Evidence-based Series #12-8 Version 2 EDUCATION AND INFORMATION 2015

A Quality Initiative of the Program in Evidence-based Care (PEBC), Cancer Care Ontario (CCO)

Liposomal Anthracyclines in the Management of Patients with HIV-positive Kaposi’s sarcoma

Members of the Systemic Treatment Disease Site Group

An assessment conducted in October 2015 put Evidence-based Series (EBS) 12-8 Version 2 in the Education and Information Section. This means that the recommendations will no longer be maintained but may still be useful for academic or other information purposes. The PEBC has a formal and standardized process to ensure the currency of each document (PEBC Assessment & Review Protocol).

EBS 12-8 Version 2 is comprised of the following 3 sections and is available on the CCO Website on the PEBC Sarcoma page.

Section 1: Clinical Practice Guideline (ENDORSED)
Section 2: Systematic Review
Section 3: Document Review Summary and Review Tool

Release Date: June 12, 2013
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Guideline Report History

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Evidence-based Series #12-8 Version 2: Section 1

Liposomal Anthracyclines in the Management of Patients with HIV-positive Kaposi’s Sarcoma: Guideline Recommendations

N. Iscoe, V. Bramwell, M. Charette, T. Oliver, B. Zanke, and members of the Systemic Treatment Disease Site Group

A Quality Initiative of the Program in Evidence-Based Care (PEBC), Cancer Care Ontario (CCO)
Developed by the Lung Cancer Disease Site Group

These guideline recommendations have been ENDORSED, which means that the recommendations are still current and relevant for decision making. Please see Section 3: Document Review Summary and Tool for a summary of updated evidence published between 2004 and 2012, and for details on how this Clinical Practice Guideline was ENDORSED.

Report Date: January 21, 2013

Guideline Question
Does liposomal anthracycline therapy have advantages over standard combination therapy for patients with human immunodeficiency virus (HIV)-positive Kaposi’s sarcoma who have aggressive cutaneous or visceral disease? Outcomes of interest are survival, time-to-treatment failure, response rates, adverse effects, and quality of life.

Target Population
These recommendations apply to patients with HIV-positive Kaposi’s sarcoma and good performance status (Eastern Cooperative Oncology Group [ECOG] 0-2) who have progressive cutaneous disease despite prior treatment with interferon and/or vinblastine, or who have visceral disease that is symptomatic or progressive.

Recommendations
- The use of conventional combination chemotherapy or single-agent liposomal anthracycline therapy, represent reasonable treatment options in the management of patients with HIV-positive Kaposi’s sarcoma.
Qualifying Statements
Many anti-viral regimens used in the treatment of HIV cause peripheral nerve damage. In patients with HIV-positive Kaposi’s sarcoma, the risk of neuropathic toxicity appears to be greater with vinca alkaloid-containing conventional treatment regimens than with single-agent liposomal anthracyclines. Therefore, if patients have neuropathy, or are at significant risk for neurotoxicity, liposomal anthracycline therapy may be preferable to conventional combination chemotherapy.

Methods
The literature was searched using the MEDLINE (Ovid) (1966 through August 2002), CANCERLIT (Ovid) (1983 through July 2002), and Cochrane Library (Issue 3, 2002) databases. In addition, the Physician Data Query clinical trials database, and abstracts published in the conference proceedings from the meetings of the American Society of Clinical Oncology (1995-2002), and the European Society for Medical Oncology (1998, 2000) were searched for reports of new or ongoing trials. The Canadian Medical Association Infobase and the National Guideline Clearinghouse databases were searched for relevant clinical practice guidelines. Reference lists from relevant articles and reviews were searched for additional trials.

Evidence was selected and reviewed by one member of the Practice Guidelines Initiative’s Systemic Treatment Disease Site Group and methodologists. This practice guideline report has been reviewed and approved by the Systemic Treatment Disease Site Group, which is comprised of medical oncologists, pharmacists, and one community representative.

External review by Ontario practitioners is obtained for all practice guidelines through a mailed survey. Final approval of the practice guideline report is obtained from the Practice Guidelines Coordinating Committee.

The Practice Guidelines Initiative has a formal standardized process to ensure the currency of each guideline report. This process consists of the periodic review and evaluation of the scientific literature and, where appropriate, integration of this literature with the original guideline information.

Update
The original literature search has been updated using MEDLINE (September 2002 through June 2004), EMBASE (September 2002 through June 2004), the Cochrane Library (Issue 2, 2004), the Physician Data Query database, the Canadian Medical Association Infobase, and the National Guideline Clearinghouse, as well as abstracts published in the proceedings of the meetings of the American Society of Clinical Oncology (2004), and the European Society for Medical Oncology (2002). Article bibliographies and personal files were also searched to June 2004 for evidence relevant to this practice guideline report. Please note that CANCERLIT is no longer included in update searches: results from an internal Practice Guidelines Initiative project indicated that the overlap with MEDLINE is 100%, making CANCERLIT database searches redundant.

Key Evidence
- In three published randomized controlled trials, liposomal anthracycline formulations have produced response rates between 25% and 59%, with response rates for the control arm combination chemotherapy regimens ranging from 23% to 28%. In two of these trials, the response rates produced with the liposomal anthracycline formulations were significantly
superior to the control chemotherapy regimens. To date, no statistically significant differences in survival or time-to-treatment failure have been seen.

Future Research
- Patients with HIV-positive Kaposi’s sarcoma should be encouraged to enter clinical trials designed to test therapies aimed at improving survival and quality of life, trials designed to assess whether there are clinically important differences between the available liposomal anthracycline formulations and trials comparing single-agent liposomal anthracyclines with single-agent non-liposomal anthracyclines.
- More information is required to provide better estimates of the risk of cardiotoxicity from liposomal anthracyclines.

Funding
The PEBC is supported by Cancer Care Ontario (CCO) and the Ontario Ministry of Health and Long-Term Care. All work produced by the PEBC is editorially independent from its funding agencies.

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Phone: 905-527-4322 ext. 42822 Fax: 905-526-6775 E-mail: ccopgi@mcmaster.ca
Liposomal Anthracyclines in the Management of Patients with HIV-positive Kaposi’s Sarcoma: A Systematic Review

N. Iscoe, V. Bramwell, M. Charette, T. Oliver, B. Zanke, and members of the Systemic Treatment Disease Site Group

A Quality Initiative of the Program in Evidence-Based Care (PEBC), Cancer Care Ontario (CCO)

These guideline recommendations have been ENDORSED, which means that the recommendations are still current and relevant for decision making.

Please see Section 3: Document Review Summary and Tool for a summary of updated evidence published between 2004 and 2012, and for details on how this Clinical Practice Guideline was ENDORSED.

Section Date: June 2004

I. QUESTION

Does liposomal anthracycline therapy have advantages over standard therapy for patients with HIV-positive Kaposi’s sarcoma who have aggressive cutaneous or visceral disease? Outcomes of interest are survival, time-to-treatment failure, response rates, adverse effects, and quality of life.

II. CHOICE OF TOPIC AND RATIONALE

Kaposi’s sarcoma (KS) is one of many malignancies that can occur with HIV infection. It is a heterogeneous disease, with a wide spectrum of disease manifestations ranging from lesions on isolated areas of the skin (cutaneous KS) to the involvement of internal organs, notably the lungs or gastrointestinal system (visceral KS). Cutaneous KS is an important cause of morbidity with significant impairment of activities of daily living leading to dependency, while visceral KS can be life-threatening (please see Appendix 1 for staging information) (1).

Treatment decisions for patients with KS must take into consideration the extent and rate of tumour growth, symptoms, immune system condition, and concurrent complications of HIV (2). The delivery of effective treatment for KS and the maintenance of adequate control of
HIV and other infections are the current goals in the treatment of this malignancy (2). Treatment with interferon or vinblastine and anti-retroviral agents can be considered for many patients with non-aggressive cutaneous KS (2). The use of combined anti-retroviral therapy has led to a decline in the incidence of KS (3). However, the possibility exists that more KS will develop in these patients if the efficacy of the anti-retroviral therapy fades over time. Radiotherapy is often used in patients with localised cutaneous disease. However, radiotherapy is unlikely to be considered as the preferred form of therapy for the patients with aggressive cutaneous or visceral disease, which is the patient population considered for this guideline.

More aggressive chemotherapy programs, such as various combination chemotherapy regimens, are generally reserved for patients with cutaneous KS resistant to interferon or vinblastine, or for patients with more life-threatening sites of disease. Anthracycline-based chemotherapy, either in single-agent form, or in combination with other drugs such as bleomycin and vincristine, has been used to treat patients with visceral or aggressive cutaneous KS. Anecdotal information suggests that combination chemotherapy with doxorubicin, bleomycin, and vincristine is the initial treatment of choice for patients with aggressive cutaneous or visceral HIV-positive KS. This is the regimen that has been used in the control arm in some of the randomized trials of liposomal anthracyclines reviewed in this report (4,5).

While anthracycline-based chemotherapy produces responses in patients with HIV-positive aggressive cutaneous or visceral KS, it may do so with some degree of toxicity for patients. Moreover, even if these regimens are well-tolerated and produce the desired responses, there are concerns that protracted exposure to the drugs in these regimens will place the patient at risk for long-term refractory organ toxicity. Patients in whom tolerance of anthracyclines is exceeded may experience cardiomyopathy. Treatment with bleomycin may result in lung dysfunction or Raynaud’s phenomenon in some patients. Vincristine use is associated with peripheral neuropathy, something these patients may be predisposed to because of the HIV or anti-retroviral agents used to control their infections. The development of a drug regimen less toxic but equally or more efficacious than current regimens would represent an improvement in the care of these patients.

Liposomal anthracycline agents were developed to deliver drugs to patients in a more selective manner. This action is related to the pharmacodynamics of these agents: they distribute themselves differently in body compartments and tissue compared to the unencapsulated (non-liposomal or free) agent. Theoretically, liposomal anthracyclines offer a therapeutic advantage over the free drug due to their prolonged circulation time and decreased drug-induced toxicity (1,2). The favourable distribution profiles for these agents should theoretically enhance their therapeutic ratios. Liposomal anthracyclines (both doxorubicin and daunorubicin) have been developed and tested in patients with HIV-positive KS in phase III trials. The costs associated with these agents and their high profile in the HIV community motivated the Systemic Treatment Disease Site Group (STDSG) to examine currently available data to determine their potential role in the management of patients with HIV-positive Kaposi’s sarcoma and to develop evidence-based recommendations for their use. The results of these studies are the subject of this report.
III. METHODS
Guideline Development

This practice guideline report was developed by the Practice Guidelines Initiative (PGI), using the methodology of the Practice Guidelines Development Cycle (6). Evidence was selected and reviewed by one member of the PGI’s STDSG and methodologists. Members of the STDSG disclosed potential conflict of interest information.

The practice guideline report is a convenient and up-to-date source of the best available evidence on chemotherapy with liposomal anthracyclines in patients with HIV-positive Kaposi’s sarcoma, developed through systematic reviews, evidence synthesis and input from practitioners in Ontario. The body of evidence in this report is primarily comprised of mature randomized controlled trial data; therefore, recommendations by the DSG are offered. The report is intended to promote evidence-based practice. The Practice Guidelines Initiative is editorially independent of Cancer Care Ontario and the Ministry of Health and Long-Term Care.

External review by Ontario practitioners is obtained for all practice guidelines through a mailed survey consisting of items that address the quality of the draft practice guideline report and recommendations, and whether the recommendations should serve as a practice guideline. Final approval of the original guideline report is obtained from the Practice Guidelines Coordinating Committee (PGCC).

The PGI has a formal standardized process to ensure the currency of each guideline report. This process consists of the periodic review and evaluation of the scientific literature, and where appropriate, integration of this literature with the original guideline information.

Literature Search Strategy

The literature was searched using the MEDLINE (Ovid) (1966 through August 2002), CANCERLIT (Ovid) (1983 through July 2002), and Cochrane Library (Issue 3, 2002) databases. In addition, the Physician Data Query clinical trials database, and abstracts published in the conference proceedings from the meetings of the American Society of Clinical Oncology (1995-2002), and the European Society for Medical Oncology (1998, 2000) were searched for reports of new or ongoing trials. The Canadian Medical Association Infobase and the National Guideline Clearinghouse databases were searched for relevant clinical practice guidelines. Relevant articles and abstracts were selected and reviewed by one member of the STDSG and methodologists, and the reference lists from these sources were searched for additional trials.

The literature search combined the disease specific terms (sarcoma, kaposi/ or kaposi:.tw. and HIV/ or HIV.mp. or HIV infections/ or human immunodeficiency virus.tw. or AIDS/) with treatment specific terms (drug therapy/ or anthracyclines/ or anthracyclines.mp. or liposome:.and doxorubicin.mp. or liposome:.and daunorubicin.mp or doxil.tw. or caelyx.tw. or liposom:.mp. or daunoxome.tw.) with search specific terms for the following study designs: practice guidelines, systematic reviews or meta-analyses, reviews, randomized controlled trials, and clinical trials.

Update

The original literature search has been updated using MEDLINE (September 2002 through June 2004), EMBASE (September 2002 through June 2004), the Cochrane Library (Issue 2, 2004), the Physician Data Query database, the Canadian Medical Association
Infobase, and the National Guideline Clearinghouse, as well as abstracts published in the proceedings of the meetings of the American Society of Clinical Oncology (2004), and the European Society for Medical Oncology (2002). Article bibliographies and personal files were also searched to June 2004 for evidence relevant to this practice guideline report. Please note that CANCERLIT is no longer included in update searches: results from an internal PGI project indicated that the overlap with MEDLINE is 100%, making CANCERLIT database searches redundant.

Inclusion Criteria

Articles were selected for inclusion in this systematic review of the evidence if they met the following criteria:
1. Randomized controlled trials (RCTs) comparing a liposomal anthracycline regimen to observation, placebo or another chemotherapy regimen for the treatment of HIV-positive Kaposi’s sarcoma.
2. Reported data on outcomes of interest including survival, time-to-treatment-failure, response rates, adverse effects, and quality of life.
3. Trials reporting on patients with aggressive cutaneous or visceral HIV-positive KS.

Exclusion Criteria

1. Phase I and II studies were not considered, because of the availability of randomized controlled trials.
2. Letters, editorials, and review articles were not included in this report.
3. Papers published in a language other than English were not considered.
4. Trials including only patients with non-aggressive cutaneous KS were not considered.

Synthesizing the Evidence

The treatment and control arms were different in each of the eligible reviewed trials. The experimental arms of the reviewed trials varied, with three trials using liposomal doxorubicin, and the fourth examining liposomal daunorubicin. The control arms also varied, with the chemotherapy regimens consisting of a combination of doxorubicin, bleomycin, and vincristine in two trials, bleomycin and vincristine in one trial, and liposomal doxorubicin, bleomycin, and vincristine in one trial. Therefore, it was judged inappropriate by the STDSG to pool the data by performing a meta-analysis.

IV. RESULTS

Literature Search Results

A total of five randomized controlled trials (RCTs) were identified (4,5,7-9) in which patients in one of the treatment arms received a liposomal anthracycline.

The randomized trial, by Uthayakumar et al (7), employed a crossover design in which patients with HIV-positive Kaposi’s sarcoma received no therapy and then went on to receive liposomal daunorubicin at the time of disease progression in the observation arm, or 12 weeks later. The patient group was restricted to individuals with non-aggressive cutaneous disease only and therefore did not meet the inclusion criteria as stated. Consequently, this study will not be discussed further.

The randomized trial reported by Mitsuyasu et al (8) is available only in abstract form at this time. In this study, patients with advanced-stage HIV-positive KS who had not received...
prior chemotherapy were randomized to receive either liposomal doxorubicin alone or combined with bleomycin and vincristine. This study is described separately in the Outcomes section, as the treatment regimen is not directly comparable to that of the other eligible trials.

The remaining three studies were randomized controlled trials with sample sizes ranging from 232 to 258 patients (4,5,9). In the study reported by Stewart et al (9), the control arm consisted of a non-anthracycline combination regimen, while a combination anthracycline regimen was used as the control treatment in the studies of Gill (4) and Northfelt (5). Two of the RCTs used liposomal doxorubicin as the liposomal anthracycline (5,9), while the third RCT used liposomal daunorubicin (4). Gill et al (4) excluded patients who had received any prior systemic chemotherapy. Northfelt et al (5) excluded patients if they had received any prior anthracycline chemotherapy, or other chemotherapy within four weeks of entry into the study. Stewart et al (9) excluded patients who had received previous cytotoxic chemotherapy or interferon treatment in the preceding four weeks before entering into the study, or more than one cycle of bleomycin or vincristine at any time. These three studies are described in Table 1, and results are presented in Table 2.

### Table 1. Description of randomized controlled trials of liposomal anthracyclines.

<table>
<thead>
<tr>
<th>Study (ref.)</th>
<th>Patient Population</th>
<th># rand. (# eval.)</th>
<th>Liposomal Anthracycline Regimen</th>
<th>Control Group Description</th>
<th>Outcome Variables</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gill et al, 1996 (4)</td>
<td>advanced KS*</td>
<td>232 (227)</td>
<td>liposomal daunorubicin 40 mg/m² IV every 2 weeks</td>
<td>doxorubicin 10 mg/m², bleomycin 15 U, vincristine 1 mg IV every 2 weeks</td>
<td>response rate, survival, adverse effects, quality of life, time-to-treatment-failure</td>
</tr>
<tr>
<td>Northfelt et al, 1998 (5)</td>
<td>progressive KS†</td>
<td>258 (258)</td>
<td>liposomal doxorubicin 20 mg/m² IV every 2 weeks</td>
<td>doxorubicin 20 mg/m², bleomycin 10 mg/m², vincristine 1 mg IV every 2 weeks</td>
<td>response rate, adverse effects, survival, time-to-treatment-failure</td>
</tr>
<tr>
<td>Stewart et al, 1998 (9)</td>
<td>progressive KS‡</td>
<td>241 (218)</td>
<td>liposomal doxorubicin 20 mg/m² IV every 3 weeks</td>
<td>vincristine 1.4 mg/m², bleomycin 15 mg/m² IV every 3 weeks</td>
<td>response rate, adverse effects, survival, time-to-treatment-failure</td>
</tr>
</tbody>
</table>

NOTE: # eval. = number of evaluable patients; # rand. = number of patients randomized; IV = intravenously; KS = Kaposi’s sarcoma.
* defined as the presence of ≥ 25 mucocutaneous lesions, symptomatic visceral involvement, or the presence of tumour-associated lymphedema.
† defined as progressive HIV-positive KS with at least 25 mucocutaneous lesions, the development of 10 or more new lesions in the preceding month or documented visceral disease.
‡ defined as progressive HIV-positive KS with at least 15 mucocutaneous lesions, the development of more than five cutaneous lesions in the preceding month or documented visceral KS with at least five assessable cutaneous lesions.

### Outcomes

In examining the results for the population of randomized patients, there were no significant differences in median survival time or time-to-treatment-failure between the treatment arms for any of the studies in which these endpoints were measured (Table 2).

In terms of response, Gill et al (4) detected no significant difference in response rates when liposomal daunorubicin was compared with a combination regimen that contained a different anthracycline given at a lower dose intensity (4). In the study reported by Northfelt et al (5), the objective response rate for patients receiving liposomal doxorubicin was 46%
versus 25% for those patients receiving a combination of doxorubicin, bleomycin, and vincristine (ABV) (p<0.001). Stewart et al (9) observed an objective response rate of 59% in patients receiving liposomal doxorubicin versus 23% in patients receiving bleomycin and vincristine (BV) (p<0.001). However, patients in the control arm of this study did not receive an anthracycline as part of their chemotherapy regimen.

The percentage of patients with visceral disease in these studies ranged from 31% in the study by Gill et al (4) to 43% in the study by Stewart et al (9). The study by Northfelt et al (5) did not separate patients with visceral disease from those patients with aggressive cutaneous disease. In the study by Gill et al (4), 45% of the patients with visceral KS randomized to receive liposomal daunorubicin had improvement in visceral disease, with 29% achieving a major response. In the group of patients receiving ABV, 55% of the patients with visceral KS had documented evidence of improvement, with 33% achieving a major response. In this study, survival was significantly improved for patients without visceral involvement at study entry (p=0.0045), independent of treatment. When survival outcome by treatment arm was evaluated separately according to baseline visceral involvement, the difference in median survival was not significant (no data reported). Stewart et al (9) were able to obtain data from 104 patients who had symptoms attributed to visceral KS. Treatment with liposomal doxorubicin decreased the incidence of symptomatic pulmonary KS from 23.1% to 10.6% (p=0.002) and symptomatic gastrointestinal KS from 16.3% to 3.8% (p<0.001). Symptoms of gastrointestinal and pulmonary KS were not found to be significantly reduced in patients treated with BV. If the examination of activity is restricted to the subgroup of patients with visceral KS, it appears there was a pattern consistent with greater improvement in visceral symptoms seen more often with the liposomal agent in the Stewart study (9) that was not repeated in the Gill report (4).

In the Mitsuyasu et al (8) study reported in abstract form, 129 patients with aggressive cutaneous or visceral HIV-positive Kaposi’s sarcoma were randomized to receive either liposomal doxorubicin at a dose of 20 mg/m² every two weeks or liposomal doxorubicin at the same dose combined with vincristine 1 mg and bleomycin 10 U/m² every two weeks. Since both treatment groups received a liposomal anthracycline, the results of this study were not directly comparable with the other reviewed randomized trials (4,5,9). Response rates were similar in both groups, with an objective response rate of 79% for the liposomal doxorubicin arm versus 80% for the combination arm. There were no significant differences between the two treatment groups for time-to-treatment failure or survival.

Table 2. Results of randomized trials of liposomal anthracyclines.

<table>
<thead>
<tr>
<th>Reference</th>
<th># entered (# eval.)</th>
<th>Treatment</th>
<th>Objective response rate (%)*</th>
<th>Time-to-treatment failure (days)</th>
<th>Median Survival (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gill et al, 1996 (4)</td>
<td>117 (116)</td>
<td>lipo daun</td>
<td>25% p=NS</td>
<td>115 99</td>
<td>369 342 p=0.19</td>
</tr>
<tr>
<td></td>
<td>115 (111)</td>
<td>ABV</td>
<td>28% p=NS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Northfelt et al, 1998 (5)</td>
<td>133 (133)</td>
<td>lipo dox</td>
<td>46% p&lt;0.001</td>
<td>124 25 128</td>
<td>160 160 p=NR</td>
</tr>
<tr>
<td></td>
<td>125 (125)</td>
<td>ABV</td>
<td>25%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stewart et al, 1998 (9)</td>
<td>121 (116)</td>
<td>lipo dox</td>
<td>59% p&lt;0.001</td>
<td>160 23 157</td>
<td>NR†</td>
</tr>
<tr>
<td></td>
<td>120 (102)</td>
<td>BV</td>
<td>23%</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NOTE: # = number; A = Adriamycin (doxorubicin); B = bleomycin; daun = daunorubicin; dox = doxorubicin; eval. = evaluable; lipo = liposomal; NS = not significant; NR = not reported; V = vincristine.

* includes complete and partial responses
† this study reported mean survival times of 239 days for liposomal doxorubicin, versus 160 days for bleomycin and vincristine.
Adverse Effects

In the study by Gill et al (4), patients treated with ABV experienced significantly more alopecia (36% versus [v.] 8%; p<0.0001) and neuropathy (41% v. 13%; p<0.0001) of any grade compared with patients treated with liposomal daunorubicin. With respect to hematologic toxicity, the incidence of grade 4 neutropenia was significantly higher in patients treated with liposomal daunorubicin versus patients treated with ABV (15% v. 5%; p=0.021). Sixteen patients treated with liposomal daunorubicin and 31 patients treated with ABV discontinued treatment. Reasons for the discontinuation of treatment with liposomal daunorubicin included death due to complications of HIV infection, patient decision, loss to follow-up evaluation, opportunistic infection, and drug toxicity. The reasons for the discontinuation of ABV were similar, with the addition of intolerable nausea and vomiting, neuropathy, alopecia, and hand-foot syndrome. Thirty-six percent of patients receiving liposomal daunorubicin developed an opportunistic infection versus 26% of patients receiving ABV chemotherapy. This difference was not statistically significant. Seventeen percent of patients in the liposomal daunorubicin group developed neutropenic fever, but no documented infection, compared with 11% of patients in the ABV group. Cardiac events (arrhythmia, palpitations, tachycardia, and hypertension) were observed in 6% of patients receiving liposomal daunorubicin and 9.9% of ABV patients.

In the study by Northfelt et al (5), thirty-seven percent of ABV patients and 11% of liposomal doxorubicin patients discontinued treatment because of an adverse event (p<0.001). One patient who received liposomal doxorubicin died as a result of cardiomyopathy. Eight patients (6%) who received liposomal doxorubicin and three patients (2%) who received ABV experienced episodes of sepsis. Opportunistic infections occurred in 37% of patients treated with liposomal doxorubicin and 30% of patients treated with ABV. The most common adverse event in both groups in the study was leucopenia, but the difference in frequency between the two study arms was not significant. However, there were significant differences between the two arms on other measures of toxicity greater than grade 3. Significantly more patients receiving ABV experienced nausea and/or vomiting (34% v. 15%; p<0.001), alopecia (19% v. 1%; p<0.001), and peripheral neuropathy (14% v. 6%; p=0.002). Mucositis was significantly more common in patients receiving liposomal doxorubicin (5% v. 2%; p=0.026), compared with patients receiving ABV. Three cases of hand-foot syndrome were observed in the liposomal doxorubicin arm versus one in the ABV arm.

Stewart et al (9) reported that the incidence of paresthesia (14% v. 3%; p<0.005), peripheral neuropathy (p<0.001), and constipation (11% v. 2%; p<0.01) were significantly higher in patients who received BV than in patients who received liposomal doxorubicin. The incidence of grade 3 leucopenia (72% v. 51%; p<0.001) and oral candidiasis (29% v. 18%; p<0.05) was significantly higher in patients receiving liposomal doxorubicin. Significantly more patients randomized to receive liposomal doxorubicin experienced an opportunistic infection, compared to patients who were randomized to receive BV (50% v. 30%; p=0.002). Other adverse effects were reported in similar frequencies in the two groups. In this study, patients who received BV were more likely to withdraw from the study prematurely due to a chemotherapy-related event (27%) versus those randomized to receive the liposomal anthracycline (11%).

Only one fully reported study (5) used equimolar doses of an anthracycline and the liposomal agent. In this study, a small and non-significant increase in leucopenia was noted.
for the liposomal arm. In the other studies where increased myelosuppression was noted, it may have been a reflection of the dose of the myelosuppressive agent used in the study.

The use of the other agents in combination with an anthracycline was associated with greater degrees of neuropathy and paresthesia, which appears to have been a contributing factor in discontinuing therapy in at least one study (5). It is important to report that one cardiotoxic death was recorded in a patient receiving liposomal doxorubicin in the Northfelt study (5). This event is consistent with the manufacturer’s statement that the use of the liposomal formulation is not a guarantee against the possibility of anthracycline cardiotoxicity. The impact of the cardioprotectant dexrazoxane on the frequency of cardiotoxicity in patients treated with liposomal anthracycline formulations is unknown.

Quality of Life

Quality of life (QOL) was assessed in the study by Gill et al (4) at each treatment cycle, using the Karnofsky performance status (KPS) score. In addition, a QOL patient questionnaire was completed every other cycle, consisting of questions that encompassed a general health survey, daily activities, treatment-specific symptoms and overall physical and emotional well-being. In this trial, baseline data on KPS and QOL scores were available from over 200 patients. At the end of 20 cycles of treatment, data on KPS scores were available on 11 patients and data on QOL scores were available on 6 patients. There were no statistically significant differences in KPS or QOL scores at any of the time points measured for patients treated with liposomal daunorubicin compared with patients treated with ABV, however, given that the authors of this trial did not provide details on missing data, quality of life results from this trial must be interpreted with caution.

Quality of life was also assessed in the Northfelt study (5). The data related to quality of life were reported in a separate publication (10). Quality of life assessments were carried out using a validated 30-item, self-report, AIDS-modified questionnaire with eleven domains. Baseline data were available on 118 patients in the liposomal doxorubicin arm and 114 patients in the ABV arm, and data at end of treatment were available on over 70% of the treatment population. When the change from baseline to the end of treatment was compared between the two treatment arms, patients receiving liposomal doxorubicin showed significant improvements in four of the eleven domains (general health, pain, social functioning, and energy/fatigue) compared to patients receiving ABV. The domains with the greatest improvement in the liposomal doxorubicin arm compared to the ABV arm were general health and pain.

V. INTERPRETIVE SUMMARY

As previously mentioned, liposomal anthracyclines were developed to be delivered to patients in a more selective manner than standard anthracyclines, theoretically offering a therapeutic advantage due to prolonged circulation time and decreased drug-induced toxicity. However, as none of the identified randomized trials included a control arm of single-agent anthracycline therapy, it is difficult to determine any incremental benefit of liposomal agents over conventional anthracyclines alone in terms of efficacy, toxicity or quality of life. While this is an important area for future research, the focus of this report remains on the currently available evidence of three randomized trials of single-agent liposomal anthracyclines compared with combination chemotherapy containing vincristine and bleomycin with or without doxorubicin.
In three published randomized controlled trials, liposomal anthracycline formulations produced response rates between 25% and 59%, compared with 23% to 28% for the control arm combination chemotherapy regimens. In two of these trials, the response rates were significantly superior with liposomal doxorubicin versus combination chemotherapy. Of these two trials, one trial included an anthracycline in the control arm while the other did not. No statistically significant differences in median survival or time-to-treatment-failure were detected in any of the three trials.

In terms of adverse events, rates of severe toxicity and opportunistic infection appear to be roughly equivalent between liposomal anthracycline therapy and conventional chemotherapy. However, it is clear that vincristine and bleomycin contribute significantly to toxicity, notably neurotoxicity. Therefore, if patients have neuropathy, or are at significant risk for neurotoxicity, the use of a liposomal anthracycline agent is a very attractive alternative to the commonly used combination regimen of doxorubicin, bleomycin, and vincristine. While not all patients with Kaposi’s sarcoma develop neurotoxicity on conventional chemotherapy, many are on anti-retroviral regimens that may cause peripheral nerve damage, and many develop signs and symptoms of neurotoxicity as a result of these therapies.

Of the two randomized trials that report data on quality of life, one trial did not detect any significant differences in quality of life measures for patients in either treatment arm. Evidence from the other trial supports that aspects of quality of life are significantly better when patients are treated with liposomal anthracycline therapy compared to conventional combination therapy. However, it is unclear to what extent the changes described are clinically meaningful.

Based on this limited available information, the use of liposomal therapy or conventional combination therapy represents equally valid approaches in the treatment of patients with HIV-positive Kaposi’s sarcoma. Patients should be informed of the harms and benefits associated with each treatment regimen and patient preference should be taken into account when making treatment decisions.

If a liposomal agent is to be used, there is insufficient information available to decide if one agent is superior to the other, or if liposomal anthracyclines are better than single-agent anthracycline therapy alone. These would be fruitful avenues for future research.

VI. ONGOING TRIALS

The STDSG is aware of the following ongoing trials evaluating liposomal anthracyclines in patients with Kaposi’s sarcoma:

RPCI-DS-96-28, NCI-G97-1241, SEQUUS-30-38: Phase III randomized study of liposomal doxorubicin in patients with AIDS-related Kaposi’s sarcoma (11). Patients will be randomly assigned to receive liposomal doxorubicin or liposomal daunorubicin in a 3:1 ratio. Eighty patients will be studied to determine tumour response, safety and clinical benefit of liposomal doxorubicin. Preliminary results of this trial have been reported in abstract form (12). The Systemic Treatment DSG will monitor the literature for mature results from this trial.

E-1D96: Phase III randomized study of paclitaxel versus liposomal doxorubicin in patients with advanced AIDS-associated Kaposi’s sarcoma (13). Two hundred and forty patients will be
accrued and randomized to receive either paclitaxel or liposomal doxorubicin. Progression-free survival, quality of life, toxicity, and response rates will be measured. The summary was last modified on the PDQ web site in July 2002.

VII. DISEASE SITE GROUP CONSENSUS PROCESS

A preliminary draft of this practice guideline report was circulated to the members of the STDSG for comment. The discussions at the DSG meetings highlighted the need to identify the patient group to whom this guideline was directed. The discussion also focused at some length on the interpretation of the data. Special care was taken to ensure that the information was conveyed in a manner that would be helpful to practitioners. As a result of these discussions, the initial draft of the practice-guideline-in-progress was modified. The modified version was recirculated to the STDSG for further comments before being sent for feedback from physicians involved in the care of patients with HIV-positive Kaposi’s sarcoma.

VIII. EXTERNAL REVIEW OF THE PRACTICE GUIDELINE REPORT

Draft Recommendations

Based on the evidence above, the STDSG drafted the following recommendations:

Target Population

These recommendations apply to patients with HIV-positive Kaposi’s sarcoma and good performance status (ECOG 0-2) who have progressive cutaneous disease despite prior treatment with interferon and/or vinblastine, or who have visceral disease that is symptomatic or progressive.

Recommendations

Key recommendations

• The first choice of therapy for these patients should be conventional anthracycline regimens. However, in circumstances where the risk of toxicity from standard chemotherapy is likely to compromise a patient’s health, a liposomal anthracycline represents an appropriate alternative treatment choice.

Qualifying statements

• Many anti-viral regimens used in the treatment of HIV cause peripheral nerve damage, and the risk of neuropathic toxicity appears to be greater with vinca alkaloid-containing conventional treatment regimens than with single-agent liposomal anthracyclines in patients with HIV-positive Kaposi’s sarcoma. Therefore, liposomal anthracycline formulations represent a reasonable alternative to currently available chemotherapy regimens for patients with pre-existing neuropathy or those at high risk of neuropathy.

Future Research

• Patients with HIV-positive Kaposi’s sarcoma should be encouraged to enter clinical trials designed to test therapies aimed at improving survival and quality of life, trials designed to assess whether there are clinically important differences between the available liposomal anthracycline formulations and trials comparing single-agent liposomal anthracyclines with single-agent non-liposomal anthracyclines.

• More information is required to provide better estimates of the risk of cardiotoxicity from liposomal anthracyclines.
Practitioner Feedback
Based on the evidence and the draft recommendations presented above, feedback was sought from Ontario clinicians.

Methods
Practitioner feedback was obtained through a mailed survey of nine practitioners in Ontario (seven medical oncologists and two hematologists). The survey consisted of 21 items evaluating the methods, results, and interpretive summary used to inform the draft recommendations outlined and whether the draft recommendations above should be approved as a practice guideline. Written comments were invited. Follow-up reminders were sent at two weeks (post card) and four weeks (complete package mailed again). The results of the survey have been reviewed by the Systemic Treatment Disease Site Group.

Results
Key results of the practitioner feedback survey are summarized in Table 3. Six (67%) surveys were returned. Six (100%) respondents indicated that the practice-guideline-in-progress report was relevant to their clinical practice and completed the survey.

Summary of Main Findings
Two (33%) respondents provided written comments. The main points were:
1. One respondent asked whether the KS that might return after the current anti-retroviral therapies fail will be clinically the same as that treated in the studies reported, or whether the results of this guideline will be relevant to those events with the testing of new therapies.
2. A second respondent noted there was not strong support for the liposomal formulations in terms of survival and asked about quality of life benefits.

Modifications/Actions
1. While this may be true, there is merit in having a guideline for the current cohort of patients. No changes were made to the document.
2. The current guideline does not support the use of liposomal anthracyclines on the basis of survival enhancement, but does note the potential for benefit in a subset of patients at risk for complications from conventional combination chemotherapy, based on the reports of increased neurotoxicity for these treatments (9). No changes were made to the document.

Table 3. Practitioner responses to eight items on the practitioner feedback survey.

<table>
<thead>
<tr>
<th>Item</th>
<th>Number (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>The rationale for developing a clinical practice guideline, as stated in the “Choice of Topic” section of the report, is clear.</td>
<td>6 (100) 0 0</td>
</tr>
<tr>
<td>There is a need for a clinical practice guideline on this topic.</td>
<td>5 (83) 1 (17) 0</td>
</tr>
<tr>
<td>The literature search is relevant and complete.</td>
<td>5 (83) 0 0</td>
</tr>
</tbody>
</table>
The results of the trials described in the report are interpreted according to my understanding of the data.  

<table>
<thead>
<tr>
<th>Statement</th>
<th>Yes</th>
<th>No</th>
<th>Unsure</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>The results of the trials described in the report are clear.</td>
<td>6</td>
<td>0</td>
<td>0</td>
<td>6</td>
</tr>
<tr>
<td>I agree with the draft recommendations as stated.</td>
<td>5</td>
<td>0</td>
<td>1 (17)</td>
<td>6</td>
</tr>
<tr>
<td>This report should be approved as a practice guideline.</td>
<td>3 (50)</td>
<td>1 (17)</td>
<td>0</td>
<td>4 (67)</td>
</tr>
</tbody>
</table>

If this report were to become a practice guideline, how likely would you be to make use of it in your own practice?  

<table>
<thead>
<tr>
<th>Likelihood</th>
<th>Yes</th>
<th>No</th>
<th>Unsure</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Very likely or likely</td>
<td>4</td>
<td>1</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Unsure</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Not at all likely or unlikely</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

NOTE: Some percentages do not add to 100 because of missing data.

Practice Guidelines Coordinating Committee Approval Process

The practice guideline report was circulated to members of the Practice Guidelines Coordinating Committee (PGCC) for review and approval. Nine of eleven members of the PGCC returned ballots. Five PGCC members approved the practice guideline report as written, and four members approved the guideline conditional on the Systemic treatment DSG addressing specific concerns. PGCC members requested that the following issues be addressed prior to the approval of the guideline report: minor typographical errors and wording changes; more information on the differences between conventional and liposomal anthracyclines as well as an explicit description of the available evidence; a discussion of the importance of intermediate markers when important outcomes do not differ; and a more complete rationale for recommending, with qualifications, conventional anthracyclines as the preferred treatment option.

 Modifications/Actions

Based on the comments of the members of the PGCC, the Systemic Treatment DSG modified the practice guideline report to address the above issues. As a result, changes to the interpretive summary, recommendations, and qualifying statements were made.

IX. PRACTICE GUIDELINE

These practice guideline recommendations reflect the integration of the draft recommendations with feedback obtained from the external review process. It has been approved by the Systemic Treatment DSG and the Practice Guidelines Coordinating Committee.

Target Population

These recommendations apply to patients with HIV-positive Kaposi’s sarcoma and good performance status (ECOG 0-2) who have progressive cutaneous disease despite prior treatment with interferon and/or vinblastine, or who have visceral disease that is symptomatic or progressive.

Recommendations

- The use of conventional combination chemotherapy or single-agent liposomal anthracycline therapy, represent reasonable treatment options in the management of patients with HIV-positive Kaposi’s sarcoma.

Qualifying Statements

- Many anti-viral regimens used in the treatment of HIV cause peripheral nerve damage. In patients with HIV-positive Kaposi’s sarcoma, the risk of neuropathic toxicity appears to be
greater with vinca alkaloid-containing conventional treatment regimens than with single-agent liposomal anthracyclines. Therefore, if patients have neuropathy, or are at significant risk for neurotoxicity, liposomal anthracycline therapy may be preferable to conventional combination chemotherapy.

Future Research

- Patients with HIV-positive Kaposi's sarcoma should be encouraged to enter clinical trials designed to test therapies aimed at improving survival and quality of life, trials designed to assess whether there are clinically important differences between the available liposomal anthracycline formulations and trials comparing single-agent liposomal anthracyclines with single-agent non-liposomal anthracyclines.

- More information is required to provide better estimates of the risk of cardiotoxicity from liposomal anthracyclines.

X. POLICY IMPLICATIONS

Currently there is little information available about the number of patients who might be candidates for the liposomal anthracycline therapy. It is clear from reports in the literature and the experience of physicians involved in the care of these patients that the number of patients with HIV-positive Kaposi's sarcoma has decreased markedly in the last few years. This has generally paralleled the improvements in HIV therapy. The result is that there is a large pool of potential patients who might develop progressive HIV and associated diseases, including Kaposi's sarcoma. Consequently, the demand for the liposomal anthracycline agents, while at present likely to be limited, could expand if the incidence of progressive HIV and its related conditions were to rise. Additionally, the distribution of these patients may be uneven in various treatment centres around the province. In order to spread the burden of cost related to these agents in an equitable fashion, we believe reimbursement for these agents should be through the provincial program and be subject to meeting criteria for use. A community representative on the DSG strongly believed that the access to these agents should be through the provincial program.

Table 4 outlines the cost per week (in Canadian dollars) for treating an average patient with either a liposomal anthracycline regimen or a combination regimen of doxorubicin, bleomycin, and vincristine. The acquisition costs reflect only a component of the costs of delivering therapy. Costs associated with pharmacy workload and chemotherapy administration need to be considered, but are beyond the scope of this report. If liposomal anthracycline therapy is being considered, the direct cost of liposomal daunorubicin appears more attractive to that of liposomal doxorubicin. However, for an accurate comparison, detailed cost-effectiveness analyses based on Canadian data are needed. Unfortunately no such analyses were identified in the literature. Two cost-effectiveness analyses, one Swedish by Hjortsberg et al (14) and the other American by Bennett et al (15) were identified. The authors computed the projected cost (in American dollars) of the two liposomal formulations to achieve similar rates of response as reported in two of the identified randomized trials (4,9). The authors concluded that despite higher acquisition costs, the costs for liposomal doxorubicin were actually much lower than those for liposomal daunorubicin. Given the current price differences between the two liposomal formulations, the results of the ongoing randomized trial (12) comparing the two formulations will hopefully clarify the relative merits of the two agents.
Table 4. Cost per m² per week based on treating an average patient.

<table>
<thead>
<tr>
<th>Chemotherapeutic agent</th>
<th>Format</th>
<th>Acquisition cost</th>
<th>Dose schedule</th>
<th>Cost / cycle (for a person 1 m²)</th>
<th>Unit cost per week</th>
</tr>
</thead>
<tbody>
<tr>
<td>liposomal doxorubicin</td>
<td>20 mg/ml</td>
<td>$683.00</td>
<td>20 mg/m² every 3 weeks</td>
<td>$683.00</td>
<td>$227.67</td>
</tr>
<tr>
<td>liposomal daunorubicin</td>
<td>50 mg/20 ml</td>
<td>$315.00</td>
<td>40 mg/m² every 2 weeks</td>
<td>$252.00</td>
<td>$126.00</td>
</tr>
<tr>
<td>doxorubicin</td>
<td>200 mg/100 ml</td>
<td>$1019.48</td>
<td>20 mg/m² every 2 weeks</td>
<td>$101.95</td>
<td>$50.98</td>
</tr>
<tr>
<td>bleomycin</td>
<td>15 U/ml</td>
<td>$201.16</td>
<td>10 U/m² every 2 weeks</td>
<td>$134.11</td>
<td>$67.06</td>
</tr>
<tr>
<td>vincristine</td>
<td>5 mg/5 ml</td>
<td>$84.50</td>
<td>1 mg every 2 weeks</td>
<td>$16.90</td>
<td>$8.45</td>
</tr>
<tr>
<td>ABV (combination of above three agents)</td>
<td></td>
<td></td>
<td></td>
<td>$50.98 + $67.06 + $8.45 = $126.49</td>
<td></td>
</tr>
</tbody>
</table>

NOTE: A = Adriamycin (doxorubicin); B = bleomycin; V = vincristine.

XI. JOURNAL REFERENCE

XII. ACKNOWLEDGMENTS
The Systemic Treatment Disease Site Group would like to thank Dr. Neill Iscoe, Dr. Vivien Bramwell, Ms. Manya Charette, and Mr. Tom Oliver for taking the lead in drafting and revising this practice guideline report.

*For a full list of members of the Breast Cancer Disease Site Group, please visit the CCO Web site at [http://www.cancercare.on.ca/](http://www.cancercare.on.ca/)*
REFERENCES

Appendix 1: Staging classification for Kaposi’s sarcoma.

<table>
<thead>
<tr>
<th></th>
<th>Good Risk (0)</th>
<th>Poor Risk (1)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Tumour (T)</strong></td>
<td>Confined to skin and/or lymph nodes and /or minimal oral disease*</td>
<td>Tumour-associated edema or ulceration Extensive oral KS Gastrointestinal KS KS in other non-nodal viscera</td>
</tr>
<tr>
<td><strong>Immune System (I)</strong></td>
<td>CD4 cells ≥ 200/µL</td>
<td>CD4 cells &lt; 200/µL</td>
</tr>
<tr>
<td><strong>Systemic illness (S)</strong></td>
<td>No history of opportunistic infection or thrush No “B” symptoms** Performance status ≥ 70 (Karnofsky)</td>
<td>History of opportunistic infections and/or thrush “B” symptoms present Performance status &lt; 70 Other HIV-related illness (e.g., neurological disease, lymphoma)</td>
</tr>
</tbody>
</table>

* Minimal oral disease in non-nodular KS confined to the palate.
** “B” symptoms are unexplained fever, night sweats, > 10% involuntary weight loss, or diarrhea persisting more than two weeks.

Evidence-based Series #12-8: Section 3

A Quality Initiative of the Program in Evidence-Based Care (PEBC), Cancer Care Ontario (CCO)

Liposomal Anthracyclines in the Management of Patients with HIV-positive Kaposi’s Sarcoma: Guideline Summary Review

S. Verma, N.P. Varela and Members of the Systemic Treatment Disease Site Group

Review Date: January 21, 2013

The 2004 guideline recommendations are ENDORSED

This means that the recommendations are still current and relevant for decision making.

OVERVIEW
Evidence-based Series History
This guidance document was originally released by Cancer Care Ontario’s Program in Evidence-based Care in 2004. In September 2012, the PEBC guideline update strategy was applied and the new document released in June 2013. The recommendations and the systematic review in this version are the same as June 2004 version.

Update Strategy
Using the Document Review Tool, the PEBC update strategy includes an updated search of the literature, review and interpretation of the new eligible evidence by clinical
DOCUMENT ASSESSMENT AND REVIEW RESULTS

Questions Considered

Does liposomal anthracyclines therapy have advantages over standard therapy for patients with HIV-positive Kaposi’s sarcoma who have aggressive cutaneous or visceral disease? Outcomes of interest are survival, time-to-treatment failure, response rates, adverse effects, and quality of life.

Literature Search and New Evidence

The new search (Jan 2004 to Aug 2012) yielded 3 relevant new publications representing one randomized control trial, one ongoing trial and a research letter. Brief results of these publications are shown in the Document Review Tool at the end of this report.

Impact on Guidelines and Its Recommendations

The new data supports existing recommendations. Hence, the Systemic Treatment DSG ENDORSED the 2004 recommendations on Liposomal Anthracyclines in the Management of Patients with HIV-positive Kaposi’s Sarcoma.

Document Summary and Review Tool

<table>
<thead>
<tr>
<th>Number and title of document under review</th>
<th>12-8 Liposomal Anthracyclines in the Management of patients with HIV-positive Kaposi’s Sarcoma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current Report Date</td>
<td>June 2004</td>
</tr>
<tr>
<td>Clinical Expert</td>
<td>Dr. Shailendra Verma</td>
</tr>
<tr>
<td>Research Coordinator</td>
<td>Norma P. Varela</td>
</tr>
<tr>
<td>Date Assessed</td>
<td>September 2011</td>
</tr>
<tr>
<td>Approval Date and Review Outcome (once completed)</td>
<td>January 21, 2013 [ENDORSED]</td>
</tr>
</tbody>
</table>

Original Question(s):
Does liposomal anthracyclines therapy have advantages over standard therapy for patients with HIV-positive Kaposi’s sarcoma who have aggressive cutaneous or visceral disease? Outcomes of interest are survival, time-to-treatment failure, response rates, adverse effects, and quality of life.

Target Population:
These recommendations apply to patients with HIV-positive Kaposi’s sarcoma and good performance status (Eastern Cooperative Oncology Group [ECOG] 0-2) who have progressive cutaneous disease despite prior treatment with interferon and/or vinblastine, or who have visceral disease that is symptomatic or progressive.

Study Section Criteria:

Inclusion Criteria:
Articles were selected for inclusion in this systematic review of the evidence if they met the following criteria:

1. Randomized controlled trials (RCTs) comparing a liposomal anthracycline regimen to observation, placebo or another chemotherapy regimen for the treatment of HIV-positive kaposi’s sarcoma.
2. Reported data on outcomes of interest including survival, time-to-treatment-failure, response rates, adverse effects, and quality of life.
3. Trials reporting on patients with aggressive cutaneous or visceral HIV-positive KS.

**Exclusion Criteria:**

1. Phase I and II studies were not considered, because of the availability of randomized controlled trials.
2. Letters, editorials, and review articles were not included in this project.
3. Papers published in a language other than English were not considered.
4. Trials including only patients with non-aggressive cutaneous KS were not considered.

**Search Details:**

January 2004 to August 2012 (Medline, Embase, Cochrane Library, PDQ Clinical Trials, ESMO Clinical Trials, ASCO Annual Meetings, and SAGE Cancer Guidelines).

**Brief Summary/Discussion of New Evidence:**

Of 26 hits from Medline and Embase + 5 from The Cochrane Library + 5 from SAGE Cancer Guidelines and Standards + 33 from ASCO Conference abstracts + 20 ongoing trials, 2 references (1 RCT and 1 ongoing trial) were found. In addition, a research letter (regardless of exclusion criteria in original Guideline) was included because of the lack of randomized clinical trials assessing the efficacy of liposomal anthracycline for the treatment of HIV-positive Kaposi’s sarcoma.

### Randomized Controlled Trials

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Population</th>
<th>Median follow up</th>
<th>Outcomes</th>
<th>Brief Results</th>
<th>References</th>
</tr>
</thead>
</table>
| Paclitaxel (Taxol) vs PLD (Doxil) | HIV-infected patients with advanced symptomatic Kaposi’s sarcoma. *Median age:*  
- Paclitaxel: 39 yrs.  
- PLD: 36 yrs. | 36 months [range, 0.9 to 90.9] | OR, PFS, OS, AE | *The overall response rate was 56% for the paclitaxel arm (CR = 8%) and 46% for the PLD arm (CR = 5%) (No statistically significant difference [p = 0.486]).*  
*The median PFS was 17.5 months in the paclitaxel arm and 12.2 months in the PLD arm (No statistically significant difference [p = 0.66]).*  
*The median overall survival for the paclitaxel arm was 53.6 months, and that for the PLD arm had not been reached at the time of last follow-up.*  
*The 2-year overall survival rate was 79% in the paclitaxel arm and 78% in the PLD arm (No statistically significant difference [p = 0.748]).*  
*The overall incidence if grade 3 or greater toxicity was somewhat higher in the paclitaxel arm (84% vs 66% [p = 0.077]), including neutropenia (58% vs | Cianfrocca et al., 2010 |
41% \( p = 0.184 \)).

- Grade 1 to 2 alopecia (57.8% vs 11% \( p = 0.001 \)) and sensory neuropathy (26% vs 9% \( p = 0.045 \)) were found to be significantly more common in the paclitaxel arm when compared to the PLD arm.
- Infection rates were similar (16% and 14% for paclitaxel and PLD arm, respectively), and there was 1 toxicity grade 5 in the paclitaxel arm (patient died of a pulmonary embolism on Day9).

---

### Research Letter Reporting a Randomized Controlled Trial

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Population</th>
<th>Outcomes</th>
<th>Brief Results</th>
<th>References</th>
</tr>
</thead>
</table>
| PDL + HAART vs HAART alone | HIV-infected patients with moderate-advanced Kaposi’s sarcoma | • Intent to treat  
• On-treatment  
• Complete remission  
• Partial Remission | Better response rates were observed in the HAART+PLD arm when compared to the HAART arm:  
• Intent-to-treat: 76% and 20% for HAART+PLD and HAART alone, respectively \( p = 0.003 \).  
• On-treatment: 91% and 23% for HAART+PLD and HAART alone, respectively \( p = 0.0001 \).  
• Complete Remission: 31% and 13% for HAART+PLD and HAART, respectively  
• Partial Remission: 46% and 7% for HAART+PLD and HAART, respectively. | Martin-Carbonero et al., 2004 |

### Ongoing Randomized Controlled Trials (RCTs)

<table>
<thead>
<tr>
<th>Interventions</th>
<th>Official Title</th>
<th>Status</th>
<th>Protocol ID</th>
<th>Last Updated</th>
<th>Estimated Completion Date</th>
</tr>
</thead>
<tbody>
<tr>
<td>ART + ET vs ART + BV vs ART + PLD</td>
<td>A Randomized Comparison of Three Regimens of Chemotherapy with Compatible Antiretroviral Therapy for Treatment of Advanced AIDS-KS in Resource-Limited Settings</td>
<td>Not yet recruiting</td>
<td>NCT01435018</td>
<td>May 30, 2012</td>
<td>September 2019</td>
</tr>
</tbody>
</table>

PLD (Doxorubicin HCL Liposome); OR (Overall Response); CR (Complete Response); PFS (Progression-Free Survival); OS (Overall Survival); AE (Adverse Effects); QOL (Quality of Life); HAART (highly active antiretroviral therapy); ART (Antiretroviral); ET (Etoposide); BV (Bleomycin and Vincristine).

**Clinical Expert Interest Declaration:**

No conflict of interest to declare.
1. Does any of the newly identified evidence, on initial review, contradict the current recommendations, such that the current recommendations may cause harm or lead to unnecessary or improper treatment if followed?  
   No.

2. On initial review,  
   a. Does the newly identified evidence support the existing recommendations?  
      YES  
   b. Do the current recommendations cover all relevant subjects addressed by the evidence, such that no new recommendations are necessary?  
      YES

3. Is there a good reason (e.g., new stronger evidence will be published soon, changes to current recommendations are trivial or address very limited situations) to postpone updating the guideline?  
   Answer Yes or No, and explain if necessary:  
   NO

4. Do the PEBC and the DSG/GDG responsible for this document have the resources available to write a full update of this document within the next year?  
   If necessary

**Review Outcome**
No change to existing guideline.

**DSG/GDG Approval Date**
January 21, 2013

**DSG/GDG Commentary**
Although there is improvement in PLD +HAART comparing with HAART alone the control is not a standard control and data is not fully published.

**New References Identified**


Literature Search Strategy

Kaposi OR Kaposi’s

Cochrane Library
(Kaposi OR kaposis) AND (liposomal anthracyclines OR doxil OR caelyx OR myocet OR daunoxome OR doxorubicin OR daunorubicin) AND (HIV OR AIDS OR immunodeficiency)

Medline
1. meta-Analysis as topic.mp.
2. meta analysis.pt.
3. (meta analy$ or metaanaly$).tw.
4. (systematic review$ or pooled analy$ or statistical pooling or mathematical pooling or statistical summar$ or mathematical summar$ or quantitative synthes?s or quantitative overview).tw.
5. (systematic adj (review$ or overview?)).tw.
6. (exp Review Literature as topic/ or review.pt. or exp review/) and systematic.tw.
7. or/1-6
8. (cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or scisearch or bids or sigle or cancerlit).ab.
9. (reference list$ or bibliograph$ or hand-search$ or relevant journals or manual search$).ab.
10. (selection criteria or data extraction or quality assessment or jadad scale or methodological quality).ab.
11. (study adj selection).ab.
12. 10 or 11
13. review.pt.
14. 12 and 13
15. exp randomized controlled trials as topic/ or exp clinical trials, phase III as topic/ or exp clinical trials, phase IV as topic/
16. (randomized controlled trial or clinical trial, phase III or clinical trial, phase IV).pt.
17. random allocation/ or double blind method/ or single blind method/
18. (randomi$ control$ trial? or rct or phase III or phase IV or phase 3 or phase 4).tw.
19. or/15-18
20. (phase II or phase 2).tw. or exp clinical trial/ or exp clinical trial as topic/
21. (clinical trial or clinical trial, phase II or controlled clinical trial).pt.
22. (20 or 21) and random$.tw.
23. (clinic$ adj trial$1).tw.
24. ((singl$ or doubl$ or treb$ or tripl$) adj (blind$3 or mask$3 or dummy)).tw.
25. placebos/
26. (placebo? or random allocation or random alllocated or allocated randomly).tw.
27. (allocated adj2 random).tw.
28. or/23-27
29. practice guidelines/
30. practice guideline?.tw.
31. practice guideline.pt.
32. or/29-31
33. 7 or 8 or 9 or 14 or 19 or 22 or 28 or 32
34. (comment or letter or editorial or note or erratum or short survey or news or newspaper article or patient education handout or case report or historical article).pt.
35. 33 not 34
36. limit 35 to english
37. Animal/
38. Human/
39. 37 not 38
40. 36 not 39
41. exp kaposi's sarcoma/
42. (HIV or HIV infection? or human immunodeficiency virus or AIDS).tw.
43. 41 and 42
44. ((liposome$ adj anthracycline$) or drug therapy or doxorubicin or daunorubicin$ or doxil or caelyx or daunoxencene or myocet).tw.
45. 43 and 44
46. 40 and 45
48. 46 and 47

**Embase**
1. exp meta analysis/ or exp systematic review/
2. (meta analy$ or metaanaly$).tw.
3. (systematic review$ or pooled analy$ or statistical pooling or mathematical pooling or statistical summar$ or matematical summar$ or quantitative synthesis$ or quantitative overview).tw.
4. (systematic adj (review$ or overview?)).tw.
5. exp review/ or review.pt.
6. (systematic or selection criteria or data extraction or quality assessment or jadad scale or methodological quality).ab.
7. (study adj selection).ab.
8. 5 and (6 or 7)
9. or/1-4,8
10. (cochraine or embase or psychlit or psyclit or psychinfo or psycinfo or cinhal or science citation index or scisearch or bids or single or cancercit).ab.
11. (reference list$ or bibliography$ or hand-search$ or relevant journals or manual search$).ab.
12. exp randomized controlled trial/ or exp phase 3 clinical trial/ or exp phase 4 clinical trial/
13. randomization/ or single blind procedure/ or double blind procedure/
14. (randomi$ control$ trial? or rct or phase III or phase IV or phase 3 or phase 4).tw.
15. or/12-14
16. (phase II or phase 2).tw. or exp clinical trial/ or exp prospective study/ or exp controlled clinical trial/
17. 16 and random.tw.
18. (clinic$ and trial$1).tw.
19. ((singl$ or doubl$ or tre$ or tripl$) adj (blind$3 or mask$3 or dummy)).tw.
20. placebo/
21. (placebo? or random allocation or random allocated or allocated randomly).tw.
22. (allocated adj2 random).tw.
23. or/18-22
24. practice guidelines/
25. practice guideline?.tw.
27. or/24-26
28. 9 or 10 or 11 or 15 or 17 or 23 or 27
29. (editorial or note or letter or erratum or short survey).pt. or abstract report/ or letter/ or case study/
30. 28 not 29
31. limit 30 to english
32. Animal/
33. Human/
34. 32 not 33
35. 31 not 34
36. exp kaposi’s sarcoma/
37. (HIV or HIV infection? or human immunodeficiency virus or AIDS).tw.
38. 36 and 37
39. ((liposom$ adj anthracycline$) or drug therapy or doxorubicin or daunorubicin$ or doxil or caelyx or daunoxene or myocet).tw.
40. 38 and 39
41. 35 and 40
43. 41 and 42

ASCO Annual Meeting - [http://www.ascopubs.org/search](http://www.ascopubs.org/search)
Kaposi’s AND HIV

Kaposi AND HIV AND (anthracyclines OR doxil OR caelyx OR myocet OR daunoxome OR doxorubicin OR daunorubicin)

OUTCOMES DEFINITION

1. ARCHIVED - An archived document is a document that will no longer be tracked or updated but may still be useful for academic or other informational purposes. The document is moved to a separate section of our website, each page is watermarked with the phrase “ARCHIVED”.

Section 3: Document Review Summary and Review Tool
2. **ENDORSED** - An endorsed document is a document that the DSG/GDG has reviewed for currency and relevance and determined to be still useful as guidance for clinical decision making. A document may be endorsed because the DSG/GDG feels the current recommendations and evidence are sufficient, or it may be endorsed after a literature search uncovers no evidence that would alter the recommendations in any important way.

3. **DELAY** - A delay means that there is reason to believe new, important evidence will be released within the next year that should be considered before taking further action.

4. **UPDATE** - An Update means that the DSG/GDG recognizes that there is new evidence that makes changes to the existing recommendations in the guideline necessary but these changes are more involved and significant than can be accomplished through the Document Assessment and Review process. The DSG/GDG will rewrite the guideline at the earliest opportunity to reflect this new evidence. Until that time, the document will still be available as its existing recommendations are still of some use in clinical decision making.