A 51-year-old HIV-infected man presented with a 4-week history of shortness of breath, right-sided chest pain, drenching night sweats, and 5-kg weight loss. He had been diagnosed with HIV infection 23 years previously. Investigations showed anemia (hemoglobin, 9.8 g/dL), an elevated C-reactive protein of 138.6 mg/L, a CD4 count of 550 cells/μL, HIV viral load of 6800 copies/mL, and plasma Kaposi sarcoma–associated herpes virus (KSHV)/human herpes virus 8 (HHV8) DNA of 70,000 copies/mL. A chest radiograph showed a moderately large right-sided pleural effusion. He was treated empirically for atypical pneumonia. Despite a course of antibiotics, the patient continued to complain of breathlessness. A chest radiograph 10 weeks later revealed a right-sided loculated effusion and pleural thickening. A computed tomography (CT) scan showed multiple pleural masses, the largest measuring 6.8 × 4.3 cm, located at the right posterolateral base, a pericardial nodule adjacent to the right atrium, a right-sided pleural effusion, and multiple small volume supraclavicular, axillary, retroperitoneal, pelvic, and inguinal adenopathy. Biopsy of the right base pleural mass identified a neoplastic infiltrate composed of large lymphoid cells with irregular pleomorphic nuclei. Many of the cells had eccentrically placed nuclei and abundant cytoplasm, giving the tumor a plasmablastic appearance (Fig 1A). Immunohistochemistry demonstrated that the tumor cells were positive for CD138 (Fig 1B), epithelial membrane antigen, and HHV8 (Fig 1C) and showed lambda light chain restriction. Epstein-Barr virus in situ hybridization was negative, and mindbomb homolog 1 showed a proliferation fraction of over 90%. The B-cell markers CD20 (Fig 1D), CD79a, and paired box gene 5 were negative. The radiologic, histologic, and serologic findings were those of HIV-associated primary effusion lymphoma (PEL). He was commenced on a regimen of cyclophosphamide, doxorubicin, vincristine, and prednisolone (CHOP) chemotherapy with prophylactic intrathecal methotrexate and simultaneously started on antiretroviral therapy (ART; tenofovir, emtricitabine, and efavirenz). A CT scan after six cycles of CHOP chemotherapy showed significant disease response, with the right posterior pleural mass reduced to 4.8 × 3.2 cm and complete resolution of the pleural effusion. However, a positron emission tomography/CT scan assessment after eight cycles of CHOP chemotherapy showed an enlarging [18F]fluorodeoxyglucose–avid pleural mass (5 × 5 cm). Biopsy of the posterior pleural mass showed histological appearances identical to previous biopsy, validating a diagnosis of refractory PEL. The patient started...
PEL is a rare HIV-associated malignancy representing around 3% to 4% of all HIV-related non-Hodgkin’s lymphoma (NHL) with poor clinical outcome.\(^1,2,3\) PEL is defined by serous body cavity effusions and expression of KSHV, also known as HHV8, in their tumor cells.\(^1,2,5\) Over 70% of cases display concurrent Epstein-Barr virus infection. The optimal treatment for PEL has not yet been defined. In most cases, patients are treated with CHOP chemotherapy together with introduction of highly active ART.\(^1\) Despite this, durable responses are rarely seen and survival is short with median survival averaging 3 to 6 months.\(^1,4,6,7\) To our knowledge, this is the first description of a patient with HIV-associated PEL in whom radiation treatment was associated with prolonged clinical, radiological, and serological response after failing three regimens of systemic chemotherapy treatments. Given the chemotherapy-refractory behavior of the disease, the dramatic response to radiotherapy was unexpected and surprising. Our experience suggests PEL is sensitive to radiation treatment and should be considered as part of the treatment recommendation for patients with chemotherapy-refractory PEL-associated solid masses. It would also be interesting to explore treating PEL patients with consolidation radiation therapy for any residual histologically proven masses following conventional CHOP chemotherapy or with total-body irradiation, whereas the role of high-dose chemotherapy and peripheral stem-cell transplantation in refractory or relapsed HIV-associated PEL is yet to be ascertained.\(^8\)

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ACKNOWLEDGMENT
Supported by University College London Hospital/University College London Comprehensive Biomedical Research Centre and Gerard Kingston for secretarial assistance.

AUTHORS’ DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST
The author(s) indicated no potential conflicts of interest.

REFERENCES
Chronic Lymphocytic Leukemia With Eyelid Involvement Responding to Alemtuzumab

A 61-year-old man was diagnosed in another hematological department as having chronic lymphocytic leukemia (CLL) in 1998. One year later, because of progressive stage II CLL characterized by doubling lymphocytes count after 6 months, increase of splenomegaly, and the appearance of B symptoms, the patient was treated with fludarabine, obtaining partial remission of disease. Starting from 2000, due to progression of disease, the patient underwent a cyclophosphamide, doxorubicin, vincristine, and prednisolone regimen plus interferon-α as maintenance therapy for 2 years; thereafter the successive relapse was treated with chlorambucil plus metilprednisolone in 2003 and chlorambucil in 2005, respectively. In January 2006, he presented lymphoadenopaties between 1.5 and 2.5 cm in his right armpit, and inguinal bilaterally inferior margin of spleen 2 cm after the costs margin, absent of B symptoms. A total-body computed tomography scan confirmed multiple lymphoadenomegalies in the mediastinum, abdomen, and splenomegaly. An emocromocytomet-ric analysis presented a WBC of 27,280/mm³ (neutrophil, 2,000/mm³; leukocyte, 23,000/mm³); hemoglobin of 14.3 g/dL; medium cellular volume of 84.6 fl; and platelets of 204 × 10⁹/L. A peripheral blood smear showed more than 50% of big lymphocytes (its size was double in respect to erythrocyte size); for these reasons we classified this form as atypical chronic lymphocytic leukemia (mixed CLL). Bone marrow aspiration showed 55% of infiltration by mature lymphocytes, and a bone marrow biopsy confirmed the same diffuse infiltration. Cyt-ofluorimetric analysis of peripheral blood showed positivity for CD5, CD5/CD19, CD23, and κ; at the same time the positivity of CD38 (73%) and ZAP-70 (41%) was detected. Fluorescent in situ hybridiza-tion analysis showed trisomy of chromosome 12, as expected by mor-phological analysis and 17p deletion. The mutational status of variable region of heavy chain immunoglobin discovered a mutated status (variable heavy chain 3-49). Because of stable stage II disease with bad biologic parameter in the heavily pretreated patient, a stringent follow-up was adopted. In October 2006, at routine check-up, the patient showed a bilateral painless nodular infiltration of the eyelids, similar to multiple chalazions. Two nodules were localized in each