INTRODUCTION

Neurolymphomatosis (NL) is the neoplastic endoneurial invasion of the peripheral nervous system (PNS). It differs from meningeal lymphomatosis, which is the infiltration of the PNS due to subarachnoid seeding (1). Furthermore, NL occurs more frequently with the spread of a known malignancy into the PNS from systemic sites or from the central primary nervous system (CNS) lymphoma (PCNSL) as a progression in the course of the disease or as a relapse following complete remission. More rarely, it may occur as an initial manifestation of the disease. We report a case of a 30-year-old male patient who was in complete remission from T-lymphoblastic lymphoma, presenting with clinical findings indicating initially ulnar entrapment. However, with the demonstration of brachial plexopathy with axonal loss in electrodiagnostic studies, MR imaging neurography dedicated to brachial plexus was carried out and revealed pathological enhancement associated with mild fluorodeoxyglucose (FDG) uptake on PET/CT. Moreover, NL, due to the relapse of T-cell lymphoma, was diagnosed through incisional biopsy, showing diffuse infiltration of blast cells positive for terminal deoxynucleotidyl transferase, CD3 and CD10. Further, radiotherapy and systemic chemotherapy were initiated, and symptoms recovered with regression of pathological FDG uptake.

Keywords: Neurolymphomatosis, T-cell lymphoblastic lymphoma, brachial plexopathy

Case Report / Olgu Sunumu

Relapse of T-Cell Lymphoma as Isolated Brachial Plexus Neurolymphomatosis: A Case Report

T-Hücreli Lenfomanın İzole Brakial Pleksus Nörolenfomatozisi Olarak Relapsı: Olgu Sunumu

Kenan Kıbıcı1, Berrin Erok2, Ali Önder Atça 3, Gülçin Yeğen4

1Altınbaş University Faculty of Medicine, Bahçelievler Medical Park Hospital, Department of Neurosurgery, İstanbul, Turkey
2University of Health Sciences Turkey, Prof. Dr. Cemil Taşçıoğlu City Hospital, Clinic of Radiology, İstanbul, Turkey
3Altınbaş University Faculty of Medicine, Bahçelievler Medical Park Hospital, Department of Radiology, İstanbul, Turkey
4İstanbul University Faculty of Medicine, Department of Pathology, İstanbul, Turkey

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ABSTRACT

Neurolymphomatosis (NL) is the neoplastic endoneurial invasion of the peripheral nervous system. It is a rare and challenging diagnosis, but it should be taken into account in the differential diagnosis of peripheral neuropathy, particularly in patients with a documented history of haematologic malignancy. Magnetic resonance (MR) neurography is very useful in diagnosis and especially when correlated with positron emission tomography/Computed tomography (PET/CT). In equivocal cases, nerve biopsy can be considered when the benefit outweighs the risk. We aimed to report a case of a 30-year-old male patient who was in complete remission from T-lymphoblastic lymphoma, presenting with clinical findings indicating initially ulnar entrapment. However, with the demonstration of brachial plexopathy with axonal loss in electrodiagnostic studies, MR imaging neurography dedicated to brachial plexus was carried out and revealed pathological enhancement associated with mild fluorodeoxyglucose (FDG) uptake on PET/CT. Moreover, NL, due to the relapse of T-cell lymphoma, was diagnosed through incisional biopsy, showing diffuse infiltration of blast cells positive for terminal deoxynucleotidyl transferase, CD3 and CD10. Further, radiotherapy and systemic chemotherapy were initiated, and symptoms recovered with regression of pathological FDG uptake.

Keywords: Neurolymphomatosis, T-cell lymphoblastic lymphoma, brachial plexopathy

ÖZ

Nörolenfomatoz (NL) periferik sinir sisteminin neoplastik endonöral invasyonudur. Nadir ve zorlayıcı bir tanıdır ancak, özellikle hematolojik malignite öyküsü bulunan hastalara periferik nöropatinin ayrıca tansında düşünülmelidir. Magnetik rezonans (MR) nörografide pozitron emisyon tomografisi-bilgisayarlı tomografi (PET/CT) ile beraber kullanılanında tanda çok yararlıdır. Arada kalın olgularda fayda riskten ağır bastığında periferik nöropati tani konulduğuunda bu an için periferik nöropatiden duyulan bulgulara 30 yaşındaki erkek hastanın sebebi olarak değerlendirildi. Ancak, elektrodagnostik çalışmaları aksonal kayıp، özellikle brakial pleksopatini göstermesi üzerine brakial pleksususal patoloji için yapılan MR nörografide sah brakial pleksustaki patolojik kısıtlama ve bu alanda PET/CT ile patolojik fluorodeoksiglukoz (FDG) tutulumu izlendi. Terminal deoksinükleotidil transferaz, CD3 ve CD10 için pozitif olan yağın blastik hücre infiltrasyonları göstererek T-Hücreli lenfoma relapsı ile ilgili LN tanısı insinyonel biyopsi ile konuldu. Radyoterapi ve sistemik kemoterapi tekrar başlandı ve hastanın bulgularında patolojik FDG tutulumunun gerilemesi ile beraber iyileşme olduğu tespit edildi.

Anahtar kelimeler: Nörolenfomatozis, T-Hücreli lenfoblastik lenfoma, brakial pleksopati

ORCID IDs of the authors: K.K. 0000-0002-5912-964B; B.E. 0000-0001-8036-547X; A.O.A. 0000-0002-7500-3316; G.Y. 0000-0003-2497-219X.

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A 30-year-old male patient presented with right medial forearm and hand pain and numbing in the fourth and fifth fingers, initially indicating ulnar nerve entrapment. A year ago, he was treated for T-lymphoblastic leukaemia/lymphoma and achieved complete remission with no pathological fluorodeoxyglucose (FDG) uptake in positron emission tomography/computed tomography PET/CT. In neurological examination, muscular strengths of the forearm flexors and intrinsic hand muscles were poor, and there was pain-related paraesthesia. Moreover, no disc impingement was demonstrated by cervical magnetic resonance imaging (MRI). Ulnar nerve entrapment was considered; however, electrodiagnostic studies revealed right lower truncus brachial plexopathy associated with axonal loss and active denervation. Gadolinium-enhanced MRI neurography showed a 25x32x41 mm-sized enhancing extrapulmonary lesion located at the apical part of the right hemithorax, affecting the middle and lower part of the brachial plexus (BP) (Figure 1). No pathological signal was detected in the adjacent pulmonary parenchyma. Also, there was no extension into the spinal canal. PET/CT showed a mild hypermetabolic FDG uptake (maximum standardized uptake value: 3.04) in the lesion. Moreover, NL was considered and incisional biopsy was performed to make an exact diagnosis. During the antigenic assessment, diffuse infiltration of blast cells positive for terminal deoxynucleotidyl transferase, CD3 and CD10 was demonstrated (Figure 2). Once again, radiotherapy and systemic chemotherapy was started. With the regression of pathological FDG uptake in PET/CT, his symptoms recovered with improvement of muscle strengths from 3/5 to 5/5.

Written informed consent has been taken from the patient.

**DISCUSSION**

NL is a very rare neurological manifestation of haematologic malignancies that should be distinguished from the more prevalent causes of PNS involvement, such as radiation-induced peripheral neuropathy or paraneoplastic syndromes. Most cases of NL are associated with diffuse large B-cell non-Hodgkin’s lymphoma (DLBCL) but can also occur in T-cell and NK-cell lymphomas or acute leukaemias. According to the report of the International Primary CNS Lymphoma Collaborative Group, NL developed in 50 patients over a 16-year period, of which 90% were associated with non-Hodgkin’s lymphoma (NHL) and 10% with acute leukaemia (2). In a 12-year retrospective analysis of 1,181 patients with NHL, 22 cases of NL were identified, of which 20 patients were associated with DLBCL, 1 patient with mantle cell lymphoma, and 1 patient with peripheral T-cell lymphoma (3). In another report of 23 cases with NL of lumbosacral plexus, B-cell histology was seen in 21 cases and T-cell histology in only 2 cases (4). In our patient, NL was related to a relapse of T-cell lymphoma. Since the clinical findings are not clear, it is frequently misdiagnosed, especially when the neuropathy is the initial presentation or manifests after a period of complete remission, as in our case. When the BP is affected, the presenting symptoms are weakness, paraesthesia and pain radiating down to the upper limb or localised in the forearm and hand in cases of middle and lower trunk involvement, as in our case. In contrast to the radiation-induced brachial plexopathy that commonly affects the upper trunk or metastatic brachial plexopathy that has a predilection for the lower trunk, all parts of BP may be invaded by lymphomatous cells from the roots to the distal branches with no distinctive distribution (5). As suggestive of NL, severe pain, rapid evolution and asymmetric distribution have been reported (2). However, relatively painless or symmetrical neuropathy has also been defined (6,7). Although axonal sensory motor neuropathy is more frequent in nerve conduction studies, pure demyelinating and mixed neuropathy have also been reported (7). Moreover, due to its low sensitivity, cerebrospinal fluid (CSF) cytology is used in the diagnosis but is not always helpful (6). Neuroimaging studies with MRI neurography often provide a correct diagnosis, especially when performed together with PET/CT. On MRI,
normal nerves are usually smaller than the accompanying arteries, showing gradual tapering in a well-organised fascicular distribution with isointensity to skeletal muscles on both T1w and T2w images without enhancement, except the dorsal root ganglia which should not be confused with a tumoral process. In NL, focal or diffuse nerve enlargement, fascicular disorganisation, hyperintensity on T2w or short-tau inversion recovery sequences and significant focal or diffuse gadolinium enhancement are seen. Although these are not specific for NL, they are highly suggestive when associated with PET/CT, which is the most accurate modality in the evaluation of nodal and extranodal spread of lymphoma (8,9). In nerve pathology, the gold standard method of diagnosis is nerve biopsy, but it is not performed in all patients because of a significant risk of permanent nerve damage. Moreover, it is concerned when the benefit outweighs the risk and the imaging studies along with CSF examinations are inconclusive. In general, treatment of NL is similar to that of PCNSL, which includes systemic and/or intrathecal chemotherapy with or without radiotherapy (8). In rare instances, surgical intervention, such as emergent decompression, is needed. Our patient was treated with radiotherapy and systemic chemotherapy with achievement of complete remission. Since the relapse of the lymphoma in our patient was localised to the right BP, radiotherapy was performed with limited-field radiation. Despite the treatment, the prognosis is worse in patients with NL than in those without (10). Although the lack of optimal treatment one of the causes, it may also be due to misdiagnosis that leads to delayed targeted treatment.

CONCLUSION

NL remains a rare and challenging diagnosis. It should be considered when a patient with a known history of haematologic malignancy presents with peripheral neuropathy, especially when the symptoms are asymmetric and rapidly evolving. MRI neurography and PET/CT are very helpful and should be used together to improve sensitivity. In equivocal cases, nerve biopsy can be used for early diagnosis when the benefit outweighs the risk, especially when, as in our case, there is an already existing axonal pattern of damage.

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