Oral v. pulse intravenous cyclophosphamide: A retrospective analysis of adverse events in a setting with a high burden of infectious disease

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Background. Cyclophosphamide (CPM) is still considered to be the first-line treatment for many life-threatening autoimmune conditions. It does, however, carry a significant risk of severe adverse events, especially infections. At present CPM is administered as either a daily oral dose (DOC) or an intravenous pulse (PIVC). There is uncertainty regarding the safety profiles of both regimens in settings with a high burden of infectious diseases.

Objective. To compare the frequency and nature of adverse events related to the use of DOC and PIVC in such a setting.

Methods. A cohort of patients treated with CPM for autoimmune diseases at Tygerberg Academic Hospital, Cape Town, South Africa, from 1 January 2008 to 31 May 2013 was studied. We compared participants receiving DOC and PIVC with regard to disease characteristics and the occurrence of major adverse events.

Results. A total of 134 participants (92 DOC and 42 PIVC) were included. Participants in the DOC group were treated for longer (174 v. 101 days; p<0.01) and with higher cumulative doses (17 276 v. 3 327 mg; p<0.01). Risk of infection was similar in the two groups, although there were 6 deaths from leucopenic sepsis in the DOC group (v. 0; p=0.18). Nadir leucocyte counts were also lower in the DOC group (median 3.8 v. 5.3 × 10⁹/L; p=0.02).

Conclusion. Infection rates in the two groups were similar, but DOC was associated with longer treatment duration, greater cumulative CPM doses and more severe leucopenia. If resources allow and available literature provides support for efficacy, consideration should be given to greater use of PIVC.


Autoimmune diseases affecting major organs carry significant morbidity and mortality. Immunosuppressive therapy with agents such as cyclophosphamide (CPM) significantly improves quality of life and survival in many patients suffering from such diseases.1,2

CPM is still considered to be the first-line treatment for many life-threatening autoimmune conditions owing to its well-established therapeutic benefits. It does, however, carry a significant risk of severe adverse events including cytopenias, infections, infertility and bladder toxicity. At present CPM is administered either as a daily oral dose (DOC) or as an intravenous pulse at various intervals (PIVC). The optimal dosing route and regimen remain controversial, especially in the treatment of lupus nephritis.

Large head-to-head trials comparing DOC with PIVC are lacking. Many opinions exist on the optimal dosing regimen, based on small studies or indirect conclusions from larger studies. Overall it seems that DOC has the advantages of lower cost, simpler administration and avoidance of high-dose exposure should the drug be inappropriately administered.3,4 PIVC has the advantage of a reduced incidence of short- and long-term adverse events and avoids the need for daily compliance with treatment. Data regarding the comparative efficacy of the two regimens are conflicting and briefly discussed below.

Total drug costs for treatment regimens are lower for DOC than for PIVC. PIVC also requires allocated space, consumables and personnel to administer the infusion. Patients need to take time off work and have additional travel expenses, which all contribute to the overall cost of this regimen.3,4

Although DOC is associated with an increase in long-term side-effects, adverse events due to inappropriate dosing are more easily managed. Examples include administration when a contraindication exists, mistaking sepsis for a flare of disease or not adjusting doses for renal function. Reducing or omitting further oral doses under these circumstances, which cannot be done with PIVC, reduces the toxicity.3,4

Austin et al.5 initially published data in 1986 supporting superior treatment outcomes and reduced side-effect profiles with the use of PIVC as opposed to DOC in patients with lupus nephritis. This led to PIVC becoming the standard treatment in lupus nephritis, with fewer subsequent trials and recommendations that include DOC.3,4 This practice is now contested, as subsequent trials4,7,8 and a recent Cochrane review9 showed no difference in outcomes or adverse events. An increased tendency to relapse has also been shown in patients with lupus nephritis treated with PIVC.10 DOC was associated with a higher frequency of adverse events in the same study. Another small trial showed a slightly larger increase in serum
albin in patients with membranous nephropathy treated with PIVC in comparison with DOC.\textsuperscript{[11]}

Trials in antineutrophil cytoplasmic antibody (ANCA)-associated vasculitides have also suggested equal to increased ability of PIVC to induce remission compared with DOC, but a greater tendency towards relapse.\textsuperscript{[12-14]} Episodes of leucopenia and infection were less frequent in ANCA-associated vasculitides treated with PIVC.\textsuperscript{[15-17]} Increased gonadal toxicity with oral CPM has also been shown in patients treated for ANCA-associated vasculitides.\textsuperscript{[18]} Decreased toxicity with PIVC is thought to be due to the lower cumulative CPM doses used.

No difference in outcomes has been shown between the two regimens in treating pulmonary involvement in scleroderma.\textsuperscript{[19]}

The use of CPM in a resource-constrained and infectious diseases-burdened environment, as in our setting at Tygerberg Academic Hospital (TBAH), Cape Town, South Africa, has unique challenges and considerations in respect of cost, efficacy and safety. Cost of treatment to hospitals, lack of allocated space and trained personnel to administer intravenous medication, patients’ ability to travel to a tertiary setting to receive treatment, compliance with treatment and follow-up, access to healthcare in the event of drug toxicity, and the overall increased burden of infectious diseases associated with a low socioeconomic environment are all factors to consider in choosing an appropriate treatment plan.

Of special concern in our setting is safety. The high prevalence of tuberculosis (TB) in communities serviced by TBAH\textsuperscript{[20]} and exposure to overpopulated and unhygienic environments can significantly endanger patients should they become severely immunosuppressed. To our knowledge, no studies comparing DOC and PIVC are available that address these safety issues when CPM is used in such settings.

Clinicians at TBAH make use of both intravenous pulse and oral regimens of CPM for the treatment of autoimmune diseases. Treatment choice is largely based on the condition treated and the preference of the discipline involved in initiating treatment (e.g. lupus nephritis is generally treated with DOC by nephrologists, while preference of the discipline involved in initiating treatment (e.g. rheumatologists) predominates in initiating treatment for non-rheumatological disease, including transplant recipients). Treatment choice is largely based on the condition treated and the preference of the discipline involved in initiating treatment (e.g. lupus nephritis is generally treated with DOC by nephrologists, while preference of the discipline involved in initiating treatment (e.g. rheumatologists) predominates in initiating treatment for non-rheumatological disease, including transplant recipients).

Objective

Our primary objective was to compare the frequency and nature of adverse events related to the use of oral and intravenous CPM in patients with autoimmune diseases.

Methods

A retrospective cohort study was conducted at TBAH to compare patients treated with DOC with those who were treated with PIVC. Data were extracted from hospital records of adult patients (>18 years) suffering from autoimmune diseases treated with CPM. Pharmacy records were used to identify all patients who started treatment between 1 January 2008 and 31 May 2013. Participants were only included if they had been on treatment with CPM for at least 1 month, unless adverse events due to CPM occurred earlier, in which case the participant was also included. Patients who had received CPM for indications other than autoimmune disease (e.g. chemotherapy, transplant recipients) were excluded. Patients on treatment for purely neurological diseases were also excluded owing to the different dosing regimens and treatment duration used. Data collected included demographics, comorbid disease, mass, diagnosis and disease involvement, duration of disease prior to CPM initiation, treatment prior to CPM initiation, baseline leucocyte and neutrophil counts, indication for CPM therapy, initial dosage, subsequent adjustment to dosing and indication for changes, cumulative total drug dose, concurrent immunosuppressive drug use, use of preventive therapy (isoniazid, co-trimoxazole, contraception and gonadal protection), documentation of counselling regarding contraception, occurrence of major side-effects and discontinuation due to drug toxicity. Data collected on side-effects included cytopenias, infections, bladder toxicity and infertility. Adverse events were reported and graded according to the Common Terminology Criteria for Adverse Events guidelines.\textsuperscript{[21]}

Data were captured on a Microsoft Excel spreadsheet and analysed using the Mann-Whitney U-test, while categorical data were analysed using the χ 2 and Fisher’s exact tests. A significance level of 5% was used throughout.

The study was approved by the Health Research Ethics Committee of Stellenbosch University (protocol No. S13/07/121). Owing to the retrospective nature of the study, the difficulty in tracing individual participants and the low risk to the participants, a waiver of informed consent was granted. Data were collected on standardised capture sheets using study codes assigned to each participant. Codes linking confidential data to the identity of the participants were kept separately and securely.

Results

After exclusion of oncology patients, 221 subjects were identified. A total of 134 participants were included in the study, 92 in the DOC group and 42 in the PIVC group. Eighty-seven subjects were excluded for reasons that included treatment for non-rheumatological disease, pharmacy errors and missing records, early withdrawal of treatment not related to drug side-effects, failure to follow up after initiation of treatment, concomitant participation in other drug trials, or receiving both PIVC and DOC.

The characteristics of the participants are summarised in Table 1. There were no significant differences between the groups in terms of age, mass, comorbid disease (HIV and diabetes mellitus) and leucocyte count at baseline. Overall, participants were young (mean age 34 years) with a female predominance (74.6%), especially in the PIVC group (85.7 v. 69.6%; p=0.03).

As expected, there was a marked difference in disease spectrum between the two groups, both in terms of primary diagnosis and organ involvement.

The indications for starting CPM varied significantly. The most frequent indication in the DOC group was class IV lupus nephritis (32.6%), followed by non-lupus-related renal disease including nephrotic syndrome (27.1%) and rapidly progressive (RPGN) or crescentic glomerulonephritis (GN) (21.7%). The most frequent indication in the PIVC group was neuropsychiatric lupus (45.2%), followed by interstitial lung disease due to various autoimmune diseases (28.6%), and lupus myocarditis (28.6%).

A total of 676 patient-months of treatment were observed. Treatment details are summarised in Table 2. Participants in the PIVC group were treated for significantly shorter periods (mean 101 v. 174 days; p<0.01) and had much lower mean cumulative doses (3 327 v. 17 276 mg; p<0.01).
Table 1. Participant baseline characteristics

<table>
<thead>
<tr>
<th></th>
<th>All (N=134)</th>
<th>DOC (N=92)</th>
<th>PIVC (N=42)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gender, n (%)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M</td>
<td>34 (25.4)</td>
<td>28 (30.4)</td>
<td>6 (14.3)</td>
<td>0.03</td>
</tr>
<tr>
<td>F</td>
<td>100 (74.6)</td>
<td>64 (69.6)</td>
<td>36 (85.7)</td>
<td></td>
</tr>
<tr>
<td><strong>Age (years), mean (SD)</strong></td>
<td>34.0 (12.7)</td>
<td>34.2 (13.3)</td>
<td>33.4 (11.4)</td>
<td>0.95</td>
</tr>
<tr>
<td><strong>Mass (kg), mean (SD)</strong></td>
<td>69.5 (18.4)</td>
<td>70.5 (18.0)</td>
<td>67.0 (19.5)</td>
<td>0.37</td>
</tr>
<tr>
<td><strong>Comorbid disease, n (%)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>15 (11.2)</td>
<td>9 (9.8)</td>
<td>6 (14.3)</td>
<td>0.44</td>
</tr>
<tr>
<td>HIV</td>
<td>5 (3.7)</td>
<td>5 (5.4)</td>
<td>0 (0.0)</td>
<td>0.12</td>
</tr>
<tr>
<td><strong>Primary diagnosis, n (%)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SLE</td>
<td>63 (47.0)</td>
<td>41 (44.6)</td>
<td>22 (52.4)</td>
<td>0.55</td>
</tr>
<tr>
<td>Membranous GN</td>
<td>16 (11.9)</td>
<td>16 (17.4)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>Mesangiocapillary GN</td>
<td>7 (5.2)</td>
<td>7 (7.6)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>SLE/systemic sclerosis overlap</td>
<td>5 (3.7)</td>
<td>3 (3.3)</td>
<td>2 (4.8)</td>
<td>*</td>
</tr>
<tr>
<td>Crescentic GN</td>
<td>5 (3.7)</td>
<td>5 (5.4)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>IgA nephropathy</td>
<td>5 (3.7)</td>
<td>5 (5.4)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>Systemic sclerosis</td>
<td>8 (6.0)</td>
<td>2 (2.2)</td>
<td>6 (14.3)</td>
<td>*</td>
</tr>
<tr>
<td>Granulomatosis with polyangiitis</td>
<td>11 (8.2)</td>
<td>5 (5.4)</td>
<td>6 (14.3)</td>
<td>*</td>
</tr>
<tr>
<td>FSGS</td>
<td>2 (1.5)</td>
<td>2 (2.2)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>Goodpasture's syndrome</td>
<td>1 (0.8)</td>
<td>1 (1.1)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>Mesangiproliferative GN</td>
<td>3 (2.2)</td>
<td>3 (3.3)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>Minimal-change nephropathy</td>
<td>2 (1.5)</td>
<td>2 (2.2)</td>
<td>0 (0.0)</td>
<td>*</td>
</tr>
<tr>
<td>Eosinophilic granulomatosis with polyangiitis</td>
<td>1 (0.8)</td>
<td>0 (0.0)</td>
<td>1 (2.4)</td>
<td>*</td>
</tr>
<tr>
<td>SLE/rheumatoid arthritis overlap</td>
<td>1 (0.8)</td>
<td>0 (0.0)</td>
<td>1 (2.4)</td>
<td>*</td>
</tr>
<tr>
<td>SLE/dermatomyositis overlap</td>
<td>1 (0.8)</td>
<td>0 (0.0)</td>
<td>1 (2.4)</td>
<td>*</td>
</tr>
<tr>
<td>SLE/polymyositis overlap</td>
<td>1 (0.8)</td>
<td>0 (0.0)</td>
<td>1 (2.4)</td>
<td>*</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>1 (0.8)</td>
<td>0 (0.0)</td>
<td>2 (4.8)</td>
<td>*</td>
</tr>
<tr>
<td>Vasculitis of unknown cause</td>
<td>1 (0.8)</td>
<td>0 (0.0)</td>
<td>1 (2.4)</td>
<td>*</td>
</tr>
<tr>
<td><strong>Indication for CPM, n (%)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RPGN/crescentic GN other than lupus</td>
<td>25 (18.7)</td>
<td>20 (21.7)</td>
<td>5 (11.9)</td>
<td>0.18</td>
</tr>
<tr>
<td>Nephrotic syndrome other than lupus</td>
<td>25 (18.7)</td>
<td>25 (27.2)</td>
<td>0 (0.0)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Vasculitis</td>
<td>4 (3.0)</td>
<td>0 (0.0)</td>
<td>4 (9.5)</td>
<td>0.01</td>
</tr>
<tr>
<td>Class III lupus nephritis</td>
<td>5 (3.7)</td>
<td>5 (5.4)</td>
<td>0 (0.0)</td>
<td>0.12</td>
</tr>
<tr>
<td>Class IV lupus nephritis</td>
<td>31 (23.1)</td>
<td>29 (31.5)</td>
<td>2 (4.8)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Class V lupus nephritis</td>
<td>7 (5.2)</td>
<td>5 (5.4)</td>
<td>2 (4.8)</td>
<td>0.87</td>
</tr>
<tr>
<td>Interstitial lung disease</td>
<td>20 (14.9)</td>
<td>8 (8.7)</td>
<td>12 (28.6)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Lupus myocarditis</td>
<td>16 (11.9)</td>
<td>4 (4.4)</td>
<td>12 (28.6)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Eye disease (granulomatosis)</td>
<td>1 (0.8)</td>
<td>1 (1.1)</td>
<td>0 (0.0)</td>
<td>0.50</td>
</tr>
<tr>
<td>Neuropsychiatric lupus</td>
<td>21 (15.7)</td>
<td>2 (2.2)</td>
<td>19 (45.2)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td><strong>Diagnosis to initiation of CPM (days), mean (SD)</strong></td>
<td>23.3 (44.8)</td>
<td>17.4 (42.2)</td>
<td>36.2 (48.1)</td>
<td>0.04</td>
</tr>
<tr>
<td><strong>Duration of disease prior to CPM initiation (months), n (%)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 - 3</td>
<td>70 (52.2)</td>
<td>51 (55.4)</td>
<td>19 (45.2)</td>
<td>0.04</td>
</tr>
<tr>
<td>3 - 12</td>
<td>19 (14.2)</td>
<td>16 (17.4)</td>
<td>3 (7.1)</td>
<td>0.04</td>
</tr>
<tr>
<td>&gt;12</td>
<td>45 (33.6)</td>
<td>25 (27.2)</td>
<td>20 (47.6)</td>
<td>0.04</td>
</tr>
<tr>
<td><strong>Leucocyte count at baseline (× 10⁹/L), mean (SD)</strong></td>
<td>9.9 (7.7)</td>
<td>9.2 (4.3)</td>
<td>11.4 (11.9)</td>
<td>0.72</td>
</tr>
<tr>
<td><strong>Initiation dose (mg/kg), mean (SD)</strong></td>
<td>1.6 (0.4)</td>
<td>10.3 (4.4)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

SD = standard deviation; M = male; F = female; SLE = systemic lupus erythematosus; FSGS = focal segmental glomerulosclerosis.

*Owing to low frequency of observations, p-values not determined.
†No comparison made between DOC and PIVC owing to different dosing regimens.
The most frequent indication for stopping therapy was the successful completion of induction therapy. This was significantly more frequent in the PIVC group, while cessation of treatment due to treatment failure was more frequent in the DOC group (Table 2). Loss to follow-up, infection and leucopenia were also frequent reasons for discontinuation of treatment in both groups. There were nine patients remaining on CPM at the time the study concluded, while one patient had requested to be transferred to a private physician for further management.

Prescription of prophylactic antimicrobials was consistently achieved, with 91.8% and 94.8% of all participants receiving isoniazid and co-trimoxazole, respectively. There was no difference between the groups. All patients received concomitant prednisone. Only 43.3% of the participants who required contraception were documented to have been prescribed contraceptive therapy.

A total of 64 episodes of infection were documented, of which 37 were considered serious and required hospitalisation. The profile of infections is illustrated in Fig. 1. There were no statistically significant differences in the number of infections (42 v. 22; \( p=0.53 \)) or hospitalisations (25 v. 12; \( p=0.12 \)) between the DOC and PIVC groups.

The most frequent infection was urinary tract infection, followed by pneumonia. The most common opportunistic infection was herpes zoster in the PIVC group and CMV pneumonitis in the DOC group. There were no new cases of TB diagnosed in either group. Infections occurred early during therapy (median 57.5 days) in both groups, with no significant difference between the groups (63.0 v. 55.0 days; \( p=0.72 \)), but occurred at a higher cumulative dose at the time of infection in the DOC group (7 964 mg v. 2 126 mg; \( p<0.01 \)).

Overall, 61.2% of all participants (59.8% of the DOC group and 64.3% of the PIVC group; \( p=0.70 \)) had one or more treatment-related adverse events (infection or leucopenia). The relative risk for developing an adverse event in the DOC group v. the PIVC group was 0.93 (95% confidence interval 0.70 - 1.23).

Over a third (38.1%) of participants developed one or more infections during CPM therapy. In the DOC group, 35.9% of participants developed infections compared with 42.9% in the PIVC group (\( p=0.45 \)). The overall risk of a participant having one or more hospitalisations as a result of serious infection was 24.6%; this was slightly higher in the DOC group (26.1 v. 21.4%; \( p=0.67 \)).

In the DOC group, six participants (6.5%) died due to leucopenia-related sepsis. There were no treatment-related deaths recorded in the PIVC group. This difference was not statistically significant (\( p=0.18 \)). Treatment-related deaths are summarised in Table 3.

Because the frequency of monitoring of leucocyte counts was highly variable, we determined the proportion of participants who became leucopenic at any time during therapy, rather than the leucopenia event rate. The lowest recorded leucocyte counts were significantly lower in the DOC group, with a median of 3.8 v. 5.3 × 10⁹/L (\( p=0.02 \)) (Table 4). Leucopenia also occurred later in the DOC group than in the PIVC group (99 v. 76 days; \( p=0.04 \)) and at higher cumulative doses (mean 3 542 v. 1 929 mg; \( p=0.01 \)). Notably, neutrophil counts were infrequently done.

Comparing the DOC and PIVC groups, 51.1% v. 33.3% (\( p=0.06 \)) of participants developed leucopenia during treatment. The mean duration of leucopenia before the CPM dose was adjusted or the drug stopped was 5 days, with a maximum delay of up to 28 days. However, delays in stopping CPM appeared not to have contributed to any of the leucopenia-related deaths.

**Discussion**

Adverse events due to the use of CPM were frequent in both groups, emphasising...
studies,
occurred early in the course of treatment. Most adverse events in our study, as in other
longer time and may therefore have been more likely to experience
non-significant increased frequency of all infections. DOC was also associated with more frequent and more severe
treatment-related deaths occurred in the DOC group and none in the

In our setting, it appears that PIVC may be safer than DOC. Six
treatment-related deaths occurred in the DOC group and none in the
PIVC group, although this difference was not statistically significant.
DOC was also associated with more frequent and more severe
leucopenia, and with a non-significant increase in hospitalisations
suggesting more serious infections, while PIVC was associated with a
non-significant increased frequency of all infections.

The participants in the DOC group were treated for a significantly
longer time and may therefore have been more likely to experience adverse events. Most adverse events in our study, as in other
studies, occurred early in the course of treatment.

It is important to appreciate that in our study the condition being
treated determined the route of CPM administration and, by default,
also selected for differences in management. Different teams of
clinicians and differences in follow-up and monitoring may therefore have affected the results.

The high cumulative doses used in the DOC group could have
increased efficacy, but are the likely explanation for the increased risk of leucopenia associated with DOC. Another factor was that PIVC
was only administered if there was an acceptable leucocyte count
on the day of each pulse infusion. With DOC, a leucocyte count
was only administered if there was an acceptable leucocyte count
of leucopenia associated with DOC. Another factor was that PIVC
increased efficacy, but are the likely explanation for the increased risk
of leucopenia associated with DOC. Another factor was that PIVC

Table 3. Leucopenia-related deaths

<table>
<thead>
<tr>
<th>Age (years), gender</th>
<th>Diagnosis</th>
<th>Duration of treatment (days)</th>
<th>Lowest leucocyte count (x 10^9/L)</th>
<th>Cause of death</th>
</tr>
</thead>
<tbody>
<tr>
<td>41, F</td>
<td>Mesangiocapillary GN with crescents</td>
<td>59</td>
<td>0.79</td>
<td>Varicella zoster pneumonia</td>
</tr>
<tr>
<td>21, F</td>
<td>SLE with class IV nephritis and myocarditis</td>
<td>21</td>
<td>3.31</td>
<td>Severe pneumonia and renal failure*</td>
</tr>
<tr>
<td>23, F</td>
<td>SLE with class IV nephritis</td>
<td>69</td>
<td>0.54</td>
<td>CMV pneumonitis</td>
</tr>
<tr>
<td>51, F</td>
<td>Crescentic GN</td>
<td>25</td>
<td>0.65</td>
<td>Neutropenic sepsis (unknown source)</td>
</tr>
<tr>
<td>37, F</td>
<td>SLE with class IV nephritis and myocarditis</td>
<td>31</td>
<td>1.58</td>
<td>CMV pneumonitis</td>
</tr>
<tr>
<td>30, F</td>
<td>SLE with class IV nephritis</td>
<td>49</td>
<td>0.34</td>
<td>Overwhelming Acinetobacter baumanii sepsis*</td>
</tr>
</tbody>
</table>

SLE = systemic lupus erythematosus; CMV = cytomegalovirus.
*Subject had renal failure, which significantly contributed to death.
†Leucocyte count at baseline 2.66 x 10^9/L.

the reality that CPM is a potent cytotoxic drug that should be
prescribed with caution and with careful follow-up and monitoring.
To our knowledge this is the first study from a developing country
comparing DOC and PIVC. Other studies have reported leucopenic
episodes in 17 - 26% of PIVC and 19 - 45% of DOC participants and
infection in 10 - 39% of PIVC and 14 - 57% of DOC participants.
Similar infection rates were observed in our study (DOC 36% v. PIVC 43%),
while leucopenia was more frequent (DOC 51% v. PIVC 33%).

In our setting, it appears that PIVC may be safer than DOC. Six
treatment-related deaths occurred in the DOC group and none in the
PIVC group, although this difference was not statistically significant.
DOC was also associated with more frequent and more severe
leucopenia, and with a non-significant increase in hospitalisations
suggesting more serious infections, while PIVC was associated with a
non-significant increased frequency of all infections.

The participants in the DOC group were treated for a significantly
longer time and may therefore have been more likely to experience adverse events. Most adverse events in our study, as in other
studies, occurred early in the course of treatment.

It is important to appreciate that in our study the condition being
treated determined the route of CPM administration and, by default,
also selected for differences in management. Different teams of
clinicians and differences in follow-up and monitoring may therefore have affected the results.

The high cumulative doses used in the DOC group could have
increased efficacy, but are the likely explanation for the increased risk of leucopenia associated with DOC. Another factor was that PIVC
was only administered if there was an acceptable leucocyte count
on the day of each pulse infusion. With DOC, a leucocyte count
was only administered if there was an acceptable leucocyte count
of leucopenia associated with DOC. Another factor was that PIVC

Table 4. Lowest recorded leucocyte count in participants who
developed leucopenia

<table>
<thead>
<tr>
<th>Severity class*</th>
<th>All n (%)</th>
<th>DOC n (%)</th>
<th>PIVC n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (3.01 - 4.00)</td>
<td>20 (32.8)</td>
<td>14 (29.8)</td>
<td>6 (42.9)</td>
</tr>
<tr>
<td>2 (2.01 - 3.00)</td>
<td>13 (21.3)</td>
<td>8 (17.0)</td>
<td>5 (35.7)</td>
</tr>
<tr>
<td>3 (1.01 - 2.00)</td>
<td>13 (21.3)</td>
<td>11 (23.4)</td>
<td>2 (14.3)</td>
</tr>
<tr>
<td>4 (&lt;1.00)</td>
<td>9 (14.8)</td>
<td>8 (17.0)</td>
<td>1 (7.1)</td>
</tr>
<tr>
<td>5 (death)</td>
<td>6 (9.8)</td>
<td>6 (12.8)</td>
<td>0 (0.0)</td>
</tr>
</tbody>
</table>

*Leucocyte count (x 10^9/L) according to Common Terminology Criteria for Adverse Events.

An unexpected finding was that no new episodes of TB were
documented in any participant. This is especially noteworthy
considering the high prevalence of TB of up to 32/1 000 population
in communities serviced by this hospital. This suggests that current
screening protocols for active TB prior to initiation of CPM and the
regular use of isoniazid prophylaxis are effective. Cyclomeagalovirus
(CMV) pneumonitis was a significant opportunistic infection in the
DOC group and a cause of leucopenia-associated death. Patients
on CPM who present with leucopenic sepsis should be screened and
treated early (and probably empirically) for CMV infection.
Current evidence to support prophylactic antiviral treatment for
CMV is lacking, and it is unlikely to be cost-effective in a resource-
constrained setting. Consideration could be given to determining
patients’ CMV status prior to CPM initiation. Knowing the baseline
CMV serology and viral load may aid in the decision whether or
not to treat for suspected CMV disease when patients on CPM
present with leucopenia and signs of infection. However, the cost-
effectiveness of such an approach would need to be determined.

Alarmingly, almost 10% of participants initiated on CPM were
lost to follow-up immediately after discharge from hospital. Another 10% were lost to follow-up later in the study. Considering the poor
prognosis of the underlying conditions if left untreated, it is possible
that many of these patients died as a result of their autoimmune disease.
However, adverse events due to CPM are probably a contributing factor.
The need for excellent record keeping and tracking of patients on CPM
must be emphasised, and clinic staff should be prompted to contact
patients should a follow-up visit be missed.

If resources allow, consideration should be given to the greater use
of PIVC where its use in the treatment of the specific condition in

SLE = systemic lupus erythematosus; CMV = cytomegalovirus.
*Subject had renal failure, which significantly contributed to death.
†Leucocyte count at baseline 2.66 x 10^9/L.
question is supported by current literature. This would be especially useful in cases where compliance with daily medication is in doubt. Owing to the differences in conditions treated, we did not evaluate the efficacy of treatment, which might be the decisive factor in choosing between DOC and PIVC. As pointed out, in many conditions there is a lack of compelling evidence regarding efficacy to preferentially advocate the use of DOC or PIVC.

Study limitations
This study relied on the completeness and accuracy of TBAH's pharmacy database to identify all patients receiving CPM during the study period. The study was retrospective with data extracted from patient records. A heterogeneous group of conditions were included and allocation to PIVC and DOC was not random but determined by the disease process and the preference of the clinicians involved (rheumatology v. nephrology). The primary disease process as well as differences in practices by clinicians are likely to influence treatment outcomes and risk of adverse events. The long-term complications of CPM (e.g. bladder toxicity, infertility) were not studied.

Conclusion
This study has provided new data on adverse events due to CPM in a developing world setting with a very high burden of infectious diseases. We found that a greater proportion of participants receiving DOC developed severe leucopenia, which may be explained by the higher cumulative doses taken over longer periods. There was, however, no significant difference in the proportions of participants who developed infections or required hospitalisation as a consequence. Six participants in the DOC group and none in the PIVC group died as a result of complications of CPM treatment; this difference was not statistically significant.

Local randomised trials are needed to compare PIVC and DOC where follow-up is standardised and where homogeneous groups of conditions are compared. This will provide not only more data with regard to safety, but also much-needed data on differences in treatment outcomes.

Acknowledgements
We thank Ms Radhika Muruvan for help in providing pharmacy data and Dr Justin Harvey from the Stellenbosch University Centre for Statistical Consultation for assistance with data analysis.

References