Visceral Leishmaniasis (VL) or kala-a-zar is endemic in Bangladesh. Historically VL was first described in 1824, in Jessore district of the then Bengal now Bangladesh. A gradual spread thereafter the disease embraced gradually almost the entire subcontinent. In 1903 Donovan in Madras and Leishman in London independently identified the parasite from the splenic tissues in autopsies of patients who died from VL and it was Ronald Ross who proposed the name Leishmania donovani for the newly found parasite. Since long the treatment of choice for VL, has been pentavalent antimony which was discovered a century ago. Trivalent antimony was also used with similar outcome. Of Pentavalent antimonials Urea stibamine and sodium stibogluconate (Sb) was the most widely used and reliable. But these well known drugs is gradually loosing its long time reputation due to the various side effects, its duration and method of treatment, the cumbersome procedure of prolonged hospitalization and the recent reports of drug resistance developed by the parasite in different regions. Reports have shown a staggering 65% resistance in Northern Bihar of India and are pacing up in other regions of India as well. The resistance of VL to antimony compounds has also been found in other geographical locations as well and the measure to overcome the situation by increasing the dose has been back fired by the serious side effects of the drug leading to sudden deaths.

The social reason underlined behind the development of drug resistance in Bihar India, a hyper endemic zone for kala azar prevails quite similarly in Bangladesh which includes discontinuation of treatment on part of the patient, different dosage schedules and methods of administration adapted by the physicians and presence of substandard drugs in the market without a proper monitoring system regarding the indiscriminate use and selling of the drugs in the endemic areas.

Alternate therapy with amphotericin B has shown some promising results in Bangladesh and India alike. The cure rate was >97% at doses of 0.75mg/kg for 15 infusions on alternate days. Side effects such as infusion reaction and thrombophlebitis are universal along with occasional hypokalaemia, thrombocytopenia, myocarditis and death. Nephrotoxicity is extremely rare. Though relapses might occur but patients have been successfully retreated with the same drug. The extremely high cost of the safe and efficacious Liposomal Amphotericin B has made its use impractical in the kalaazar endemic regions. Thus amphotericin B may be a promising drug either alone or in combination provided proper trials conducted in Bangladesh.
Miltefosine an effective substitute for Sb (V) is gradually making its way in the treatment formulation of Kala azar. At a dose of 100mg daily for patients weighing ≥ 25 kg and 50 mg for those <25 kg orally for 4 weeks with a cure rate of 95% and minimal side effects apart from its teratogenicity, was thought to be an affordable and acceptable solution to the ongoing problem faced with Sb(V). But unfortunately reports regarding treatment failure with Miltefosine from different parts of India and Bangladesh have become a major concern.

Paramomycin, an amino glycoside is yet to be investigated thoroughly though it showed glimpses of hope to the researchers with 93% cure rate in one study in India but unfortunately the manufacturers of the drug discontinued its production.

Ketoconazole an orally administrable anti fungal drug has also been tried in the treatment of kala azar. In one study in Bangladesh 4 out of 16 cases achieved parasitological cure and 12 cases achieved clinical cure. While another study in India 4 of 5 cases responded to the drug. The researchers from Bangladesh had at that time suggested a large scale clinical study with this drug especially in those who showed resistance to Sb (V).

Researchers like Shyam Sundar, TK Jha and CP Thakur have been working relentlessly to find out the pattern of drug resistance in Indian Kala azar and an effective drug and dosage schedule to combat the situation. Measures may be taken to develop multi drug therapy for VL treatment which may show results like we have seen in tuberculosis and leprosy.

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References: