A Case of Primary Aggressive Non Hodgkin’s Lymphoma of Parotid Gland

Abstract

Introduction: Primary parotid Non Hodgkin lymphoma is rare and is generally overlooked in differential diagnosis. A high index of suspicion is therefore warranted for the diagnosis. In the present case report, we describe a case of high grade diffuse large B cell Non-Hodgkin’s lymphoma of the parotid gland.

Case presentation: A 24 Yrs old male non smoker without any known co-morbidities presented with painless swelling of left parotid region of four months duration which was insidious in onset and gradually progressive. There was no history of any constitutional symptoms, facial nerve involvement and rapid increase in the size of swelling. There was no other significant family or personal history. On examination, the swelling was 5 x 4 cm, firm in consistency and fixed to the overlying skin just below the pinna and adherent to upper part of sternomastoid muscle. There was puckering of the overlying skin in the centre with surrounding areas of erythema and dark pigmentary changes. The facial nerve was intact. No other abnormality was found in the oral cavity, nasopharynx, larynx or the ears. There was no palpable enlargement of cervical lymph nodes. Routine biochemical lab tests were within normal limits (Figures 1 & 2).

Conclusion: Lymphomas arising from the parotid gland are an uncommon entity, said to account for only 0.6–5% of tumors or tumor-like lesions of the parotid, and are therefore commonly overlooked. This misdiagnosis often leads to unnecessary diagnostic procedures, delaying the initiation of proper treatment.

Keywords: Non Hodgkin’s Lymphoma; Parotid gland; Diffuse large B cell lymphoma; Tumors; Parotidectomy; Dorsolumbar spine; Follicular lymphoma; B-cell lymphoma; Chemotherapy; Radiotherapy; Cervical lymph nodes; Erythema; Malignant; Oral cavity; Nasopharynx; Larynx; Ear; Swelling; Hypo intense lesion; Ultrasonography; Preauricular skin incision; Submandibular glands; Pigmentation

Abbreviations: DLBL: Diffuse Large B Cell Lymphoma; MALT: Mucosa Associated Lymphoid Tissue; FNAB: Fine Needle Aspiration Biopsy; CECT: Contrast Enhanced Computed Tomography

Introduction

Primary lymphomas arising from the parotid gland are uncommon and usually account for only 0.6–5% of tumors or tumor-like lesions of the parotid, and therefore are commonly overlooked. Most cases involve the major salivary glands, primarily the parotid (50–93%) and the submandibular glands [1]. The misdiagnosis often leads to unnecessary diagnostic procedures thereby delaying the initiation of appropriate treatment. It is a common knowledge that it is often difficult to make distinction between lymphoma developing primarily in the parotid gland tissue or de novo in the intra-parotid lymph nodes. It has been reported that primary tumours of the parotid gland show no characteristic features on diagnostic imaging, reflecting none of their histological findings [2]. The prognosis and morphology, however, is more or less similar between the two origins [3]. In our case the main tumour was arising from the parenchyma of the deep lobe of the parotid gland. Most cases of non-Hodgkin’s lymphoma arising in salivary glands are of B-cell lineage including low-grade B-cell lymphomas of mucosa-associated lymphoid tissue (MALT), diffuse large B-cell lymphomas and follicular lymphomas [4]. It is difficult to differentiate between benign conditions from malignant tumours and therefore most patients are subjected to varied surgical procedures before a definitive diagnosis has been made. The head and neck region is the most common site where malignant lymphomas occur, but malignant lymphoma of the parotid gland is relatively rare [5]. Generally, surgeons do not anticipate primary Non Hodgkin’s Lymphoma (NHL) in the salivary glands pre-operatively and pathologists too find it difficult to give a definitive and diagnostic report based on either frozen section or fine needle aspiration biopsy (FNAB) [6]. In contrast to other extra nodal locations of NHL, parotid gland involvement is more likely to be of low grade and the patients have a better prognosis [7]. However, in our case it was of high grade. The treatment of primary parotid lymphoma includes chemotherapy and may be radiotherapy. Surgical excision is advocated only as a diagnostic tool.

Case Report

A 24yr old male non smoker without any known co-morbidities presented with painless swelling of left parotid region of four months duration which was insidious in onset and gradually progressive. There was no history of any constitutional symptoms, facial nerve involvement and rapid increase in the size of swelling. There was no other significant family or personal history. On examination, the swelling was 5 x 4 cm, firm in consistency and fixed to the overlying skin just below the pinna and adherent to upper part of sternomastoid muscle. There was puckering of the overlying skin in the centre with surrounding areas of erythema and dark pigmentary changes. The facial nerve was intact. No other abnormality was found in the oral cavity, nasopharynx, larynx or the ears. There was no palpable enlargement of cervical lymph nodes. Routine biochemical lab tests were within normal limits (Figures 1 & 2).
A total parotidectomy with preauricular skin incision was planned. However, intra-operatively involvement of surrounding muscles, periparotid lymph nodes and level IIb / III lymph nodes was noted. Facial nerve and its terminal branches were preserved with careful dissection; however there was adherence of inferior division of facial nerve to the tumour mass which was cleared carefully under microscopic control. The retromandibular vein was ligated and a globular mass seen arising from the deep lobe of the parotid. The tumour was removed in to along with other smaller mass from the deep lobe. Total parotidectomy was performed along with excision of unhealthy overlying skin, part of upper and anterior border of Sternocteolomastoid muscle and a part of posterior belly of Digastric muscle. Level IIb and III lymph node clearance was done. A suction drain was put for 48 hours. Post operative recovery was uneventful with minimal deviation of angle of the mouth which recovered fully after two weeks. Histopathology of the mass revealed diffuse large B cell NHL showing sheets and strands of atypical lymphoid cells with involvement of superficial, deep lobe of parotid, Sternocteolomastoid muscle and Digastric muscle. Skin and level IIb and III lymph nodes also revealed similar population of atypical cells with perinodal extension. The tumour cells were positive strongly for CD20, LCA and CD79a on Immunohistochemistry (IHC). However, the cells were negative for CD5, CD3, EMA, CK and TdT on IHC. The Ki-67 labelling index was 100% (Figures 3-8).

The patient complained of low back ache in the workup period for which a dorso-lumbar spine X-Ray was obtained which was normal. A Contrast enhanced computed tomography (CECT) chest and abdomen was advised keeping a high index of suspicion in view of a high grade DLBCL. The CECT revealed multiple ill defined lytic lesions noted in the bodies of C7, DV4-DV6, DV10, DV12, LV2, SV1 and bilateral iliac bones. Also, a compression fracture of DV4 was noted. Rest of the metastatic workup including the bone marrow study was normal. With the clinical stage IV disease, patient was started on RCHOP chemotherapy regimen. As part
of the RCHOP regimen the patient received eight cycles of Inj Rituximab (375mg/m2), Inj Cyclophosphamide (750mg/m2), Inj Doxorubicin (50mg/m2), Inj Vincristine (1.4mg/m2) and Tab Prednisolone 40mg/m2/day for 5 days. The chemotherapy was well tolerated. Presently, the patient is disease free after one year and is on regular follow up (Figures 9 & 10).

Figure 4: Intraoperative image showing: Deep lobe tumor of parotid being delivered, sternomastoid muscle, facial nerve and Spinal accessory nerve.

Figure 5: Under microscopic control the adherent lower division of facial nerve was freed from the deep lobe tumour.

Figure 6: Resected specimen: Superficial lobe of parotid, deep lobe of parotid, parotid tumour mass, lymph nodes, skin, part of SCM and post belly of digastrics muscle.

Figure 7: HPE Specimen (H&E, x200) - Infiltration of salivary gland parenchyma with atypical lymphoid cells having high N: C ration and moderate degree of nuclear pleomorphism.

Figure 8: Postoperative image after removal of deep lobe and superficial lobe of parotid showing accessory parotid remnant, sternomastoid muscle, internal jugular vein, retromandibular vein, spinal accessory nerve and facial nerve.

Figure 9: Post operative picture of the patient and after the completion of chemotherapy with complete recovery of angle of mouth weakness.

**Discussion**

Most of the extra nodal lymphomas of the head and neck are NHL and in 4%-5% of cases the parotid gland is involved [8,9]. Non-Hodgkin’s lymphoma of the parotid gland may be classified as extranodal if the origin is from the mucosa associated lymphoid tissue (MALT) or nodal if the true origin is from lymph node within the gland. The lesions, in the opinion of few authors, arise from the intra-parotid lymph nodes associated within the gland and involving the gland parenchyma secondarily. The theory makes it difficult to establish the true origin of the disease [10]. Although considered insufficient by some authors [11], Hyman & Wolff [12] have suggested some criteria to consider a lesion as primary parotid gland lymphoma:

I. Involvement of the gland should be the first clinical manifestation of the disease.

II. Histological, the disease should involve the gland parenchyma and not adjacent lymph nodes or soft tissue alone.

III. There should be confirmation of the malignant nature of the lymphoid infiltrate.

Parotid lymphoma most commonly presents as a painless mass indistinguishable from other non-malignant or other more common epithelial tumors. This explains why this diagnosis is commonly overlooked and patients are often subjected to unnecessary procedures and a delay in diagnosis. In most cases the facial nerve is not jeopardized [13]. Diagnosing parotid lymphoma can be a difficult task, as evidenced by the unnecessary radical operations to the patient with all its associated risks involved. Also, attention is drawn towards the warranted metastatic workup keeping a high index of suspicion in a case of high grade DLCBCL of the parotid gland.

**Conclusion**

This case has been described to highlight the relatively rare presentation of primary NHL of the parotid gland. This difficulty in pre and intra-operative diagnosis often results in unnecessary radical operations to the patient with all its associated risks involved. Also, attention is drawn towards the warranted metastatic workup keeping a high index of suspicion in a case of high grade DLCBCL of the parotid gland.

**References**