Reduction in the nephrotoxicity of amphotericin B when administered in 20% intralipid

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**BRIEF REPORT**

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S SALAMA, C ROTSTEIN. Reduction in the nephrotoxicity of amphotericin B when administered in 20% intralipid. Can J Infect Dis 1997;8(3):157-160. The administration of amphotericin B (AmB) is often limited by the development of nephrotoxicity. In a pilot crossover trial, aqueous AmB followed by a new preparation of a mixture of AmB with 20% intralipid (AmB-IL) was administered to 10 immunocompromised patients for systemic fungal infections caused by Candida species. Mean total dose and duration of therapy with AmB-IL exceeded that of aqueous AmB (649±165 mg versus 394±105 mg, P=0.061 and 13.2±2.5 days versus 9±2.1 days, P=0.31). However, mean creatinine clearance of the patients rose during AmB-IL therapy by 10.7±7.7 mL/min (P=0.03). AmB-IL warrant further investigation to assess its stability and efficacy for treating serious fungal infections.

Key Words: Amphotericin B, Candida species, Nephrotoxicity

**Réduction de la néphrotoxicité de l’amphotéricine B lors de son administration avec l’intralipide à 20 %**

RÉSUMÉ : L’administration d’amphotéricine B est souvent limitée par l’apparition de signes de néphrotoxicité. Dans le cadre d’un essai pilote avec permutation des groupes, de l’amphotéricine B aqueuse (AmB), suivie d’une nouvelle préparation d’amphotéricine B mélangée à l’intralipide à 20 % (AmB-IL) a été administrée à dix patients immunodéficients pour des infections fongiques à Candida généralisées. La dose moyenne totale et la durée du traitement par AmB-IL ont été supérieures à celles du traitement par AmB aqueux (649 ± 165 mg contre 394 ± 105 mg, P = 0,061 et 13,2 ± 2,5 jours contre 9 ± 2,1 jours, P = 0,31). Toutefois, la clairance moyenne de la créatinine des patients a augmenté de 10,7 ± 7,7 mL/min (P = 0,03) lors du traitement par AmB-IL. Les résultats enregistrés avec l’AmB-IL justifient la poursuite des recherches pour évaluer sa stabilité et son efficacité dans le traitement des infections fongiques graves.

**Fungal infections have begun to rival those caused by bacteria as significant nosocomial problems (1). Improvement in the prognosis of patients with cancer, immunodeficiencies and connective tissue disorders has created a patient population that more susceptible to infection. The increase in the incidence of nosocomial fungal infection since 1980 has mirrored the expansion of the immunocompromised patient population (2). These infections are often severe, progress rapidly and are refractory to therapy. Despite the availability and relative success of newer sys-**
emic antifungal agents such as fluconazole (3) and itraconazole (4), amphotericin B (AmB) has remained the treatment of choice for systemic fungal infections. However, the clinical use of AmB is complicated by the development of a variety of adverse events including fever, chills, nausea, vomiting, hypokalemia, hypomagnesemia and anemia (5). The most formidable of these effects, nephrotoxicity, may actually restrict AmB’s effectiveness because of premature discontinuation of antifungal therapy or suboptimal dosing (6).

One approach to circumventing nephrotoxicity has been the formulation of preparations of AmB that are less toxic (7). As a result, AmB has been entrapped in liposomes, ambisome or amphotericin B lipid complex (ABLC), and complexed with cholesteryl sulfate, amphotericin B colloidal dispersion (ABCD). Ambisome is a liposomal preparation of small unilamellar vesicles with a particle size 80 nm, containing a 10% concentration of AmB in the membrane (8). ABLC is an AmB lipid complex with a particle size of 1.6 to 11 µm that includes an approximately 33% concentration of AmB in microparticulate polymorphic sheets and ribbons (8). Finally, ABCD is a stable complex of AmB and cholesteryl sulfate intercalated in a 1:1 ratio in disc-like structures with a diameter of 122 nm (8). All three formulations have been administered at higher dosages, 5 to 7 mg/kg/day, than AmB (9). These preparations have proven to be efficacious with reduced renal toxicity for the treatment of fungal infections in immunocompromised hosts (10-12). Nevertheless, the formulations are very expensive. An alternative, less expensive delivery system has been studied by Anaissie et al (11), Chavanet et al (13) and Caillot et al (14). This lipid emulsion mixture with AmB was demonstrated to reduce toxicity while maintaining efficacy (11,13,14). We, therefore, decided to conduct our own pilot study to assess the effect of AmB mixed with 20% intralipid on the renal function of immunocompromised patients.

### PATIENTS AND METHODS

Ten immunocompromised patients (eight with an underlying hematological malignancy, one with diabetes mellitus and rheumatoid arthritis, and one with Crohn’s disease) with a systemic fungal infection due to a *Candida* species were initially given AmB in aqueous solution. The presence of a systemic fungal infection was determined by the patient’s attending physician based on microbiological documentation of *Candida* species from a body site in the clinical setting of fever, presumed to be infectious in etiology.

Aqueous AmB was given at a dose of 0.6 mg/kg/day intravenously after an initial test dose of 1 mg in 100 mL of dextrose and water solution infused intravenously over 1 h. AmB was reconstituted in 10 mL of dextrose and water solution and subsequently diluted in 500 mL of the same intravenous solution. The aqueous AmB solution was then infused intravenously daily over 4 h. Premedication with acetaminophen 650 mg orally, intravenous diphenhydramine 25 or 50 mg and intravenous meperidine 25 or 50 mg was given to each patient.

Each patient was crossed over from aqueous AmB to AmB in 20% intralipid (AmB-IL). This occurred during the treatment course in six cases and at the onset of nephrotoxicity in four patients. The same premedication was employed. Nephrotoxicity was defined as a 50% decrease from the baseline value in calculated creatinine clearance (15).

AmB-IL was prepared daily in the following manner. Based

### TABLE 1

<table>
<thead>
<tr>
<th>Patient number</th>
<th>Age (years)</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>Type of organism</th>
<th>Site of infection</th>
<th>Amphotericin B derived nephrotoxicity</th>
<th>Lipid</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>44</td>
<td>Female</td>
<td>Acute leukemia</td>
<td><em>Candida albicans</em></td>
<td>Urine, mouth, lungs</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>2</td>
<td>40</td>
<td>Female</td>
<td>Acute leukemia</td>
<td><em>Candida albicans</em></td>
<td>Urine</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>3</td>
<td>59</td>
<td>Male</td>
<td>Acute leukemia</td>
<td><em>Candida albicans, Torulopsis glabrata</em></td>
<td>Gastrointestinal tract, mouth</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>4</td>
<td>67</td>
<td>Female</td>
<td>Acute leukemia</td>
<td><em>Candida tropicalis</em></td>
<td>Blood</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>5</td>
<td>53</td>
<td>Female</td>
<td>Lymphoma</td>
<td><em>Candida albicans</em></td>
<td>Blood, urine, lungs</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td>57</td>
<td>Male</td>
<td>Acute leukemia</td>
<td><em>Candida albicans, Candida lambica</em></td>
<td>Gastrointestinal tract</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>7</td>
<td>32</td>
<td>Female</td>
<td>Lymphoma</td>
<td><em>Candida lusitaniae</em></td>
<td>Vagina</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>8</td>
<td>73</td>
<td>Male</td>
<td>Diabetes mellitus, rheumatoid arthritis</td>
<td><em>Candida albicans</em></td>
<td>Blood, urine</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>9</td>
<td>60</td>
<td>Female</td>
<td>Acute leukemia</td>
<td><em>Candida albicans</em></td>
<td>Urine, mouth, gastrointestinal tract</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>10</td>
<td>36</td>
<td>Male</td>
<td>Crohn’s disease</td>
<td><em>Candida albicans</em></td>
<td>Blood</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

– Nephrotoxicity absent; + Nephrotoxicity present
on a dosage of 1 mg/kg/day, the daily requirement of AmB to be administered was calculated. AmB (50 mg/vial) was then reconstituted in 10 mL of sterile water for injection to yield a final concentration of 5 mg/mL. The required daily dosage of AmB was then transferred from the reconstituted preparation of 5 mg/mL to a 100 mL bottle of 20% intralipid (Kabi Pharmacia Canada Inc) fat emulsion for parenteral use. This was mixed and shaken thoroughly for 2 mins. The mixture was protected from light and refrigerated. The AmB-IL formulation was infused intravenously over 2 h within 24 h of its preparation. No premedication was administered.

Patient characteristics were recorded including age; sex; underlying diagnosis; site of infection; organism type; baseline and end of therapy serum creatinines, potassium, magnesium, aspartate aminotransferase, alanine aminotransferase, gamma-glutamyl transpeptidase, alkaline phosphatase and bilirubin; duration of therapy with aqueous AmB and AmB-IL. The total dose administered of each formulation was also recorded. Mean total dose of each formulation of AmB, mean duration of treatment and mean change in calculated creatinine clearance based on initial and end of therapy values for each formulation of AmB, respectively, were compared by paired Student’s t tests.

RESULTS

Ten immunocompromised patients were initially treated with aqueous AmB and were then crossed over to AmB-IL. Table 1 details the characteristics of the patients. Six patients with underlying acute leukemia and two patients with lymphoma had recently completed antineoplastic chemotherapy. One patient, with a history of diabetes mellitus, rheumatoid arthritis and renal insufficiency, developed candiduria and candidemia. Another patient developed a central venous access device-related candidemia while receiving total parenteral nutrition for Crohn’s disease. There were six female and four male patients, mean age 52 years. Causative pathogens were *Candida albicans* in six patients, *Candida tropicalis* in one, *Candida lusitaniae* in one, *C albicans* with *Torulopsis glabrata* in one, and *C albicans* with *Candida lambica* in one. Sources of infection were urine, bloodstream and gastrointestinal tract (neutropenic enteropathy).

The mean total dose of AmB-IL exceeded that of aqueous AmB (649±165 mg versus 344±105 mg, P=0.06), as did the mean duration of treatment (13.2±2.5 versus 9±2.1 days; P=0.31) (Table 2). However, during the aqueous AmB treatment, the mean creatinine clearance for the group dropped by 23.5±9.2 mL/min (Figure 1). Thereafter, during AmB-IL therapy, the mean creatinine clearance rose by 10.7±7.7 mL/min.

**TABLE 2**

<table>
<thead>
<tr>
<th></th>
<th>Mean total dose (mg ± SD)</th>
<th>Mean treatment duration (days ± SD)</th>
<th>Mean change in creatinine clearance (mL/min ± SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amphotericin B</td>
<td>344±105</td>
<td>9±2.1</td>
<td>-23.5±9.2</td>
</tr>
<tr>
<td>Amphotericin B in intralipid</td>
<td>649±165</td>
<td>13.2±2.5</td>
<td>+10.7±7.7</td>
</tr>
<tr>
<td>P</td>
<td>0.064</td>
<td>0.307</td>
<td>0.026</td>
</tr>
</tbody>
</table>

**Table 3**

<table>
<thead>
<tr>
<th>Biochemical parameter</th>
<th>Aqueous amphotericin B</th>
<th>Amphotericin B in intralipid</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum potassium (mmol/L)</td>
<td>0.14±0.3</td>
<td>0.4±1.2</td>
<td>0.95</td>
</tr>
<tr>
<td>Serum magnesium (mmol/L)</td>
<td>0.036±0.1</td>
<td>0.009±0.12</td>
<td>0.93</td>
</tr>
<tr>
<td>Asparate aminotransferase (U/L)</td>
<td>-2.0±5.1</td>
<td>10.4±17.6</td>
<td>0.97</td>
</tr>
<tr>
<td>Alanine transferase (U/L)</td>
<td>7.9±22.7</td>
<td>15.2±31.7</td>
<td>1.03</td>
</tr>
<tr>
<td>Gamma-glutamyl transpeptidase (U/L)</td>
<td>-2.3±56.3</td>
<td>49.3±123.8</td>
<td>1.15</td>
</tr>
<tr>
<td>Alkaline phosphate (U/L)</td>
<td>21.2±28.3</td>
<td>11.4±44.8</td>
<td>1.04</td>
</tr>
<tr>
<td>Total bilirubin (µmol/L)</td>
<td>6.1±10.3</td>
<td>3.3±23.3</td>
<td>0.59</td>
</tr>
</tbody>
</table>

**Figure 1** Baseline and ending creatinine clearance values for patients treated with aqueous amphotericin B. Creatinine clearance was calculated using the Cockcroft and Gault formula.

**Figure 2** Baseline and ending creatinine clearance values for patients treated with amphotericin B in intralipid. Creatinine clearance was calculated using the Cockcroft and Gault formula.
was recognized, other factors such as concomitant hypoten-
tions. First, because aqueous AmB was initiated when sepsis
the addition of flucytosine (11). This latter finding may have been related to
significant reductions in creatinine clearance with a fat emulsion
administration, even with administration of
more of the latter preparation. Some investigators have dem-
strated a renal sparing effect with AmB-IL compared with
aqueous AmB (10); nevertheless, others have still noted sig-
nificant reductions in creatinine clearance with a fat emulsion
preparation (11). This latter finding may have been related to
the addition of flucytosine (11).

However, there are a number of limitations to our observa-
tions. First, because aqueous AmB was initiated when sepsis
was recognized, other factors such as concomitant hypoten-
sion or other nephrotoxins may have contributed to the neph-
rotoxicity observed. Second, we could not assess the efficacy
of the AmB-IL preparation. This is relevant because it has been
speculated that AmB-IL produces less toxicity by trapping and
binding AmB molecules in the lipid portion, thus reducing the
availability of free and active AmB compound (7). Similarly,
serum concentrations of free AmB in the AmB-IL mixture are
lower than aqueous AmB (10). It may be necessary to infuse
higher doses of AmB-IL to attain the efficacy rates achieved by
aqueous AmB (7,10). Third, we did not demonstrate the per-
manent stability and absolute safety of our mixture. Concerns
have been raised that AmB-IL may contain particles at least 10
µm in diameter (16) and that these particles may lodge in
blood vessels. Aqueous AmB consistently had particles of less
than 10 µm. Finally, it would have been advantageous for all
our patients to cross over from aqueous AmB to AmB-IL at a
specified AmB dose.

In this crossover pilot study where patients acted as their
own controls, AmB-induced nephrotoxicity was significantly
reduced with AmB-IL. This mixture warrants further evalu-
ation of its long term stability, safety, clinical efficacy and
maximum tolerated dose. Because the costs of AmB-IL are
substantially less than ABLC, ABCD or liposomal AmB, the
mixture may prove to be a welcome addition to our antifungal
armamentarium.

REFERENCES