Visceral leishmaniasis (VL), which is endemic in the Indian subcontinent (42,623 reported cases/year of which 34,918 were in India), the Mediterranean region (875 cases/year), East Africa (8,569 cases/year), and Brazil (3,481 cases/year) has undergone a revolution in chemotherapy in the last 15 years. Treatment had been with the classical agents pentavalent antimony and amphotericin B deoxycholate, but with >90% of Indian disease occurring in antimony-resistant regions, the sole effective drug in this key region of the world was amphotericin B deoxycholate. Amphotericin B deoxycholate is extraordinarily effective for Indian VL. In phase 3 studies amphotericin B deoxycholate levels, the total dose of amphotericin B that might diminish toxicity, thereby permitting larger individual doses and a shorter total treatment period. Most reports concern liposomal amphotericin B (Ambisome), but there are a few reports on amphotericin B lipid complex (ABLC/Abeclat), amphotericin B colloidal dispersion (ABCD/Amphocil), amphotericin B lipid emulsion (ABLE), and in this issue of the Journal, an Indian-formulated liposomal amphotericin B (Fungisome).

Ambisome: Although adverse reactions to Ambisome are qualitatively similar to those of amphotericin B deoxycholate, their frequency and severity are diminished.6 Improved tolerance led to higher daily doses, a trial of 7.5 mg/kg total dose over 5 days, and then a trial of 7.5 mg/kg in merely one injection, but both regimens were only 90–93% effective (Table 1 regimens 3 and 4).7,8 To bring efficacy up to amphotericin B deoxycholate levels, the total dose of Ambisome was increased to 10 mg/kg. Both 10 mg/kg over 5 days and finally 10 mg/kg administered once showed a high efficacy rate of 96% (Table 1 regimens 5 and 6).5,9

ABLC/Abeclat: In a head-to-head comparison, Abeclat was inferior to Ambisome in efficacy (Table 1 regimen 7 versus regimen 5) and in tolerance (fever/chills were experienced by 76% of ABLC and 29% of Ambisome patients).7

ABCD/Amphocil: 97% efficacy was shown in a large trial using <1 week of therapy (Table 1 regimen 8).10

ABLE: 15 mg/kg in one injection was only 85% effective (Table 1 regimen 9).11

Fungisome: In this issue of the Journal, Sundar and others report the efficacy of one injection of 10 mg/kg or of 15 mg/kg in an early phase 2 study of 15 patients per cohort.15 One patient in each cohort relapsed therefore the cure rate was 14 of 15 (93%) for each regimen (Table 1 regimen 10). There was a 90% incidence of infusion-related fever and chills, and an ~25% incidence of diarrhea and vomiting. The 5 SAEs (2 nephrotoxicity, 2 thrombocytopenia, 1 pulmonary edema), even though reversible, may be considered frequent for a small 30-patient database.

In sum, systematic evaluation of Ambisome has led to a very high dose of 10 mg/kg administered in a very short period of time of 1 day, which in a large study showed 96% efficacy. This regimen is overall superior to the standard regimen of amphotericin B deoxycholate (1 mg/kg every other day for 15 infusions) on the basis of efficacy (almost equal), tolerance (superior), feasibility (far superior), and cost (~$200 for Ambisome at the developing-world favorable price). Ambisome 10 mg/kg once is now the treatment of choice for Indian subcontinent VL.12,13 ABCD/Amphocil showed excellent efficacy in a relatively short course; whether single dose Amphotilc is competitive with single dose Ambisome is not known. Other amphotericin B formulations were either inferior in efficacy (ABLC/Abeclat, ABLE) or have not yet been evaluated in large trials (Fungisome).

Other approaches toward replacing amphotericin B deoxycholate for Indian subcontinent VL are a parenteral agent that can be administered intramuscularly (paromomycin) and an oral agent (miltefosine).

In a large phase 3 study, 95% of Indian per-protocol patients were cured with a regimen of paromomycin 11 mg/kg/day for 21 days intramuscularly (Table 1 regimen 11).4 An attempt to shorten the inconvenient 21-day intramuscular treatment course to 14 days revealed low efficacy (84%) for the 2-week course (Table 1 regimen 12).14

In a large phase 3 trial, 97% of Indian per-protocol patients were cured with a regimen of miltefosine 2.5 mg/kg/day for 28 days (Table 1 regimen 13) after which miltefosine was made the VL treatment of choice in India, but after 10 years of use, the efficacy rate has fallen to 90% (Table 1 regimen 14).15 Gastrointestinal side effects are frequent, and miltefosine is contraindicated in pregnancy.

Combining short courses of two drugs will decrease the length of parenteral therapy and should protect against resistance including that to miltefosine and the aminoglycoside paromomycin. When combinations of short courses of 2-drug combinations of Ambisome, miltefosine, and paromomycin were evaluated, each combination was 99% effective (Table 1...
Drugs for visceral leishmaniasis in otherwise immunocompetant patients in the Indian subcontinent

| Regimen no. | Drug and route          | Per-protocol efficacy 6'-9' after RX† | Reference |
|-------------|-------------------------|--------------------------------------|-----------|
| 1           | Amphotericin B deoxycholate IV | 15 mg/kg over 30 days                | 96/96 (100%) | 3         |
| 2           | Amphotericin B deoxycholate IV | 15 mg/kg over 30 days                | 163/164 (99%) | 4         |
| 3           | Liposomal amphotericin B /Ambisome IV | 7.5 mg/kg over 5 days               | 26/28 (93%) | 7         |
| 4           | Liposomal amphotericin B /Ambisome IV | 7.5 mg/kg once                    | 183/203 (90%) | 8         |
| 5           | Liposomal amphotericin B /Ambisome IV | 10 mg/kg over 5 days               | 49/51 (96%) | 5         |
| 6           | Liposomal amphotericin B /Ambisome IV | 10 mg/kg once                    | 291/304 (96%) | 9         |
| 7           | ABLC/Abeclert IV           | 10 mg/kg over 5 days                | 47/51 (92%) | 8         |
| 8           | ABCD/Amplocil IV           | 7.5 mg/kg over 6 days               | 131/135 (97%) | 10        |
| 9           | ABLE                     | 15 mg/kg once                       | 317/373 (85%) | 11        |
| 10          | India-formulated Liposomal Amphotericin B/Fungisome IV | 10 mg/kg or 15 mg/kg once        | 14/15 (93%) | 12        |
| 11          | Paromomycin IM            | 11 mg/kg/d x 21 days                | 474/501 (95%) | 4         |
| 12          | Paromomycin IM            | 11 mg/kg/d x 14 days                | 183/217 (84%) | 14        |
| 13          | Miltefosine oral          | 2.5 mg/kg/day x 28 days             | 282/291 (97%) | 3         |
| 14          | Miltefosine oral          | 2.5 mg/kg/day x 28 days             | 512/567 (90%) | 15        |
| 15          | Ambisome IV + Miltefosine oral | 5mg/kg once + 2.5 mg/kg/d for 7 days | 155/157 (99%) | 16        |
| 16          | Ambisome IV + Paromomycin IM | 5mg/kg once + 11 mg/kg/d for 10 days | 153/155 (99%) | 16        |
| 17          | Miltefosine oral + Paromomycin IM | 2.5 mg/kg/day for 10 days + 11 mg/kg/d for 10 days | 156/158 (99%) | 16        |

Drugs for visceral leishmaniasis in otherwise immunocompetant patients in other endemic regions

| Regimen no. | Drug and route          | Per-protocol efficacy 6'-9' after RX† | Reference |
|-------------|-------------------------|--------------------------------------|-----------|
| 18          | Ambisome IV in East Africa | 7.5 mg/kg once                    | 8/20 (40% ) | 17        |
| 19          | Ambisome IV in East Africa | 21 mg/kg over 21 days              | 46/54 (85%) | 17        |
| 20          | Paromomycin in East Africa | 11 mg/kg/d x 21 days              | 80/121 (66%‡) | 19        |
| 21          | Ambisome IV in Mediterranean | 18 mg/kg over 10 days            | 41/42 (98%‡) | 18        |
| 22          | Pentavalent Antimony in East Africa | 20 mg/kg/day x 30 days       | 104/112 (93%‡) | 19        |
| 23          | Pentavalent Antimony in Mediterranean | 20 mg/kg/day x 30 days       | 47/52 (90%) | 20        |

*mg drug = mg active ingredient (amphotericin B, paromomycin base, miltefosine, antimony) in the formulation.
†Efficacy = no. cured/no. evaluable (%).
‡HIV-positive patients omitted from calculation.

regimens 15–17) in substantial numbers of patients. Choosing between these three combinations is difficult; the main reason not to use a combination involving Ambisome, miltefosine, or paromomycin is the need to maintain a cold-chain, female contraception, and 10 days of injections, respectively.

A fundamental issue with anti-VL chemotherapy is that although the incidence of VL in India is such that high-quality studies can be performed, efficacy against Indian VL does not convey to disease from other endemic regions. Ambisome and paromomycin regimens that were 90% and 95% effective against Leishmania donovani in India (Table 1 regimens 4 and 11) were 40% and 66% effective against Mediterranian disease caused by L. donovani (Table 1 regimen 22). Even 21 mg/kg Ambisome was only 85% effective in East Africa (Table 1 regimen 19), although such a dose seems effective against Mediterranean disease caused by Leishmania infantum (Table 1 regimen 21). On the other hand, the classic agent pentavalent antimony is still effective in East Africa (Table 1 regimen 22) and in the Mediterranean (Table 1 regimen 23). These discrepancies are important not just for non-Indian endemic regions, but for VL seen in the developed world. Visceral leishmaniasis is uncommon in the United States; however, the disease that this reviewer has seen in the US was contacted in the Mediterranean, East Africa, and Brazil. Other issues for the United States are FDA approval (Ambisome, Abeclert, and miltefosine are approved products in the United States) and pricing that will not be the same as in India.

Remarkable progress has been made in VL chemotherapy with the advent of an oral agent and then short-course Ambisome in the Indian subcontinent. The future of anti-VL chemotherapy is likely to involve further fine-tuning of lipid formulations of amphotericin B such as Fungisome to possibly compete with Ambisome in the Indian subcontinent; use of single agent approved products in the developed nations; and combination therapy worldwide, to include clinically resistant disease in patients with underlying immunodeficiencies.

REFERENCES

1. Alvar J, Vélez ID, Bern C, Herrero M, Desjeux P, Cano J, Jannin J, den Boer M; Leishmaniasis Control Team WHO, 2012. Leishmaniasis worldwide and global estimates of its incidence. PLoS ONE 7: e35671.
2. Sundar S, 2001. Drug resistance in Indian visceral leishmaniasis. Trop Med Int Health 6: 849–854.
3. Sundar S, Jha TK, Thakur CP, Engel J, Sindermann H, Fischer C, Junge K, Brycecion A, Berman J, 2002. Oral miltefosine for Indian visceral leishmaniasis. N Engl J Med 347: 1739–1746.
4. Sundar S, Jha TK, Thakur CP, Sinha PK, Bhattacharya SK, 2007. Injectable paromomycin for visceral leishmaniasis in India. N Engl J Med 356: 2571–2581.
5. Sundar S, Mehta H, Shuresh AV, Singh SP, Rai M, Murray HW, 2004. Amphotericin B treatment for Indian visceral leishmaniasis: conventional versus lipid formulations. Clin Infect Dis 38: 377–383.
6. Ambisome label. 2000. Available at: http://www.accessdata.fda.gov/drugsatfda_docs/nda/2000/50-740S001_AmBisome_prntlbl.pdf. Accessed December 10, 2014.
7. Sundar S, Jha TK, Thakur CP, Mishra M, Singh VR, Buffels R, 2002. Low-dose liposomal amphotericin B in refractory Indian visceral leishmaniasis: a multicenter study. Am J Trop Med Hyg 66: 143–146.
8. Sundar S, Jha TK, Thakur CP, Mishra M, Singh VP, Buffels R, 2003. Single-dose liposomal amphotericin B in the treatment of visceral leishmaniasis in India: a multicenter study. Clin Infect Dis 37: 800–804.
9. Sundar S, Chakravarty J, Agarwal D, Rai M, Murray HW, 2010. Single-dose liposomal amphotericin B for visceral leishmaniasis in India. N Engl J Med 362: 504–512.
10. Sundar S, Mehta H, Chhabra A, Singh V, Chauhan V, Desjeux P, Rai M, 2006. Amphotericin B colloidal dispersion for the treatment of Indian visceral leishmaniasis. Clin Infect Dis 42: 608–613.
11. Sundar S, Pandey K, Thakur CP, Jha TK, Das VN, Verma N, Lal CS, Verma D, Alam S, Das P, 2014. Efficacy and safety of amphotericin B emulsion versus liposomal formulation in Indian patients with visceral leishmaniasis: a randomized, open-label study. PLoS Negl Trop Dis 8: e3169.
12. Sundar S, Singh A, Rai M, Chakravarty J, 2015. Single-dose indigenous liposomal amphotericin B in the treatment of Indian visceral leishmaniasis: a phase 2 study. Am J Trop Med Hyg 92: 513–517.
13. Mondal D, Alvar J, Hasnain MG, Hossain MS, Ghosh D, Huda MM, Nabi SG, Sundar S, Matlashewski G, Arana B, 2014. Efficacy and safety of single-dose liposomal amphotericin B for visceral leishmaniasis in a rural public hospital in Bangladesh: a feasibility study. Lancet Glob Health 2: e51–57.
14. Sundar S, Agrawal N, Arora R, Agarwal D, Rai M, Chakravarty J, 2009. Short-course paromomycin treatment of visceral leishmaniasis in India: 14-day vs. 21-day treatment. Clin Infect Dis 49: 914–918.
15. Sundar S, Singh A, Rai M, Prajapati VK, Singh AK, Ostyn B, Boelaert M, Dujardin JC, Chakravarty J, 2012. Efficacy of miltefosine in the treatment of visceral leishmaniasis in India after a decade of use. Clin Infect Dis 55: 543–550.
16. Sundar S, Sinha PK, Rai M, Verma DK, Nawin K, Alam S, Chakravarty J, Vaillant M, Verma N, Pandey K, Kumari P, Lal CS, Arora R, Sharma B, Ellis S, Strub-Wourgaft N, Balasgaram M, Olliaro P, Das P, Modabber F, 2011. Comparison of short-course multidrug treatment with standard therapy for visceral leishmaniasis in India: an open-label, non-inferiority, randomized controlled trial. Lancet 377: 477–486.
17. Khalil EA, Weldegebreal T, Younis BM, Omollo R, Musa AM, Hailu W, Abuzaid AA, Dorlo TP, Hurissa Z, Yifruf S, Haleke W, Smith PG, Ellis S, Balasagaram M, EL-Hassan AM, Schoone GJ, Wasunna M, Kimutai R, Edwards T, Hailu A, 2014. Safety and efficacy of single dose versus multiple doses of AmBisome for treatment of visceral leishmaniasis in eastern Africa: a randomized trial. PLoS Negl Trop Dis 8: e2613.
18. Davidson RN, di Martino L, Gradoni L, Giacchino R, Gaeta GB, Pempinello R, Scotti S, Cusco A, Castagnola E, Maisto A, Gramiccia M, di Caprio D, Wilkinson RJ, Bryceson AD, 1996. Short-course treatment of visceral leishmaniasis with liposomal amphotericin B (AmBisome). Clin Infect Dis 22: 938–943.
19. Hailu A, Musa A, Wasunna M, Balasagaram M, Yifruf S, Mengistu G, Hurissa Z, Hailu W, Weldegebreal T, Tesfaye S, Makonnen E, Khalil E, Ahmed O, Fadlalla A, El-Hassan A, Raheem M, Mueller M, Koummuki Y, Rashid J, Mbui J, Mucee G, Njoroge S, Manduku E, Musibi A, Mutuma G, Kirui F, Lodenyo H, Mutea D, Kirigi G, Edwards T, Smith P, Muthami L, Royce C, Ellis S, Abolo M, Omollo R, Kesujo J, Owiti R, Kinuthia J: Leishmaniasis East Africa Platform (LEAP) group, 2010. Geographical variation in the response of visceral leishmaniasis to paromomycin in East Africa: a multicentre, open-label, randomized trial. PLoS Negl Trop Dis 4: e709.
20. Syriopoulou V, Daikos GL, Theodoridou M, Pavlopoulou I, Manolaki AG, Sereti E, Karamboula A, Papathanasiou D, Krikos X, Saroglou G, 2003. Two doses of a lipid formulation of amphotericin B for the treatment of Mediterranean visceral leishmaniasis. Clin Infect Dis 36: 560–566.