Natural Killer Cell Lymphoma: A Case with Classification Dilemma

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ABSTRACT
Non-Hodgkins lymphoma of the Natural Killer (NK) cell type is rare. World Health Organisation recognises 3 NK-cell phenotypic entities; extranodal NK/T cell lymphoma, nasal type (ENK/TL); aggressive NK cell leukaemia (ANKL); and chronic lymphoproliferative disorders of NK cells (CLPD-NK) which is classified as a provisional entity. Though specific clinical, morphological and immunophenotypic criteria have been laid down to diagnose these conditions there may however, be considerable variations in the clinical presentation making diagnosis difficult. We present a case with contrasting clinical and haematopathological findings posing difficulty in its diagnosis and classification, and despite the aggressive presentation showing favourable response to treatment.

CASE REPORT
A 30-year-old man was admitted with 1 month history of fever, weakness, lethargy, non quantifiable weight loss and cervical lymphadenopathy in a private hospital. The patient was diagnosed with diffuse large B-cell lymphoma on fine needle aspiration and was started on CHOP regimen. However, following two cycles of chemotherapy, no response was seen and he was referred to our institute. Details pertaining to diagnosis and immunophenotyping were not available with the patient on initial presentation.

The patient on arrival was febrile, pulse 90/min, BP 110/70 mm Hg. On examination bilateral cervical, axillary and inguinal lymph nodes were noted along with hepatosplenomegaly. There was also diffuse midline swelling consistent with enlargement of thyroid gland. Examination of the nasal cavity was unremarkable. On investigation, he was mildly anaemic (haemoglobin 10.8 g/dl), total bilirubin 0.7 mg/dl, ALT 27 U/L, AST 22 U/L, blood urea nitrogen 24 mg/dl and serum creatinine 0.5 mg/dl. Serum electrolytes were normal and serum LDH was raised. Peripheral blood examination showed normocytic normochromic anaemia and no atypical cell was noted.

On imaging, multiple abdominal and mediastinal lymph nodes were noted, along with multiple foci of hepatic involvement. Fine needle aspiration (FNA) performed at the private hospital, from the cervical lymph node; thyroid and liver were retrieved and reviewed. All showed highly cellular smear with sheets of medium to large cells consistent with the diagnosis of non-Hodgkins lymphoma [Table/Fig-1a]. Flow cytometry (FCM) was performed using FACSCalibur (Becton Dickinson, San Jose, CA, USA) on a fresh cervical lymph node aspirate. Cells were gated on SSC/CD45 dot plot. The gated population showed normocytic normochromic anaemia and no atypical cell was noted.

Based on the clinical presentation and the investigation, a diagnosis of non-Hodgkins lymphoma of NK-cell phenotype, Ann-Arbor stage IVB was made. However, we were not able to subclassify our case into any of the NK-cell lymphoma entity described in the 2008 WHO classification. The patient was started on VPD chemotherapy regimen. Ifosfamide at 1200mg/m²/day as a slow infusion over 3 hours with mesna injection at 0, 4 and 8 hours, etoposide at 100 mg/m²/day and cisplatin at 30 mg/m²/day over 2 hours were given for 3 days. Intravenous dexamethasone, 40 mg was given on day 1 to 4. He has received 3 cycles of chemotherapy. All the cycles were uneventful and presently, as assessed by the revised response evaluation criteria in solid tumour (RECIST) guidelines [1], the patient is under partial remission and regular follow up.

DISCUSSION
Lymphoma of the NK cell phenotype is rare. International T-cell lymphoma project reports that 10.4% of T cell lymphomas are of NK/T cell type (NK/TCL). WHO recognized entities include extranodal NK/T cell lymphoma, nasal type (ENK/TL), which is predominantly nasal or may rarely be extranasal; aggressive NK cell leukaemia (ANKL) and chronic lymphoproliferative disorders of NK cells (CLPD-NK). These disorders are uncommon and are seen in Asians, Central and South Americans, and Mexicans. Among
these entities, CLPD-NK is indolent but the other two follows an aggressive course with variable prognostic and short survival [2-4]. The 5 year overall survival for NK/TCL is 32% and this drops down to 9% for extranasal, aggressive and unclassifiable NK cell lymphoma [5]. Diagnosis of NK cell lymphoma can be challenging. Unlike T-cell lymphomas, NK-cell lymphoma fails to show receptor gene rearrangement. So, immunophenotyping of the NK-lymphoma cells play a critical role in disease characterisation [6]. ENKL shows an Asian and South American predilection, occurs exclusively in extranodal fashion involving nasal and upper aerodigestive tract with rare advanced cases showing disseminated lymphadenopathy, hepatomegaly or leukemic phase. Rarely, exclusive extranasal site like skin, salivary glands, testis, and gastrointestinal tract may be involved. They present mostly in stage I/II, with the extranasal cases being more aggressive, presenting at stage III/IV. They are associated with Epstein-Barr virus (EBV), and quantification of EBV DNA helps to assess tumour load, and disease prognosis. Morphologically, it exhibits ulceration, geographic necrosis with angiocentricity and angiodestruction and these cells shows cytoplasmic CD3 and CD56 positivity [2,7]. ANKL are aggressive involving peripheral blood (PB), bone marrow (BM), spleen, liver, presenting with fever, jaundice with variable cytopenia, leukemic blood picture, sometimes associated with haemophagocytosis and generally progresses to multigorgan failure. They are associated with EBV. It occurs at a younger median age than ENK/TCL or CLPD-NK, with frequent BM involvement and rare cutaneous involvement. These features are contrary to the suggestion that ANKL are merely the leukemic phase of ENK/TCL [3,8]. The disease is aggressive, not responsive to conventional chemotherapy with median survival of 2 months [9]. CLPD-NK is rare and indolent. Most patients are asymptomatic or show variable cytopenia. PB and BM are the predominantly involved sites, with hepatosplenomegaly, lymphadenopathy and cutaneous involvement being infrequent. They may occur in association with other medical conditions, and rarely may transform to aggressive forms [4,10].

The present case had a unique presentation of a young patient, in a disseminated state of nodal and multiple extranodal site involvement, without evidence of nasal or PB disease. Immunophenotype was consistent with NK cell but based on the clinical presentation, we were unable to place the case in any of the WHO described entities. The survival rate according to literature for such aggressive clinical presentation usually does not exceed 2 months. Despite the aggressive presentation, the patient responded to the prescribed chemotherapy regimen. After six months of diagnosis and 3 cycles of chemotherapy, the patient significantly improved with disappearance of all the peripherally palpable target lesions, including lymph nodes and thyroid. Presently, the abdominal lymph nodes and hepatic lesion show partial remission.

CONCLUSION

We wish to highlight that a revision including wider clinical presentations, with specific diagnostic markers maybe necessary to increase the diagnostic accuracy and reproducibility for such cases, as this might have profound impact on their survival and disease prognostication.

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REFERENCES


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