1 | INTRODUCTION

A 26-year-old woman developed bilateral blindness after a complete remission for ALK-positive ALCL. Initial workup showed neurolymphomatosis involving the bilateral optic nerves. She underwent hematopoietic cell transplantation, and she remained in complete remission after 1 year; however, her vision never improved. ALCL likely led to the destruction of the optic pathway.

Neurolymphomatosis is a rare clinical entity in which lymphoma cells with neurotropism infiltrate and destroy peripheral nerves, spinal nerve roots, nerve plexuses, and cranial nerves.1-3 Neurolymphomatosis associated with T-cell lymphoma is very rare.1 In large series, T-cell lymphoma was diagnosed in 2.5%-10% of neurolymphomatosis cases.1-3 We hereby report a very unusual case of CNS involvement by ALCL with bilateral optic neurolymphomatosis resulting in permanent blindness.

2 | CASE

A 26-year-old woman was diagnosed about a year ago with stage IIA ALK+ ALCL with right axillary and cervical lymphadenopathy and was initiated on CHOP (cyclophosphamide, doxorubicin, etoposide, vincristine, and prednisone) chemotherapy. She achieved complete remission following three cycles of treatment. Soon thereafter, she developed sudden loss of vision in both eyes associated with fever, and night sweats. Vision loss started in the left eye then progressed to the right eye, with severity at a loss of light perception level. She was transferred to our institution. Significant
clinical findings include loss of vision bilaterally with limited ability to appreciate light and dark stimulus. Pupils were dilated bilaterally both with a positive direct and indirect light reflex. She had left ocular dysmotility consistent with CN6 palsy. No focal weakness or sensory deficit bilaterally in the face and extremities. She had no lymphadenopathy, or hepatosplenomegaly. Serum lactate dehydrogenase enzyme level was elevated at 406 U/L (122-222 U/L). MRI brain imaging showed bilateral nodular leptomeningeal enhancements in the posterior fossa and enhancement around both optic nerves (Figure 1) consistent with neurolymphomatosis. CSF cytology was positive for atypical large cells consistent with ALCL (Figure 2). Bone marrow biopsy did not show any involvement with lymphoma. She did not have any clinical evidence of neurolymphomatosis involving peripheral nerves. Review of her outside right axillary lymph node biopsy (Figure 3) confirmed the diagnosis of ALCL with the proliferation of large, atypical lymphocytes with horseshoe-shaped nuclei on hematoxylin and eosin stain and strong expression of ALK-1, CD5, and CD30, and negative expression of PAX 5, CD20, or CD3 by immunohistochemistry studies.

She was initially treated with dexamethasone 40 mg/d for 4 days followed by a taper and the Ferrari regimen with high-dose methotrexate 3500 mg/m² and high-dose cytarabine 2000 mg/m² with intrathecal methotrexate 12.5 mg. This was followed with 10 sessions of radiation to the bilateral orbits and optic nerves, a total cumulative dose of 3000 cGy. There was no significant improvement in her vision after the initial treatment. She continued with the Ferrari regimen every 3 weeks for a total of four cycles. After two cycles, there was no radiological evidence of residual lymphoma on follow-up brain MRI imaging (Figure 4). Furthermore, repeat CSF cytology was negative for malignant cells. The patient subsequently underwent high-dose chemotherapy with BCNU and thiopeta followed by autologous stem cell transplant. One year after her transplant, she continues to be in complete remission for both systemic and CNS disease and had small improvement in light and color perception. In spite of her CNS disease being responsive to treatment and achieving complete remission, her vision never improved significantly.

3 DISCUSSION

Our patient has a very unique clinical presentation with bilateral optic neurolymphomatosis in the setting of treatment sensitive systemic ALK+ ALCL. Although she was treated promptly with radiographic complete response of CNS disease, her vision never recovered resulting in permanent blindness. It is understandable that the optic nerves and tracts were infiltrated and destroyed rather quickly and irreversibly. Our case appears to be the first case of bilateral optic neurolymphomatosis with ALCL reported in the literature as our extensive search did not turn up any such case.

Neurolymphomatosis of cranial nerves by T cell lymphoma is rare. In a large study reported by International Primary CNS Lymphoma Collaborative Group (IPCG), the involvement of cranial nerves was seen in 46% of 50 patients. In another large study from Massachusetts General Hospital (MGH), cranial nerves were involved in 51% of 72 patients. However, a report from Mayo Clinic showed 10% (4/40) cranial nerve involvement. Histologically, most cases of neurolymphomatosis are related to B-cell lymphomas at 82% of cases (IPCG and MGH), and 97.5% of cases (Mayo). T-cell lymphomas account for 10% of cases (5/46

**FIGURE 1** Pretreatment gadolinium-enhanced T1 MRI: optic nerve infiltration by lymphoma (A and B, arrows). Leptomeningeal lymphomatosis (C, arrowheads)
cases, IPCG study), 5% of cases (4/72 cases, MGH study), and 2.5% of cases (1/40 cases, Mayo study).1-3 Interestingly, one case series reported neurolymphomatosis of cranial nerves with leptomeningeal involvement in 4/5 patients with primary T-cell CNS lymphoma.4 Our patient had a similar distribution of CNS involvement.

In our patient, the diagnosis of optic neurolymphomatosis was supported by diffuse bilateral enhancement of optic nerves and tracts on MRI together with positive CSF cytology and known systemic ALK+ ALCL based on right axillary lymph node biopsy. We suggest that enhancement of cranial nerves on MRI with evidence of lymphoma in CSF or peripheral sites should be regarded as diagnostic of cranial neurolymphomatosis. The diagnosis can be quite difficult requiring meningeal and nerve biopsies when CSF studies and peripheral site biopsies are negative.1-3

Primary or secondary CNS involvement of ALK+ ALCL is not common.5 In a retrospective analysis of 600 patients with PTCL, the 5-year cumulative incidence of CNS relapse in ALK+ ALCL is 5.4% (4/74 patients).1-3 There has been no standardized protocol to assess the CNS risk in patients with peripheral T-cell lymphoma. Involvement of more than one extra nodal site is a significant risk factor for CNS relapse with bone, subcutaneous tissue, spleen, skin, lung, and liver regarded as high-risk sites.6 Bone marrow involvement was reportedly associated with higher risk of CNS involvement in some reports.7 Extra nodal involvement has a 1-year cumulative incidence of 17% for CNS relapse.6,8 Other risk factors for CNS relapse include elevated serum lactate dehydrogenase (LDH), and high International Prognostic Index (IPI) score ≥3, B symptoms, ALK+ histologic type, and stage III-IV disease.6,8 CNS prophylaxis for PTCL is controversial with little data reported.9 Some authors advocate for evaluation of CNS at the time of diagnosis and possible CNS-directed prophylaxis in patients with ALK positivity and extra nodal involvement.6 In our patient, the presence of

**FIGURE 2** The cytospin slide from CSF fluid (modified Giemsa stain ×20) contains many large atypical cells (see green arrowheads); and occasional mitosis is present (see black arrow)

**FIGURE 3** The H and E section of right axillary lymph node core needle biopsy (A, ×20) shows the numerous large atypical cells with horseshoe-shaped nuclei and prominent nucleoli. Immunohistochemistry studies show the neoplastic lymphocytes positive for CD 30 (B, ×20), ALK-1 (C, ×20), and CD 5 (D, ×20); negative for PAX5, CD20, CD3
ALK positivity, elevated LDH, and B symptoms may have increased her risk for CNS relapse although she did not have extranodal involvement on initial diagnosis.

Interestingly, she developed CNS disease after achieving complete response of systemic disease to three cycles of CHOEP chemotherapy. It is possible that CHOEP chemotherapy did not have prophylactic or therapeutic impact on CNS disease, or the damage may have already occurred. It is not surprising as chemotherapeutic agents in the regimen are known for poor CNS penetration. However, she had complete CNS response to CNS-penetrating agents, high-dose methotrexate, and high-dose Ara-c. We recommend that CNS-penetrating agents should be used as early as possible in PTCL patients with high CNS risk or CNS disease. High-dose methotrexate-based treatment has been associated with improved survival in neurolymphomatosis and primary CNS T-cell lymphoma.3,9

In conclusion, our case highlights devastating consequences secondary to CNS involvement by ALK+ ALCL. Novel biomarkers and clinical markers need to be identified for better CNS risk assessment in patients with ALCL. Early treatment with high-dose methotrexate-based regimen should be instituted in patients with CNS involvement or those at an increased CNS risk.

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CONFLICT OF INTEREST
Mohamed A. Kharfan-Dabaja: reports consultancy for Pharmacyclics and Daiichi Sankyo. All other authors have no conflict of interest to declare.

AUTHOR CONTRIBUTIONS
KS, DF, KK, and JH: made substantial contributions to conception and design, acquisition of data, and drafting the manuscript. MAM: made substantial contributions to acquisition of data and drafting the manuscript. MAK-D, EA, VG, HWT, and LJ: made substantial contributions to conception and design, and revised it critically for important intellectual content.

ETHICAL APPROVAL
Not needed.

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