NK/T Cell Lymphoma - A Case Report and Literature Review

Abstract

We present here the case of an 87 year old Caucasian female who complained of nasal tumour causing partial nasal obstruction without bleeding. The diagnosis nasal T/NK cell lymphoma was established by immunohistochemistry. The case was diagnosed as stage IE lymphoma because no other site was involved. The patient was treated using 6-15MeV linear accelerators at 2 Gy per daily fraction for a total dose of 44 Gy. The nasal T/NK cell lymphoma is a rare, special entity of non hodgkin lymphomas and is known to be very aggressive in nature. The choice of treatment depends upon the stage of the lymphoma. In the initial stages locorregional radiotherapy alone or combined with chemotherapy is recommended. For the stage III/IV NK/T Cell lymphomas combined modality therapy consisting of high-dose chemotherapy followed by auto-PBSCT can be an alternative. In short, a high index of clinical suspicion is the key required, in order to reach early diagnosis that could open the doors of treasure i.e. improved disease outcome.

Keywords: Nasal NK/T cell lymphoma; Immunohistochemistry; Tumour; Nasal congestion

Case Report

A Caucasian 87 year old woman was referred to department of Teletherapy, Maria Skłodowska Curie Centre and Institute of Oncology for the radical treatment of T cell lymphoma of nasal type. The patient noticed about 8 months ago, a small mass on the right side of the nasal vestibule that caused partial nasal obstruction. She was referred to the surgical outdoor by her general practitioner where she underwent surgical resection of tumour. Postoperatively, after about 3 weeks she observed reappearance of the tumour which aggressively increased in size resulting in about 3 cm cauliflower shaped irregular mass, this time involving the left vestibule and the nasal septum.

Patient at the time of admission exhibited extensive nasal crusting with clinical symptoms like nasal congestion, pain on touching without bleeding, sensation of increased pressure in the nose. There had been no trauma prior to her complaints and she had no allergies. There were no history of substance abuse and no other general complaints. Patient had a history of travel to U.S.A 5 years back.

ENT examination revealed an extensive amount of friable granular tissue along the left side of the nasal vestibule with infiltration of nasal septum, columella and skin of the nose. There were no other abnormalities detected during ENT and general physical examination and laboratory blood tests were as shown in below:

Laboratory findings on admission and on discharge (in brackets)

Peripheral blood

White blood cells 6400/L (6600), Red blood cells 4.95 x 10^6 (4.86), Hemoglobin 14.1g/dl (13.7), Platelets 26.7x10^4/L (14.5), Hematocrit 43.1 (42.2)

Biochemical data

Na 145mEq/L (142), K 4.4mEq/L (4.5), Blood urea nitrogen 28 mg/dl (30), Creatinine 0.8 mg dl (0.64), Uric acid 3.6 mg/dl (3.8), Aspartate transaminase 25U/L (28), Alanine aminotransferase 23U/L (20), Lactate dehydrogenase 629 U/L (655), Alkaline phosphatase 104 (92), GGTP 21 (21), Bilirubin 8.8 (14), Glucose 89 (87), Total protein 73, D-dimer 817

Immunohistochemical staining was consistent with nasal T cell lymphoma. It showed that the infiltrate consisted of T cells (CD3, CD30 positive). The B cell markers CD 20 were negative. Cytotoxich markers like CD56 were negative. Anticytokeratin antibodies which serve to identify micrometastases to lymph nodes were also negative (CKAE1/CKAE3 negative). Ki67 was + in 80% cells showing very aggressive nature of the tumour. CT of the head and neck revealed a localized mass over the nasal columella infiltrating nasal vestibules bilaterally without any involvement to the lymph nodes. CT of the chest to pelvis was negative for tumour involvement. Accordingly, the patient was diagnosed as stage IE nasal T cell lymphoma.

We qualified patient for the radical treatment with radiotherapy alone without preceding CHOP (Cyclophosphamide, doxorubicin, Vincristine and Prednisolone) regimen. IMRT was delivered using 6-15MeV linear accelerators at 2 Gy per daily fraction for a total dose of 44 Gy. The radiation therapy alone resulted in CR (complete remission). During treatment patient had experienced dryness of nasal cavity resulting in severe nasal crusting. Patient was treated with vitamin E nasal drops and ointment. She also had inflammatory changes in buccal cavity and oropharyngeal fungal infection. Accordingly was treated with antifungal and antibacterial preparations.

CT of nasopharynx (with contrast) which was done at the end of treatment revealed that we had achieved complete remission
of the lymphoma. According to the recommendations of NCI post follow up investigations i.e. routine blood tests, ENT examination and follow up CT of nasopharynx were done every 4 weeks in our institute. Unfortunately patient died after four years due to distant metastasis post palliative chemotherapy.

Discussion

Non-Hodgkin’s lymphoma (NHL) is a heterogeneous group of malignancies of the lymphoid system characterized by an abnormal clonal proliferation of B cells, T cells or both. Presented here is a rare entity of NHL - the Nasal NK/T cell lymphoma. Nasal natural-killer- (NK)/T-cell lymphoma is an extranodal lymphoma of the nasal cavity and nasopharynx. This disease for a number of years was believed to be caused by uncertain cause but is now believed to represent a T cell lymphoma. It has been named differently in the past as lethal midline granuloma, midline malignant reticulosis, polymorphic reticulosis, angiocentric lymphoma (in the revised European-American (REAL) classification), extranodal NK/T-cell lymphoma, nasal and nasal type (in the new World Health Organization classification). However for staging NHL Ann Arbor classification is applied routinely [1-5].

The prevalence of nasal lymphomas is estimated at 0.17-1.5% for all non-Hodgkin’s lymphomas (NHL), of which 45% are thought to be of origin from NK/T cell lymphomas of nasal cavity are seen more commonly in Asia (China, Hong Kong, Taiwan), Mexico, and in native population in Peru. A consistent association with EB virus infection is demonstrated in the lymphoma cells, suggesting a probable pathogenic role of EBV. The disease affects men more than women. The age of incidence is generally in the fifth decade. It is characterized by poor prognosis with an aggressive clinical course involving unrelenting ulceration and necrosis. Extra nodal sites invariably involved include nose, palate, upper airway, GIT and skin. Disease appears to spread locally with only a small predilection for regional or systemic failure. International Prognostic index developed by 16 different institutes of U.S., Europe and Canada is used which includes adverse factors like age >=60 years, Ann Arbor stage III/IV, serum LDH level above normal, number of extranodal sites of involvement >=2, performance status >=ECOG 2 or equivalent [2,3,6,7]. Treatment approaches include Radiotherapy alone, chemotherapy alone or the combined modality treatment with the two different sequence of Chemotherapy which could prove beneficial in these cases i.e. either chemotherapy after Radiotherapy or alternate scheduling of RT and Chth in contrast to induction chemotherapy followed by RT in other head and neck lymphomas. High dose Chemotherapy is indicated for patients with stage III-IV and patients with local recurrence or systemic tumor spread after locoregional radiotherapy. Intrathecal methotrexate can be considered in the case of paranasal sinus and orbit involvement to decrease the risk of leptomeningeal spread [1,2,8]. In a recent study L-asparaginase was an effective option to improve the chemotherapeutic efficacy on NK/T cell lymphoma, nasal type, and is worth further study. There have been some reports that high dose Chth and autologous or allogeneic bone marrow transplantation has been successful for NK/T-cell lymphomas, and recently Sasaki et al. [9] reported that double auto-PBSCT achieved CR in a therapy resistant nasal NK/T-cell lymphoma [5,8,10] (Figure 1 & 2).

Conclusion

Extranodal NK/T cell nasal-type pursues an aggressive clinical course with poor prognosis. Several recent clinical investigations have reported 5-year overall survival (OS) rates ranging from 36% to 49.5% for patients with stage I-IV disease. Up to now, optimal treatment strategies have not been fully recognized. Although radiotherapy and chemotherapy are both effective for NK/T cell nasal-type lymphoma, approximately 50% of the patients still fail in locoregional recurrences or systemic disease progression. Thus, an innovative therapy is urgently needed. However, immunohistochemical staining has enabled a better estimation of disease course and prognosis but despite these improvements high index of clinical suspicion is imperative to ensure early diagnosis and ultimately improve disease outcome [11-15].

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