LANGERHANS CELL HISTIOCYTOSIS: A CASE REPORT AND UPDATE ON TREATMENT MODALITIES


ABSTRACT
Various treatment modalities exist for Langerhans cell histiocytosis(LCH) due to its uncertain aetiology. LCH patients are clinically divided into a single and multisystem group and each group warrants a different approach to treatment. However, histologically, CD1a antigen and Birbeck's granules have been implicated in both types of the disease. This article therefore describes clinical presentations of different forms of the disease and the treatments contrived for them. The single system group is treated conservatively with either curettage of the localised lesion, intralesional steroid injection or in some cases radiotherapy. For the multisystem group, treatment consists of chemotherapeutic agents as either a single or combination therapy.

Key Words: Langerhans cell histiocytosis, Letterer-Siwe disease, Birbeck's granules.

INTRODUCTION
Langerhans cell histiocytosis(LCH) have been an enigma since its discovery by Hand in 1893 (1). It remained so for a very long time due to its wide clinical presentations, variable histopathological features and an uncertain aetiology. It was Lichtenstein in 1953 (2) who analyzed the range of clinical and pathological variations and concluded that the malady of diseases were in fact a single entity and termed it "Histiocytosis X". Histiocytosis for the proliferation of histiocyte-like cells while "X" denotes the unknown aetiology. Nezelof in 1973 (3) was first to identify that these proliferative histiocytes were Langerhans cells. Subsequent developments in immunocytochemistry have positively proven this by the discovery that these histiocytes share a common surface antigen with normal Langerhans cells, the CD1a antigen (4). It is now accepted that Langerhans cell histiocytosis (LCH) is the most appropriate term to refer to the above disease.

CLINICOPATHOLOGICAL PRESENTATIONS
Little is known of the epidemiology of LCH primarily because it is a relatively rare disease. However Cartensen and Ornvold (5) found that the incidence rate of LCH is 1.08 in 200,000 in a survey involving a total population of 1 million children under the age of 15 in Denmark. The incidence of adult however is more difficult to fathom due to wider clinical presentations, often symptomless and under reporting by clinical specialists.

LCH is classified traditionally into 3 distinctive clinical presentations. The mildest of which is termed eosinophilic granuloma where symptoms of solitary or multiple bone lesions are usually reported. The second is the chronic disseminated form of LCH which is clinically known as Hand-Schuller-Christian syndrome characterized by a triad of symptoms which includes intranasal lesion, exophthalmos and diabetes insipidus. The third is the acute or subacute form of the disease which is known as Letterer-Siwe syndrome. It is characterized by rapid onset with widespread skeletal and extra-skeletal lesions. It occurs usually in infants less than 3 years of age with severe clinical course and sometimes results in a fatal outcome. Not all cases however satisfy the above criteria. Lieberman (6) proposed a simpler classification in which LCH is classified as having either a unifocal or multifocal entity.

The histological picture is possibly the common denominator of the above diseases. Under light microscopy, the lesion is often described as an aggregation of histiocytic like cells arranged in a sheet like fashion superimposed with eosinophils in a fibrous connective tissue matrix. Under electron microscopy, Birbeck's granules are characteristically present. Immunohistochemistry would show a positive labelling for CD1a surface antigen.

CASE REPORT
A two year old Chinese boy was referred from Kuching General Hospital to both the Paediatric Department, Faculty of Medicine and the Oral and Maxillofacial Surgery Department, Faculty of Dentistry, University of Malaya, Kuala Lumpur, Malaysia in February 1992. The figure 1: Shows cervical lymphadenopathy
patient suffered recurrent bouts of fever and cervical lymphadenopathy. He was initially admitted to the Kuching General Hospital and treated with antibiotics but the bouts of fever persisted. He then went on to develop hepatosplenomegaly. At this point he was transferred to Kuala Lumpur for further assessment.

He is the youngest of nine siblings. His father is of Chinese origin while the mother is of Iban extract. No known medical problems of other members of the family were found. On examination we found that he had persistent and recurrent bouts of pyrexia, bilateral cervical lymphadenopathy (Fig. 1), hepatosplenomegaly (Fig. 2) and seborrhoic dermatitis of the scalp. Intraorally we found areas of granulomatous like lesion of the hard palate and the deciduous teeth were very mobile (Grades 2 and 3). Radiographic investigations were ordered - a Posteroanterior (P.A.) skull radiograph and lateral cephalogram revealed a poorly defined radiolucency of the maxilla. The teeth showed characteristic “floating in air” appearance (Fig. 3). A full body skeletal survey, bone scan with TC 99 MDP showed “hot spots” localized to the maxillary bone. No other bone were found to be affected. A provisional diagnosis of Letterer-Siwe disease was considered most consistent to the above history and examination. However the occurrence of lymphoma was not ruled out. Biopsies were then taken from the palate and cervical lymph nodes which subsequently confirmed the diagnosis of LCH.

A full dearance of all deciduous teeth was done under general anaesthesia because it was decided that they could be a source of infection or the teeth may exfoliate and there is this risk of their being aspirated. The disease was treated with a single course of VP16 (Etopipodophylotoxin Etoposide). Despite treatment, the patient died 4 months later from the disseminated form of the disease.

DISCUSSION
The treatment of LCH is somewhat a controversial issue. There are many variations amongst clinicians to their approach of treatment modalities. This is perhaps due to the uncertain etiology of the disease. Recent developments have shown that LCH is a clonal disorder (7) suggesting that it may be neoplastic in origin. The variability of clinical manifestations from symptomless single system lesion to that of a life threatening multiple system disorder thus complicates the management even further.

Current approaches to the management of LCH is to divide the lesion into either a single or multiple system disease. A single system disease usually affect a single organ i.e. bone, lymph nodes. It is usually benign in nature and remission is likely. Therefore treatment for this disease has been more conservative. Curettage of the bone will usually lead to spontaneous remission.

Jones et al (8) and Ong et al (9) in their respective case reports have shown that intralesional infiltration of steroid ranging from 40 to 200 mg methylprednisolone have been successful in the resolution of the isolated osteolytic lesion of the mandibular bone.

The use of radiotherapy have gradually gone out of favour amongst clinician possibly due to better understanding of the nature of the disease and the risk (5%) of second tumour development (10). However, the use of low dose radiation is justifiable if LCH threatens critical organs such as lesions encroaching upon the optic nerve or the spinal cord where local infiltration of corticosteroid may not be possible.

In the multisystem disease, 2 major approaches exist - a conservative approach involving minimal therapy and a more intensive chemotherapy plan. Based on observation by Broadbent (11), minimal therapy will lead to spontaneous recovery for chronic form of LCH. However,
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evidence of fever, pain, immobility, failure to thrive and worsening condition of vital organs may indicate for a more aggressive approach (12). This may include the administration of methylprednisolone (60mg/m2/day) given for a short period of time and followed by VP16, vincristine or vinblastine.

Another alternative approach stipulated by the German-Italian prospective cooperative multicentre study group, the AIEOP-CNR-HX 83 (13), divides patients with multisystem disease into good and poor prognostic groups according to the absence or presence of organ dysfunction at the time of diagnosis. Treatment of good prognosis patients is to administer single agents sequentially to obtain favourable response. These are vinblastine (6.5 mg/m2 i.v. weekly for 3 months) followed by doxorubicin (20 mg/m2 i.v. for 2 consecutive days every 3 weeks for 3 months) and finally VP16 (200 mg/m2 i.v. for 3 consecutive days, 3 weekly, at least for 3 months). For poor risk group, they will receive 9 courses of a 4 week treatment with a combination chemotherapy including vincristine, cydophosphamide, doxorubicin and prednisolone. This study concludes that response for single agents were effective in patient with good prognosis. In poor risk cases however, prognosis remain bad despite multi-agent therapy.

With greater understanding of the disease process researches have tried other approaches in the treatment of LCH. The Histiocyte Society have also established the Salvage Therapy Group to propose a study for the use of experimental drugs when the normal regimen have been proven ineffective (14). These approaches include the use of cyclosporin A (15) on the rationale that selective inhibition of the cellular immune response and cytokine mediated cellular activation would suppress the immunoregulatory effect of the disease. Alpha interferon have also been used in the hope of increasing NK cytotoxic activity. There is a case report of success in using this approach (16). However, no prospective clinical trial, as yet, have been initiated to show the efficacy of this form of treatment.

Bone marrow ablative therapy with bone marrow transplant (BMT) in the recurrent multisystem disease have been tried out in clinical practice A study (17) has reported some degree of success but the mortality rate due to either the recurrence of disease or the side effect of treatment is still unacceptably high.

Monoclonal antibody therapy of LCH patients exploits the potential role of monoclonal antibody directed against the CD1a surface antigen of Langerhans cells. Kelly and Pritchard (18) were first to investigate the possibility of such treatment. The clinical application potential of this technique is immense either through a cytotoxic or immunoregulatory mode of treatment.

CONCLUSION

Langerhans cell histiocytosis have puzzled clinicians and scientists alike. Recent scientific breakthrough have shown that it is of neoplastic origin rather than a reactive disease. Clinical diagnosis may be difficult and often it is the dental surgeon who are first in line to notice this disease. LCH may mimic a wide range of clinical entities. It may present itself as juvenile or refractory periodontitis. It may be present as diabetes insipidus or a clinical condition similar to lymphoma. Therefore suspicious localized or generalized periodontal pocketing or bone loss must be investigated and the differential diagnosis of LCH must not be discounted.

Treatment of LCH is a contentious issue. However the consensus is to divide patients into single or multiple systems group. The single system group is treated more conservatively with either curettage of the localised lesion, intralungal steroid injection or in some cases radiotherapy, if the lesion threatens any vital organs. For the multisystem group, a more comprehensive approach is implemented primarily utilising VP16, vincristine or other chemotherapy agents as either a single or combination therapy. It must be pointed out that in situations where there is organ dysfunction, prognosis remain poor despite utilising all chemotherapy options. Novel therapeutics offer new potentials by opening new fronts in combatting LCH. This is only possible by further research in the quest for better understanding of this disease.

Acknowledgement

The authors wish to thank Sister Rathiyah Ahmad and her staff for their sincere efforts in assisting in the clinical work and the UMNO club of London for its facilities.

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