EVALUATING AN ALTERNATIVE TREATMENT REGIMEN FOR VISCERAL LEISHMANIASIS IN EASTERN AFRICA: A Phase III trial to assess the safety and efficacy of miltefosine plus paromomycin

DNDi and AfriKADIA consortium partners conducted a Phase III clinical trial for visceral leishmaniasis patients in Ethiopia, Kenya, Sudan, and Uganda to assess the safety and efficacy of miltefosine plus paromomycin (MF+PM) as an alternative to the standard of care, sodium stibogluconate plus paromomycin (SSG+PM), in Eastern Africa.

The trial found that the MF+PM combination therapy was as effective as SSG+PM but with fewer injections, a shorter treatment duration, and no risk of SSG-associated cardiotoxicity. MF+PM also reduced the risk of subsequent post-kala azar dermal leishmaniasis, which is a source of ongoing VL transmission in communities. Based on the evidence that it is an equally effective but more patient-friendly treatment, MF+PM could be a future alternative therapy for patients with VL in Africa.

WHY WAS THIS CLINICAL TRIAL CONDUCTED?

Since 2010, the first-line treatment for VL in Eastern Africa has been a 17-day sodium stibogluconate plus paromomycin (SSG+PM) combination therapy. It is an improvement over the previous 30-day SSG monotherapy, but the treatment is still lengthy and painful, requiring patients to be hospitalized for the entire 17-day duration of treatment, with two daily injections per day, given one after another. More importantly, although rare, the antimonial drug SSG may present life-threatening side effects such as cardiotoxicity, hepatotoxicity, and pancreatitis.

There is an urgent need for alternative treatments that are suitable to be used in the remote areas where VL typically occurs and with children, who represent more than half of patients needing treatment, as well as the elderly, who are at highest risk of SSG-related toxicity.

Miltefosine (MF) is the only oral treatment available for VL. In India and Bangladesh, a 10-day combination of MF with paromomycin (PM) has shown cure rates at 6 months ranging from 96.9% to 98.7%, making it an attractive option to evaluate in Eastern Africa. In addition, the absence of antimonial-related toxicity, fewer required injections, and shorter treatment duration would make MF+PM a better option for patients with VL.

This trial assessed the safety and efficacy of MF+PM as an alternative to the standard of care, SSG+PM, in Eastern Africa.
remote settings with limited access to health facilities. Based on the evidence that it is an equally effective but more patient-friendly treatment, MF+PM could be a future alternative therapy for patients with VL.

**WHAT ARE THE NEXT STEPS?**

The paper reporting on the results of this Phase III trial to assess the safety and efficacy of miltefosine plus paromomycin has been accepted for publication by *Clinical Infectious Diseases* journal (see link below).

DNDi and its partners will work to ensure availability of MF+PM as a VL treatment in Eastern Africa through the Leishmaniasis East Africa Platform (LEAP). DNDi will also work with local authorities and WHO to support evidence review and updating of the VL treatment guidelines with the ultimate goal of reducing morbidity and mortality from visceral leishmaniasis in the region.

**HOW AND WHERE WAS THIS TRIAL CONDUCTED?**

The purpose of this trial was to determine the safety and efficacy of a combination regimen of oral miltefosine plus paromomycin (MF+PM) for 14 days as compared to the standard of care SSG+PM 17-day treatment for primary VL patients.

The study began in 2018, conducted by DNDi and partners from the AfriKADIA Consortium with funding from the European and Developing Countries Clinical Trials Partnership (EDCTP). There were 439 participants in total, both children and adults, from seven sites in Sudan (Doka, Um el Kher and Tabarak Allah), Kenya (Kacheliba), Ethiopia (Gondar and Abdurafi), and Uganda (Amudat).

Study participants were randomly assigned to receive one of the two treatments:

- **Paromomycin (20 mg/kg/day)** via intramuscular injection for 14 days and miltefosine (allometric dosing) twice per day taken orally for 14 days, or
- **Sodium stibogluconate (20 mg/kg/day) IV/IM and paromomycin (15 mg/kg/day)** via intramuscular injection for 17 days.

The primary efficacy endpoint was definitive cure at six months follow-up, defined as absence of clinical signs and symptoms of VL after 210 days and no rescue treatment during the trial.

**WHAT WERE THE RESULTS?**

The study showed that a MF+PM combination achieved similar cure rates at six months follow-up as the standard of care SSG+PM in adult and paediatric patients with VL in Eastern Africa (91.2% and 91.8%, respectively). MF+PM also reduced the risk of subsequent post-kala azar dermal leishmaniasis (PKDL) from 20.9% in those treated with SSG+PM to just 4.4% in Ethiopia and Sudan. PKDL is a source of ongoing VL transmission in communities.

MF+PM was well tolerated; the most common adverse event was mild vomiting related to miltefosine. MF+PM eliminates one painful daily injection and the potential life-threatening toxicity associated with the antimonial SSG, and it reduces the length of hospital stay from 17 to 14 days. Given that most patients are children, these are important benefits, making the treatment more patient-friendly than the current treatment. The main disadvantage is that MF+PM cannot be taken by pregnant women, and any women of child-bearing potential must take contraceptives during treatment and for five months after treatment.

The new MF+PM combination would provide the Eastern African region with the first non-antimony-based treatment for uncomplicated VL, with increased suitability to be used in remote settings with limited access to health facilities. Based on the evidence that it is an equally effective but more patient-friendly treatment, MF+PM could be a future alternative therapy for patients with VL.

**READ THE PUBLISHED STUDY**


**FOR MORE INFORMATION**

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Photo credits: Mercy Mumo-DNDi

**THE SCIENCE EXPLAINED**

A packet of miltefosine capsules with the trade name Impavido used to treat VL patients.