antimonials are used as the first line of treatment of all clinical forms of leishmaniasis, persistent PKDL and ML patients are treated with liposomal amphotericin B (Ambisome) as a second-line drug. In this work, we report the development of allergic reactions by a PKDL and a ML Sudanese patient to Ambisome. The findings warrant future close supervision of patients to be treated with the drug.

The patient was referred to our laboratory in Khartoum for further research and treatment. Skin snips were taken from the PKDL lesions for parasite culture in Novy-MacNeal-Nicolle (NNN) media and parasite growth was positive after 7 days of incubation. The patient was referred for Liposomal Amphotericin B (Ambisome; Gilead Sciences Inc., Foster City, CA) at Umdurman Tropical Diseases Hospital in Khartoum. The patient was a known asthmatic since 7 years of age, and he was not on regular medication for asthma.

At presentation to the hospital, he had no constitutional symptoms apart from the facial rash. He had an extensive maculo—a popular rash confined to the face, mainly on the forehead and around the lips (Figure 1). On examination all other systems were normal, his weight was 65 kg. He was scheduled for Ambisome treatment at a dose of 2.5 mg/kg in 5% dextrose on alternate days for 20 days.

On the fifth treatment dose the patient developed facial flushing, breathlessness, and generalized expiratory wheezes. The asthmatic attack was controlled and he was placed on a regular Salbutamol inhaler thereafter. One week later, Ambisome (Gilead Sciences Ltd., Cambridge, UK) was rechallenged under close supervision. After 10 cc of the infusion, he felt chest tightness but there were no wheezes and facial flushing. The treatment was immediately stopped and the patient was considered allergic to the drug.

Mucosal leishmaniasis patient. A 54-year-old Sudanese male had a history of living in a VL-endemic area in Eastern Sudan. The patient developed a chronic ulcer on the lower lip that persisted for more than 6 months (Figure 2). The patient was suspected of having ML. A tissue biopsy was surgically taken for parasite culture and smear preparation. The biopsy was cultured in NNN media and incubated at 24 °C. Mucosal leishmaniasis was confirmed by successful parasite growth from the biopsy. Because of the severity of the lesion and the age of the patient, he was scheduled for Ambisome treatment at a dose of 2.5 mg/kg for 4 weeks and given every other day.

During the ninth dose, the patient developed generalized rash and vesicles, generalized wheezes, and gasped for oxygen that necessitated nublized solbutamol. The patient felt chilly with rigors and complained of chest tightness. His total white blood cell count was 7,200 with 9% eosinophilia. Ambisome was immediately stopped and the patient was given 10 mg antihistamine and 100 mg hydrocortisone intravenously. The patient was

* Address correspondence to Maowia Mukhtar, Institute of Endemic Diseases, University of Khartoum, P.O. Box 11463, Khartoum, Sudan. E-mail: mmukhtar@tropmedicine.org
observed for 3 days, no further event occurred and the patient was discharged in good condition.

DISCUSSION

Post kala-azar dermal leishmaniasis and ML are two serious clinical forms caused by *Leishmania donovani* in Sudan. Because of the persistent lesions of PKDL and ML that require prolonged treatment courses with pentavalent antimony, the two patients were treated with liposomal amphotericin B (Ambisome). The first patients developed PKDL lesions following VL treatment with SSG. Zijlstra and Elhassan reported an exceptionally high incidence of PKDL among VL Sudanese patients during or following treatment of VL with SSG. The patient reported in this work received 160 doses of SSG for PKDL treatment in the peripheral hospital where no alternative treatment was available. On the basis of his past repeated SSG treatment, he was recommended for Ambisome treatment. The patient tolerated 4 doses of Ambisome, and unexpectedly, developed symptoms of allergic reaction during the fifth dose. His allergic symptoms included facial flushing, breathlessness, and generalized expiratory wheezes that prompted stopping the Ambisome treatment and use of anti-allergic drugs. The fact that the patient had a history of asthma and allergic reaction to SSG might have increased the risk for his allergic reaction to Ambisome. Unfortunately, no further treatment was provided for the patient who was discharged showing reasonable clinical improvement.

The second patient who presented with a mucosal ulcer developed an allergic reaction to Ambisome treatment, although he had no history of allergy. He tolerated 8 doses of Ambisome, and then developed a serious allergic reaction during the ninth dose accompanied by high eosinophilia suggestive of hypersensitivity to Ambisome. His allergic reaction prompted discontinuation of Ambisome treatment and use of an anti-allergic drug.

Ambisome is known to have a wide margin of safety; nevertheless, allergic and infusion reactions to the drug were previously reported. The documented allergic reaction to Ambisome during treatment of PKDL and ML warrants close supervision of patients, especially since Ambisome is currently considered to be a first-line treatment of leishmaniasis in several endemic regions.

Received September 10, 2010. Accepted for publication June 8, 2011.  
Disclaimer: The study was approved by the institute ethical committee and written consents were obtained from the two patients.

Authors’ addresses: Maowia Mukhtar, Institute of Endemic Diseases, University of Khartoum, Khartoum, Sudan, E-mail: mmukhtar@tropmedicine.org. Mona Aboud, Umdurman Tropical Diseases Hospital, Khartoum, Sudan, E-mail: monaaboud@live.com. Musa Kheir, Department of Medicine, Faculty of Medicine, University of Khartoum, Khartoum, Sudan, E-mail: kheirmusa@hotmail.com. Sahar Bakhit, Institute of Endemic Diseases, University of Khartoum, Khartoum, Sudan, E-mail: nazik.abdullah@yahoo.com. Ahmed Ali, Faculty of Medicine, Al Nilan University, Khartoum, Sudan, E-mail: draali@gmail.com. Nadia Hassan, Department of Medicine, Faculty of Medicine, Al Nilan University, Khartoum, Sudan, E-mail: nadiahassan@yahoo.com. Elwaleed Elamine, Faculty of Medical Laboratory Sciences, Al Zaim Alazhary University, Khartoum, Sudan, E-mail: wmelamin@hotmail.com. Atif Elagib, Tropical Medicine Research Institute, Ministry of Science and Technology, Khartoum, Sudan, E-mail: atifelagib@hotmail.com.

REFERENCES