**CASE REPORT**

**Successful treatment of AIDS-related pulmonary Kaposi's sarcoma with liposomal daunorubicin**

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Pulmonary Kaposi's sarcoma (KS) is a life-threatening complication in patients with the acquired immunodeficiency syndrome (AIDS). The prognosis is poor, with a median survival after diagnosis of 4–10 months [1–3]. The response to chemotherapy is usually limited, and current treatment protocols are associated with considerable toxicity [3, 4].

The use of targeted liposomes as a carrier for cytotoxic drugs may reduce the toxic effects on normal tissue, whilst increasing drug concentrations and effectiveness at the tumour site [5]. Recently, it has been reported that liposomal daunorubicin may be a safe and effective formulation for treating AIDS-related KS [6]. Experience with the usefulness of this drug formulation is limited. We report an AIDS patient with rapidly progressive pulmonary KS, who responded to treatment with a liposomal daunorubicin formulation (DaunoXome®, Vestar, San Dimas, CA, USA).

**Case report**

A human immunodeficiency virus (HIV) seropositive male homosexual patient, aged 35 yrs, presented in October 1991 in poor condition, with a 4 week history of cough, dyspnoea and weight loss. HIV infection had been diagnosed in 1985. A right-sided local cutaneous thoracic KS was diagnosed 14 months earlier, which had temporarily responded to irradiation. The patient had a history of thoracic herpes zoster, recurrent colitis due to *Giardia lamblia*, and disseminated *Mycobacterium avium* complex (MAC) infection, the latter having been the initial AIDS diagnosis in March 1990.

The patient had received zidovudine for 15 months. In addition, he was receiving 25 mg pyrimethamine/500 mg sulfadoxine (one tablet Fansidar®), plus 15 mg folic acid twice a week for prophylaxis against *Pneumocystis carinii* pneumonia and toxoplasma encephalitis, 200 mg fluconazole twice a week for prevention of oral candidiasis, and 450 mg rifabutin plus 1,600 mg ethambutol daily as maintenance treatment for MAC infection.

Pertinent laboratory values included: CD4+ lymphocyte count 0.003×10^9 l⁻¹ (3 µl⁻¹), haemoglobin level 117 g l⁻¹, platelets 90×10^9 l⁻¹, and leucocyte count 1.7×10^9 l⁻¹ (absolute granulocyte count 1.04×10^9 l⁻¹). Chi-square test showed extensive reticulonodular perivascular infiltrates in the left lower lobe and lingula. A vascularized tumour was revealed by bronchoscopy (fig. 1). Computed tomography showed extensive reticulonodular perivascular infiltrates in the left lower lobe and lingula. A vascularized tumour was revealed by bronchoscopy, causing a 90% occlusion of the lingula bronchus. Biopsies were diagnostic of KS. Vital capacity (VC) was reduced to 71% of the predicted value. Chest X-ray 10 days later showed rapid progression.

Treatment was initiated with 40 mg·m⁻² DaunoXome® every 14 days. Dose modification with a dose reduction to 50% for a granulocyte count less than 1.0×10^9 l⁻¹ and with dose delay for 1 week for a granulocyte count less than 0.5×10^9 l⁻¹ was planned. Antiretroviral treatment and chemoprophylaxis of opportunistic infections were continued with the same doses and schedules as before chemotherapy.

Respiratory symptoms resolved completely within 4 weeks. VC returned to 110% of the predicted value after five cycles. Response to treatment was demonstrated by chest X-ray after two cycles. After six cycles, maximal resolution had been achieved, with only small residual alterations being demonstrated by chest X-ray (fig. 2) and thoracic computed tomography. Re-evaluation by bronchoscopy revealed almost complete resolution of endobronchial lesions, with only slightly erythematous...
endobronchial foci being found. Biopsies were negative for KS.

Leucopenia required delay of cycle 3 for 1 week, 50% dose reduction during cycles 5 and 7, delay of cycle 13 for two weeks, and 50% dose attenuation for cycles 14–17. After cycle 15, progression was diagnosed, with reappearance of respiratory symptoms and of radiographic manifestations. Granulocyte colony-stimulating factor (G-CSF) (5–10 µg·kg⁻¹ body weight daily) was given between 4–10 days after cycle 17 between all the following cycles, enabling continuation of the original dose schedule. With continuation of full doses, remission was reinduced and maintained until the patient's death, which was due to progression of his HIV infection.

The case reported shows that liposomal daunorubicin may be a relatively safe and effective long-term treatment regimen for pulmonary KS. The patient tolerated chemotherapy well for a long duration, in spite of a poor pretreatment condition, advanced AIDS and a markedly reduced bone marrow reserve. The only toxicity seen during the course of treatment was marrow suppression. Importantly, dose reduction required by bone marrow toxicity was associated with progression of KS. However, the use of G-CSF to reduce bone marrow toxicity enabled continuation of the original dose schedule. With continuation of full doses, remission was reinduced and maintained until the patient's death, which was due to progression of his HIV infection.

The efficacy of liposomal daunorubicin in treating AIDS-related KS, including pulmonary involvement, and its good tolerability has recently been confirmed [7]. The response rate in 25 patients treated with the same formulation of liposomal daunorubicin (DaunoXome®) as the patient reported here, was 57%. Two of these 25 patients had pulmonary involvement. One patient showed partial remission and the other complete remission of pulmonary KS. Chemotherapy of AIDS-related KS must be given lifelong, since the tumour progresses rapidly after cessation of treatment [3, 4]. However, toxicity limits the benefits of chemotherapy, making long-term treatment difficult [3]. Thus, there is need for an effective regimen with low toxicity. Liposomal daunorubicin may be a promising alternative to the chemotherapy protocols in current use.

Discussion

The case reported shows that liposomal daunorubicin may be a relatively safe and effective long-term treatment regimen for pulmonary KS. The patient tolerated chemotherapy well for a long duration, in spite of a poor pretreatment condition, advanced AIDS and a markedly reduced bone marrow reserve. The only toxicity seen during the course of treatment was marrow suppression. Importantly, dose reduction required by bone marrow toxicity was associated with progression of KS. However, the use of G-CSF to reduce bone marrow toxicity enabled continuation of the original dose schedule. With continuation of full doses, remission was reinduced and maintained until the patient's death, which was due to progression of his HIV infection.

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References